

## **Vertical gradients in physiological function and resource allocation of white spruce diverge at the northern- and southern-most range extremes**

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## ABSTRACT (200 words)

Light availability drives vertical canopy gradients in photosynthetic functioning and carbon (C) balance, yet patterns of variability in these gradients remain unclear. We measured light availability, photosynthetic CO<sub>2</sub> and light response curves, foliar C, nitrogen (N) and pigment concentrations, and the photochemical reflectance index (PRI) on upper and lower canopy needles of white spruce trees (*Picea glauca*) at the northern and southern extremes of the species' range. We combined our photosynthetic data with previously published respiratory data to compare and contrast canopy C balance between latitudinal extremes. We found steep canopy gradients in irradiance, photosynthesis, and leaf traits at the southern range limit, but clear convergence across canopy positions at the northern range limit. Thus, unlike many tree species from tropical to mid-latitude forests, high latitude trees do not require vertical gradients of metabolic activity to optimize photosynthetic C gain. Consequently, accounting for self-shading is less critical for predicting gross primary productivity at northern relative to southern latitudes. Northern trees also had a significantly smaller net positive C balance than southern trees suggesting that, regardless of canopy position, low photosynthetic rates coupled with high respiratory costs may ultimately constrain the northern range limit of this widely distributed boreal species.

SHORT TITLE: Variation in photosynthetic canopy gradients of white spruce at the species' range limits.

KEY WORDS: *Picea glauca*, canopy gradients, carbon balance, photosynthesis, photochemical reflectance index, Arctic treeline

## SUMMARY STATEMENT

Canopy gradients in photosynthetic capacity of white spruce diminish at high compared to low latitudes. Low carbon balance in high latitude trees may determine the extent of northern treeline.

## INTRODUCTION

Illuminating the factors driving species' distributional ranges, i.e., the boundaries within which a species can grow and reproduce successfully, remains a fundamental ecological, evolutionary, and biogeographic challenge. Extensive research has uncovered a number of factors that play important roles. These include environmental tolerance, niche breadth, metapopulation dynamics, genetic diversity, phenotypic plasticity and dispersal ability (Brown, Stevens & Kaufmann 1996; Gaston 1996; Lowry & Lester 2006). The diversity of possible factors highlights that no universal cause of species' distributions has yet emerged, but rather that a combination of historical, physiological, and biotic filters may define a species' distributional range (Lambers & Oliveira 2019). Nevertheless, the ability of a plant to tolerate the local growth environment remains paramount. This ability is intimately linked to a species' physiological performance. The physiological processes of photosynthesis and respiration, resulting in the exchange of carbon, are heavily affected by environmental conditions. Together, these two processes determine the carbon balance of a plant which must remain positive if a plant is to grow, survive and reproduce in a given environment. Thus, photosynthesis and respiration fundamentally act as a physiological filter determining a species' distributional range (Lambers & Oliveira 2019). They are also critical to quantifying carbon exchange as well as the ecological effects of climate change on future carbon sequestration.

The evolution from modeling vegetation as a single “big leaf” to “two big leaves” including distinct parameters for sun and shade leaves, to multi-layer canopies including gradients of physiological function (Field 1983; Sands 1995) has improved estimates of vegetative carbon gain in many ecosystems (Harley & Baldocchi 1995; de Pury & Farquhar 1997; Bonan, Oleson, Fisher, Lasslop & Reichstein 2012; Rogers *et al.* 2017). These multi-layer canopy models are parameterized using physiological and biochemical traits that vary predictably with vertical gradients in environmental conditions such as light, temperature, humidity and vapor pressure deficit throughout the tree crown (Sellers, Berry, Collatz, Field & Hall 1992; Bond, Farnsworth, Coulombe & Winner 1999). Of particular importance to modeling photosynthesis are gradients in light availability. For example, traits such as maximum photosynthesis ( $A_{\max}$ ) and leaf nitrogen content, as well as leaf mass per area (LMA), have been found to correlate positively with changes in irradiance (Field 1983; Hirose & Werger 1987; Ellsworth & Reich 1993; Niinemets, Kull & Tenhunen 1998; Niinemets 2007) leading to the

common finding that high canopy, high light foliage has higher photosynthetic capacity. Pigment pools have similarly been found to vary with canopy position and irradiance with high light foliage frequently containing larger pools of photoprotective pigments that can dissipate excess light energy and protect chlorophyll-filled photosystems (Demmig-Adams & Adams III 1992). Even the ratio of Chla:Chlb frequently reflects canopy gradients of light availability because shade-adapted leaves contain more light harvesting antenna complexes (LHCII) associated with photosystem II that are high in Chlb (Dale & Causton 1992; Hikosaka & Terashima 1995; Kitajima & Hogan 2003; Ruban 2015; Magney *et al.* 2016; Scartazza, Di Baccio, Bertolotto, Gavrichkova & Matteucci 2016). Thus, the incorporation of canopy gradients into carbon exchange modeling comes from a strong foundation of work that demonstrates how differences in photosynthetic functioning and resource allocation depend on canopy position and light availability. At the same time, we note that this foundation is born from a wealth of studies on temperate and tropical forests and tree crowns that have inherent vertical gradients in irradiance, as well as physiological and biochemical traits. As irradiance and photoperiod also vary latitudinally, we question whether these assumptions hold true when a species' distribution spans a large latitudinal distance.

White spruce (*Picea glauca* (Moench) Voss) has a massive, transcontinental range distribution stretching from the Forest Tundra Ecotone (FTE) in northern Alaska where it is one of the dominant species, to mixed forests on the eastern seaboard of Canada and New England (Viereck, Van Cleve & Dyrness 1986; US Geological Survey 1999; Eitel *et al.* 2019). At the northern edge of the distribution, white spruce are exposed to unique environmental challenges for plant growth and survival including low winter temperatures, shallow soil depth underlain by permafrost, and brief growing seasons. During the growing season, white spruce canopies are exposed to a unique light environment caused by high solar zenith angles and continuous photoperiod (i.e. 24 hours of light for several weeks). Furthermore, tree crown structure at the FTE exhibits an almost ubiquitous tall, vertical, narrow and untapering crown shape (Kuuluvainen 1992) which creates an open canopy forest structure with widely spaced trees. By contrast, white spruce at the southern range extreme are exposed to much lower solar zenith angles and a shorter photoperiod during the growing season. The southern tree crowns are also generally much denser, wider and more tapered, and typically grow with a higher stand density than in the north. These contrasting light environments may impact canopy gradients of light

interception, which may, in turn, drive canopy gradients of photosynthetic physiology, biochemistry and pigment concentrations. In addition to photosynthetic, biochemical and pigment traits, we also examine canopy differences in the photochemical reflectance index (PRI). This index is a more successful photosynthetic proxy than the commonly used normalized difference vegetation index (NDVI) in conifer species because it is sensitive to changes in pigment pools such as the ratio of chlorophylls to carotenoids (Gamon *et al.* 2016). PRI has been successfully employed to track seasonal changes in photosynthetic phenology in coniferous forests (Wong & Gamon 2015a, b; Gamon *et al.* 2016) and at the FTE (Eitel *et al.* 2019, 2020), and a broadband analog Chlorophyll Carotenoid Index (CCI), can be derived from global Moderate Resolution Imaging Spectroradiometer (MODIS) imagery (Gamon *et al.* 2016). Thus, PRI presents a promising prospective avenue for further understanding carbon dynamics across broadly distributed conifer forests.

In this study we focus specifically on the photosynthetic traits of white spruce and their relationship to light environment. However, to gain a full picture of the physiological limits at latitudinally distant range extremes, a carbon balance approach should be employed. In fact, recent work on the physiology of white spruce has focused specifically on the respiratory cost associated with growth and survival at the FTE. In a previous study, we found that white spruce in this environment have extremely high respiration compared to white spruce growing at the southernmost range extreme as well as higher respiration than the average respiration of the boreal biome and needle-leaved evergreen trees (Griffin *et al.* 2021). Griffin *et al.* (2021) posit that this high respiratory cost may constrain the northern limit of white spruce and the Arctic treeline in general. With the addition of photosynthetic traits measured on the same trees, we are now in a unique position to examine the degree to which this respiratory cost is matched by photosynthetic carbon gains and whether a lower carbon balance may contribute to the northern limits of white spruce's latitudinal range.

Thus, the overarching aims of this research are twofold: first, to provide a comprehensive assessment of the photosynthetic physiology of white spruce and its relationship to canopy and latitudinal light environment at its range extremes; and second, to examine whether the carbon balance of this species constrains its northernmost range limit. First, we hypothesize that foliar light availability, physiological traits, biochemical traits and PRI will converge across high and low vertical canopy positions in white spruce growing at the FTE. In contrast, steep canopy

gradients of light availability and all foliar traits will be apparent at the southernmost range extreme. Second, we hypothesize that the carbon balance of white spruce at the northern location will be much lower than the southern location reflecting the challenging growth environment and providing a physiological mechanism for the northernmost range extreme of this important boreal species.

## MATERIALS AND METHODS

*Site Descriptions:* This study was conducted in two locations at the northern and southern range extremes of white spruce (*Picea glauca* (Moench) Voss). In the north, data were collected from six sites chosen to represent the FTE (Eitel *et al.* 2019). These sites are located along a 5.5 km long stretch of the Dalton Highway, in the Brooks Range in northern Alaska (67°59'40.92" N latitude, 149°45'15.84" W longitude). At the FTE, white spruce is the dominant tree species with deciduous shrubs and sedges present in the understory (Eitel *et al.* 2019). Three white spruce study trees (with DBH greater than 10cm) were chosen at each site for a total of 18 study trees at the FTE (tree characteristics are described in Table 1). Mean annual precipitation was 485.4 mm and mean annual temperature was -8.1°C as determined from a SNOTEL site at the nearby Atigun Pass (<https://wcc.sc.egov.usda.gov/nwcc/site?sitenum=957>) between 2007 and 2016. During the Alaska measurement campaign in July 2017, photoperiod ranged from 22 to 24 hours. All measurements were taken at a high and low south-facing canopy position on each of 18 trees for a total of 36 measurements. The low canopy position was at approximately 1.37 m (diameter at breast height), and the high canopy position was approximately 1m below the apical meristem.

At the southern edge of the species' range, data were collected from six trees located in Black Rock Forest, New York (BRF; 41°24'03.91" N latitude, 74°01'28.49" W longitude; Table 1). Mean annual precipitation from 1982 – 2010 was 1285 mm and mean annual temperature was 10.9°C (Arguez *et al.* 2012). During our BRF measurement campaign in June 2017, photoperiod was approximately 15 hours. BRF is an oak-dominated northern temperate deciduous forest (Schuster *et al.* 2008; Patterson *et al.* 2018). Measurements were also taken at high and low south-facing canopy positions on each tree. Due to the lower count of available trees, three measurements were taken at the low and high canopy positions on each of the six trees for a total of 36 measurements. As at the FTE, the low canopy position was at

approximately 1.37 m and the high canopy position was approximately 1m below the apical meristem.

*Environmental Measurements:* Hemispherical photographs were taken with a digital camera (CoolPix 4500, Nikon Corporation, Tokyo Japan) and an attached fisheye lens to assess canopy openness at the high and low canopy positions chosen for each tree (described above). The camera was positioned immediately adjacent to the branch chosen for measurements, with the camera pointing north. Once the camera was level, the hemispherical canopy photo was taken. Using these photographs, light environment was assessed and modeled using the free R code and documentation of ter Steege (2018). These scripts are based on the widely used Hemiphot and Winphot programs (ter Steege 1993, 1997). Modeling of the light environment using this code takes into account GPS location as well as the measurement dates in order to correct for differences in solar angle (see ter Steege (2018) for a fuller description). Average Photosynthetic Photon Flux Density (PPFD ; mol m<sup>-2</sup> day<sup>-1</sup>) at the high and low canopy positions was calculated by averaging the diurnal light courses over the measurement campaign days (June 21 – June 27 at BRF, and July 4 – July 20 at the FTE). At the FTE, air temperature (°C) was collected using a meteorological sensor (VP-4, METER, Pullman, WA) and local solar open sky radiation readings were collected using a PYR Solar Radiation Sensor (METER, Pullman, WA). Solar radiation (W m<sup>-2</sup>) was converted to PPFD (μmol m<sup>-2</sup> s<sup>-1</sup>) by multiplying by 2.02 (Mavi & Tupper 2004). At BRF, air temperature and local open sky PPFD were collected from a nearby BRF-run weather station (<https://blackrock.ccnmtl.columbia.edu/portal/weather/>) equipped with a temperature/humidity sensor (HC2S3, Rotronic, Hauppauge, NY) and a quantum sensor (LI-190SB Quantum Sensor, LICOR, Lincoln, NE).

*Gas Exchange Measurements:* Gas exchange was measured at high and low canopy positions on each tree at the two locations. At both locations, measurements were taken on a branch tip inserted into the cuvette. At BRF, all measurements were taken on the tree with high canopy reached using a tree crane. At the FTE, low canopy branches were measured on the tree, but high canopy branches had to be detached using a pole-clippers as there was no way to reach samples otherwise. For these samples, cut branches were placed immediately in water, and recut underwater. CO<sub>2</sub> (*A-C<sub>i</sub>*) and light response (*A-Q*) curves were collected using two portable



photosynthesis systems (LI-6800, LiCor, Lincoln, NE). Needles were sealed in a cuvette in which the temperature and relative humidity were set to 20°C and 50% humidity at the FTE and 25°C, and 60% humidity at BRF in order to mirror ambient conditions in each location. For the  $A-C_i$  curves, irradiance was kept constant at 1000 PPFD and the CO<sub>2</sub> concentrations were stepped through the following concentrations (ppm): 1200, 1000, 800, 600, 400, 300, 200, 100, 75, 50, 25, 0. Values of  $V_{cmax}$  (the maximum rate of RuBisCO carboxylation) and  $J_{max}$  (the maximum rate of electron transport) were determined according to the photosynthesis model of Farquhar, von Caemmerer and Berry (Farquhar, von Caemmerer & Berry 1980) and adjusted to 25°C using the plantecophys package in R (Duursma 2015).

For the  $A-Q$  curves, reference CO<sub>2</sub> concentration was set to 400 ppm and photosynthesis was measured at the light intensities: 1800, 1500, 1200, 1000, 800, 400, 200, 150, 100, 90, 80, 75, 70, 65, 60, 55, 50, 45, 40, 35, 30, 25, 20, 15, 10, 5, 0  $\mu\text{mol m}^{-2} \text{s}^{-1}$ . After 15 minutes at zero irradiance, respiration in the dark ( $R_D$ ) was recorded. Maximum likelihood estimation (executed in the bblme package for R (Bolker & R Development Core Team 2020)) was used to fit the following model (describing a non-rectangular hyperbola) to the data:

$$A = \frac{\Phi * I + A_{max} - \sqrt{(\Phi * I + A_{max})^2 - 4 * \theta * \Phi * I * A_{max}}}{2 * \theta} - R_D$$

where  $\theta$  is the curvature factor and  $I$  is the irradiance. Parameters extracted and calculated from the fit of the model to each light curve included the photosynthesis at 1500  $\mu\text{mol m}^{-2} \text{s}^{-1}$  ( $A_{1500}$ ), the apparent quantum yield ( $\Phi$ ), the light compensation point (LCP – the light level required for zero net carbon flux) and the light saturation point (LSP – the light level required to obtain 75% of  $A_{1500}$ ).

*Respiration in the Light:* Respiration in the light ( $R_L$ ) was obtained from the  $A-Q$  curves according to the methods of Kok (1948); Sharp, Matthews & Boyer (1984); and Heskell, Atkin, Turnbull & Griffin (2013b). These curves contain a subtle change in slope near the light compensation point (the light intensity where photosynthesis and respiration equal each other). The break point where this change in slope occurs divides the photosynthetic response to light into two linear sections. Extrapolating from the upper linear portion of the curve to the y-axis



highlights an alternate respiration value which was interpreted by Bessel Kok to be the amount of respiration occurring in the light (Kok 1948; Heskell *et al.* 2013; Tcherkez *et al.* 2017).

An important assumption of the Kok method is that carbon assimilation is only influenced by light. Consequently, corrections were made to account for any changes in internal CO<sub>2</sub> concentration ( $C_i$ ) following the methods of Kirschbaum & Farquhar (1987) (see Ayub *et al.* (2011) for a full description). It is possible that  $R_L$  measured with the Kok method may be influenced by factors such as mesophyll conductance (Farquhar & Busch, 2017; *n.b.* Buckley *et al.*, 2017); however, such investigations are beyond the scope of this study.

*Leaf Trait Measurements:* After collecting  $A-C_i$  and  $A-Q$  curves, needles were detached from the branch, photographed to determine projected surface area (cm<sup>2</sup>) using ImageJ (Schnieder, Rasband & Eliceiri 2012), placed into coin envelopes and oven dried at 60°C for 48 hours. Leaves were reweighed post-drying for leaf dry mass (g). Specific leaf area (SLA; cm<sup>2</sup>g<sup>-1</sup>) was calculated. Leaf samples were then ground using a ball mill (SPEX 8000 Mixer/Mill, Metuchen, NJ, USA) and prepared for analysis of % carbon and % nitrogen using a carbon-nitrogen flash analyzer (CE Elantech, Lakewood, NJ, USA). Using these percentages, foliar nitrogen per area ( $N_{area}$ ; mg cm<sup>-2</sup>) and carbon to nitrogen ratio (C:N) were calculated.

*Pigment measurements:* Needles growing on branches adjacent to those selected for gas exchange at both high and low canopy positions were selected for pigment analysis. Branches were wrapped in wet paper towels and foil, and transported back to the laboratory in a cooler with ice. Foliar pigment extraction began no later than three hours after returning from the field, and any samples not processed in this time were frozen until pigment extraction could take place.

Between 5 and 10 needles were removed from each branch and their wet weight and needle area measured. Needles were ground in 100% acetone until no identifiable green needle tissue remained. A small amount of sand, MgCO<sub>3</sub> and ascorbic acid were added to help with grinding and to prevent acidification and pigment degradation. The extracts were then centrifuged and diluted. Using a spectrophotometer (Go Direct SpectroVis Plus Spectrophotometer, Vernier, OR, USA), the absorbance of the supernatant was measured at 470, 645, 662 and 710 nm. Pigment pools were calculated on a leaf area basis including chlorophyll *a* ( $Chla_{area}$ ), chlorophyll *b* ( $Chlb_{area}$ ), and bulk carotenoids ( $Car_{area}$ ; ug/cm<sup>2</sup> for all pools) were

calculated according to Lichtenthaler (1987). The total chlorophyll pool size ( $\text{Chl}_{\text{area}} = \text{Chla}_{\text{area}} + \text{Chlb}_{\text{area}}$ ) and ratios  $\text{Chla}:\text{Chlb}_{\text{area}}$  and  $\text{Chl}:\text{Car}_{\text{area}}$  were also calculated.

*Spectral reflectance: PRI:* Needle spectral reflectance was measured with a spectroradiometer (UniSpec SC, PP Systems, Haverhill MA, USA) with an attached fiber optic probe (UNI400, PP Systems, Haverhill MA, USA) and specialized needle clip (UNI501, PP Systems, Haverhill MA, USA). The spectroradiometer allowed sampling the spectrum between 310 and 1100 nm with a 3.3 nm sampling interval which was afterwards reduced to 1 nm using linear interpolation. After collecting dark and white standard (Spectralon, LabSphere, North Sutton, NY, USA) reference measurements, three spectral measurements were made on each branch to capture spatial heterogeneity. Reflectance values were calculated from the spectral measurements by dividing each foliage measurement by the measurement from the white standard. PRI was calculated according to the equation:  $(R_{531} - R_{570}) / (R_{531} + R_{570})$  (Gamon, Penuelas & Field 1992; Penuelas, Filella & Gamon 1995) where  $R_{531}$  and  $R_{570}$  are the reflectances at 531 nm and 570 nm, respectively.

*Statistical analyses:* Relationships between measured foliar parameters (physiology, pigments and leaf traits), and integrated daily PPFD and location (BRF or the FTE) were assessed using linear mixed effects models. The interaction between PPFD and location was included so as to examine the variations in slope between locations. Individual tree (FTE  $n = 18$ , BRF  $n = 6$ ) was included as a random effect to account for repeat measures on trees (Parameter  $\sim$  PPFD \* location + (1|tree)). If necessary, parameters were log-transformed to meet assumptions of normality of the residuals. Parameter estimates were considered significantly different from zero if  $P < 0.05$ . Linear mixed effects analysis was conducted using the lme4 package (Bates, Maechler, Bolker & Walker 2015) with the lmerTest package loaded to provide p-values (Kuznetsova, Brockhoff & Christensen 2017). All analyses took place in R v. 4.1.0 (R Core Team 2021).

The effects of canopy position and location on foliar traits were also assessed using mixed effects models implemented in the lme4 package (Bates *et al.* 2015). Again, individual tree was included as a random effect to account for repeat measures on trees. Pairwise comparisons between canopy position (high or low) and location (FTE or BRF) were assessed

using the emmeans package in R and found to be significant if  $P < 0.05$  (Lenth 2020). All data reported below are means  $\pm$  SE.

## RESULTS

*Tree Characteristics and Growth Environment:* Tree characteristics for the trees measured in this study have been previously reported in Griffin *et al.* (2022) but have been summarized in Table 1. Briefly, trees from BRF tended to have a larger diameter at breast height (DBH; measured at 1.37 m from the ground) with BRF trees having an average diameter of  $23.1 \pm 1.99$  cm and FTE trees having an average diameter of  $16.7 \pm 0.94$  cm (Table 1). Trees from the two locations were also extremely different in their average dripline areas ( $27.34 \pm 2.00$  m<sup>2</sup> at BRF vs.  $3.52 \pm 0.37$  m<sup>2</sup> at the FTE; Table 1). Nevertheless, despite these differences trees from the two locations had very similar average heights ( $9.91 \pm 0.73$  m at BRF and  $9.32 \pm 0.57$  m at the FTE; Table 1).

Irradiance at the range extremes of white spruce, from the northernmost location at the FTE in Alaska, to the southernmost location at BRF in New York, showed pronounced differences in both intensity and day length (Fig 1a, b, d). Average ambient PPFD over the measurement campaigns (Fig 1a; orange) estimated from daily diurnal light courses modeled from canopy photos (e.g., July 4<sup>th</sup> 2017; Fig 1b) was 36% higher at BRF than at the FTE ( $54.4 \pm 0.03$  compared to  $39.9 \pm 0.06$  mol m<sup>-2</sup> day<sup>-1</sup>). However, day length was much longer at the FTE than at BRF (Fig 1b). At the canopy positions where physiological and biochemical measurements were made, average daily PPFD was significantly greater at high canopy positions than low canopy positions at both BRF and the FTE ( $34.97 \pm 0.16$  vs.  $9.20 \pm 0.13$  mol m<sup>-2</sup> day<sup>-1</sup> at BRF, and  $29.24 \pm 0.16$  vs.  $21.14 \pm 0.16$  mol m<sup>-2</sup> day<sup>-1</sup> at the FTE; Fig 1a). Even so, the difference in the daily mean PPFD experienced from the upper to the lower canopies was much smaller at the FTE than at BRF (only 8 mol m<sup>-2</sup> day<sup>-1</sup> at the FTE vs. 25 mol m<sup>-2</sup> day<sup>-1</sup> at BRF). This convergence in the irradiance gradient across vertical canopy positions at the FTE was decoupled from canopy height, given that trees at both locations were of similar heights.

In addition to differences in the light environment, there were also pronounced differences in the daytime air temperatures between the two locations (Fig 1c). During the respective study campaigns (as marked by the green (BRF) and blue (FTE) shaded areas (Fig 1c & 1d), BRF mean daytime temperature was 23.4°C, with a minimum temperature of 12.8°C and a maximum temperature of 29.1°C, whereas the FTE mean temperature was 17°C, with a

minimum temperature of 6.5°C and a maximum temperature of 28°C. Thus, the FTE experienced colder temperatures and greater temperature variability.

*Relationships between light environment and gas exchange, biochemistry, pigments and PRI:*

In general, significant relationships were found between average PPFD and physiological traits, biochemical traits, pigment concentrations and PRI at BRF. In contrast, few significant relationships were found between average PPFD and foliar traits at the FTE (Table 2).

More specifically, a significant positive relationship was observed between average PPFD and  $A_{1500}$  at BRF with high canopy foliage having a higher  $A_{1500}$  than low canopy foliage. No significant relationship was observed between  $A_{1500}$  and average PPFD at the FTE, nor were there any significant differences between the canopy positions (Fig 2a, Table 2). Across all samples at each location, mean  $A_{1500}$  was found to be 40% higher at BRF than at the FTE ( $10.8 \pm 0.5$  at BRF vs.  $7.7 \pm 0.4 \mu\text{mol m}^{-2} \text{s}^{-1}$  at the FTE; Table 4). The differences between BRF and the FTE in the relationship of  $A_{1500}$  to PPFD correspond to differences in several other parameters commonly extracted from light response curves. Namely the following were observed: a significant negative relationship between average PPFD and  $\Phi$ , with significantly lower  $\Phi$  in the high than the low canopy (Fig 2b, Table 2); and significant positive relationships between average PPFD and LSP, LCP,  $R_D$ , and  $R_L$  with higher values in the high than the low canopies (Figs 2c & 2d, Table 2). At the FTE, as in the case of  $A_{1500}$ , no significant relationship was found between average PPFD and  $\Phi$ , LCP,  $R_D$  or  $R_L$  (Figs 2b & 2d, Table 2); however, a significant positive relationship was found between average PPFD and LSP with a correspondingly higher LSP in the high canopy than the low canopy foliage at both locations (Fig 2c, Table 2). Across all samples at each location,  $\Phi$  was significantly lower at the FTE than at BRF ( $0.03 \pm 0.001$  at the FTE vs.  $0.05 \pm 0.002$  at BRF; Tables 3 & 4), and LSP, LCP,  $R_D$  and  $R_L$  were significantly higher at the FTE than at BRF (LSP =  $644 \pm 14$  at the FTE vs.  $522 \pm 18.6 \mu\text{mol m}^{-2} \text{s}^{-1}$  at BRF; LCP =  $90.7 \pm 4.5$  at the FTE vs.  $36.8 \pm 2.1 \mu\text{mol m}^{-2} \text{s}^{-1}$  at BRF (LCP means are backtransformed);  $R_D$  =  $2.46 \pm 0.11$  at the FTE vs.  $1.98 \pm 0.14 \mu\text{mol m}^{-2} \text{s}^{-1}$  at BRF;  $R_L$  =  $2.22 \pm 0.12$  at the FTE vs.  $1.58 \pm 0.16 \mu\text{mol m}^{-2} \text{s}^{-1}$  at BRF; Tables 3 & 4). In particular, the differences in mean  $R_L$  between the two locations were dramatic, with the FTE having 41% higher  $R_L$  than at BRF.

The parameters extracted from the  $A-C_i$  curves ( $V_{\text{cmax}}$  and  $J_{\text{max}}$  normalized to 25°C) also followed the positive relationship between PPFD and  $A_{1500}$  at BRF. Both  $V_{\text{cmax}}$  and  $J_{\text{max}}$  had positive relationships with PPFD and significantly higher values in the high than the low canopy at BRF (Fig 3, Tables 2, 3 & 4). However, the expected lack of a relationship with PPFD at the FTE was not observed. Instead, FTE  $V_{\text{cmax}}$  and  $J_{\text{max}}$  both had significant negative relationships with PPFD, with significantly lower values in the high than the low canopy. Consequently, when values from canopy positions were pooled to compare the two locations, no significant differences were observed between the FTE and BRF ( $V_{\text{cmax}} = 48.5 \pm 1.7$  at the FTE and  $50.3 \pm 2.2 \mu\text{mol m}^{-2} \text{s}^{-1}$  at BRF;  $J_{\text{max}} = 80.8 \pm 2.7$  at the FTE and  $85.9 \pm 3.4 \mu\text{mol m}^{-2} \text{s}^{-1}$  at BRF; Tables 3 & 4).

In addition to gas exchange parameters, leaf traits, pigment concentrations and the photochemical reflectance index (PRI) were examined. A significant negative relationship was observed between SLA and PPFD at BRF while no relationship was observed between SLA and PPFD at the FTE (Fig 4a, Table 2). Across canopy positions, high canopy foliage had significantly lower SLA at both locations. The FTE had significantly lower SLA than BRF ( $32.0 \pm 1.1$  at the FTE vs.  $50.4 \pm 1.2 \text{ cm}^2 \text{g}^{-1}$  at BRF, Tables 3 & 4). %N showed no significant relationships with PPFD, and no significant differences between canopy positions in either location; although the FTE did have a small but significantly lower %N than BRF ( $0.98 \pm 0.03$  at the FTE vs.  $1.1 \pm 0.04$  at BRF; Fig 4c, Tables 2, 3 & 4). Similarly, C:N showed no significant relationships with PPFD and no significant differences between canopy positions in either location; however, C:N was significantly lower at BRF compared to the FTE (Tables 2, 3 & 4). Due to the strong relationship between SLA and PPFD,  $N_{\text{area}}$  also had a significant positive relationship with PPFD at BRF and no relationship at the FTE (Fig 4e, Table 2). With regard to pigment concentrations, positive relationships were seen both at BRF and the FTE in the ratio of chlorophyll *a* and chlorophyll *b* ( $\text{Chl}a:\text{Chl}b_{\text{area}}$ ) and in the Carotenoids ( $\text{Car}_{\text{area}}$ ) with higher  $\text{Chl}a:\text{Chl}b_{\text{area}}$  and higher  $\text{Car}_{\text{area}}$  in the high than the low canopies (Figs 4b & 4d, Tables 2, 3 & 4). The expected decrease in the ratio of chlorophyll to carotenoids ( $\text{Chl}:\text{Car}_{\text{area}}$ ) with increasing PPFD was observed at BRF, but no relationship was found at the FTE (Fig 2f, Table 2). However, the FTE did have 46% more carotenoids than BRF ( $14.5 \pm 0.5$  at the FTE vs.  $9.9 \pm 0.8 \mu\text{g cm}^{-2}$  at BRF; Table 4) and a significantly lower  $\text{Chl}:\text{Car}_{\text{area}}$  ( $6.7 \pm 0.2$  at the FTE vs.  $9.0 \pm 0.3$  at BRF; Tables 3 & 4). Finally, PRI had the expected positive relationship with  $\text{Chl}:\text{Car}_{\text{area}}$  at BRF

(Fig 5) and it decreased significantly with increasing PPFD at BRF (Fig 5, Table 2). However, no relationships between PRI and  $\text{Chl}:\text{Car}_{\text{area}}$  or PPFD were found at the FTE (Fig 5, Table 2).

Finally, we also examined relationships between gas exchange, pigments and nitrogen. As with PPFD, positive relationships were found between  $N_{\text{area}}$  and  $A_{1500}$ , and  $N_{\text{area}}$  and  $V_{\text{cmax}}$  at BRF while no relationships were observed at the FTE (Fig 6a & 6b, Table 5). In contrast, both BRF and the FTE had positive relationships between  $N_{\text{area}}$  and  $R_{\text{D}}$  (Fig 6c, Table 5), between  $N_{\text{area}}$  and  $R_{\text{L}}$  (Table 5) and between  $N_{\text{area}}$  and total pigments ( $\text{chl}a + \text{chl}b + \text{carotenoids}$ ; Fig 6d, Table 5).

*Carbon balance at the range extremes of white spruce:* The collection of gas exchange parameters such as  $V_{\text{cmax}}$  and  $J_{\text{max}}$ , which are commonly used to predict photosynthesis, together with values of respiration at 25°C ( $R_{25}$ ) previously reported by Griffin *et al.* (2022), provide a unique opportunity to examine the carbon balance across the range extremes of white spruce. We find that, when all parameters are normalized to 25°C, the ratios of  $V_{\text{cmax}}$  and  $J_{\text{max}}$  to  $R_{25}$  are 228% higher at BRF than at the FTE ( $V_{\text{cmax}}/R_{25} = 21.4 \pm 1.39$  at the FTE vs.  $70.2 \pm 7.47$  at BRF;  $J_{\text{max}}/R_{25} = 36.5 \pm 2.45$  at the FTE vs.  $119.9 \pm 13.21$  at BRF). No significant differences were observed in the ratios between canopy positions at each location, although there was a trend for larger ratios in the low canopy than the high canopy at BRF (Fig 7).

## DISCUSSION

We find remarkable differences between the vertical canopy gradients of foliar physiological and biochemical traits of white spruce growing at its northern- and southern-most range limits. Whereas steep canopy gradients in physiological functioning are found at the southern range limit in Black Rock Forest (BRF), a clear convergence in foliar physiological capacity is observed throughout the canopies of white spruce at its northernmost range extreme in the Forest-Tundra Ecotone (FTE). As our two study locations delimit the distributional range of white spruce and are separated by a distance of more than 5000 km, these differences are likely in response to local environmental conditions. Solar zenith angles, day length, air and soil temperatures, soil moisture and vapor pressure deficit vary substantially between these two locations. Of these environmental conditions, the impacts of local light environment are of particular interest for the following reasons: 1) light energy is a critical driver of photosynthetic



carbon gain (Hirose & Werger 1987); 2) abundant evidence shows that canopy gradients in photosynthetic traits are strongly correlated to gradients in irradiance throughout tree crowns (Field 1983; Hirose & Werger 1987; Ellsworth & Reich 1993; Bond *et al.* 1999; Niinemets, Keenan & Hallik 2015), but little work assesses the impacts of latitudinal location on these gradients within a single species; and 3) these gradients are key parameters in ecosystem models used to calculate gross primary productivity across latitudinal space (Bonan *et al.* 2012; Rogers *et al.* 2017). Day length and solar zenith angles increase and light intensity decreases as one moves towards the northernmost latitudes of the globe (Nilsen 1983; Slaughter & Viereck 1986). These local light environments are further modified by differences in forest and tree crown structure between the two locations. Trees at the southern location exist within a complex forest canopy comprised of deciduous hardwood species where they must compete for light energy and other above-ground resources. Consequently white spruce tree crowns are distinctly conical in shape with a wide dripline area in the southern location, presumably to minimize self-shading and maximize light energy capture. In contrast, trees at the northern location are widely spaced with narrow and untapering tree crowns. The open forest canopy minimizes competition for resources such as light. Furthermore, the cylindrical shape promotes more equal exposure of foliage to irradiance in a region of the globe with extremely high solar zenith angles, ultimately increasing the efficiency of light capture for photosynthesis (Kuuluvainen 1992). Overall, these solar and structural differences lead to a narrowing of the intra-canopy light gradients as one moves from the southern to the northern latitudinal limits of this species.

Inherent to the structural differences of trees in these two locations is the need for tree crowns to organize their physiological and biochemical traits in order to optimize photosynthetic carbon gain (Hirose & Werger 1987; Niinemets *et al.* 1998). Therefore, at BRF, where there are strong intra-canopy light gradients, we find strongly positive relationships between PPFD and  $A_{1500}$ ,  $J_{\max}$  and  $V_{\max}$  indicating the upregulation of the light reactions (in which RuBP is produced) and the carbon reactions (in which rubisco acts on RuBP to fix carbon) of photosynthesis in the high canopy (Givnish 1988; Valladares & Niinemets 2008). The elevated metabolic activity in the high canopy results in higher respiratory activity in order to support the costs of growth and protein maintenance (Reich *et al.* 1998; Lewis, McKane, Tingey & Beedlow P. A. 2000; Griffin, Tissue, Turnbull, Schuster & Whitehead 2001). Biochemical traits also follow the inherent light gradients. High light needles have lower SLA and a greater  $N_{\text{area}}$



reflecting the greater leaf thickness of high light leaves from added palisade layers and the presence of nitrogen in both the enzyme rubisco and in chlorophyll (Evans 1989; Smith, Vogelmann, DeLucia, Bell & Shepherd 1997; Evans & Poorter 2001). Finally, high light leaves have greater  $Car_{area}$  and a smaller Chl:Car ratio reflecting the need for greater photoprotection from excess light energy (Demmig-Adams & Adams III 1992). In all cases, biochemical and physiological traits of low latitude white spruce conform to the established literature showing strong correlations with canopy gradients of irradiance. At the high latitude location where intra-canopy light gradients become less pronounced, we see the absence of canopy gradients in many physiological and biochemical traits. The most dramatic of these is the lack of relationships between irradiance and each of photosynthesis ( $A_{1500}$ ), light respiration, and dark respiration. Together with the lack of relationships between irradiance and the apparent quantum yield, the light compensation point, specific leaf area and  $N_{area}$ , these findings suggest that the uneven distribution of metabolic activity throughout the canopy required to optimize photosynthetic carbon gain in correspondence with canopy irradiance gradients at low latitudes does not occur at high latitudes where irradiance varies minimally throughout the canopy profile.

A more nuanced interpretation of the various potential environmental controls on canopy photosynthetic physiology at the northern latitude location is possible when several additional traits are examined. Positive relationships between irradiance and the ratio of Chl $a$  to Chl $b$ , the light saturation point and carotenoid concentrations suggest that even though the magnitude of the light gradient narrows at northern latitudes, there is still a subtle intra-canopy light gradient to which pigments are responding. Thus, in the higher light upper canopy we see greater investments in Chl $a$  than Chl $b$  to maximize light capture and a greater investment in carotenoids to dissipate excess light energy and protect photosynthetic machinery (Dale & Causton 1992; Hikosaka & Terashima 1995; Kitajima & Hogan 2003; Ruban 2015; Magney *et al.* 2016; Scartazza *et al.* 2016). Despite these subtle gradients in response to the northern light environment, we see no upregulation of photosynthesis in the high canopy of white spruce in the northern latitudes, and, surprisingly, we find a downregulation of  $J_{max}$  and  $V_{cmax}$  in the high canopy foliage. It is unclear why we find these negative relationships between irradiance and the component processes of photosynthesis at the northern locations; however, there are several possible explanations and hypotheses that could be further explored. One possibility is that the combination of diminutive vertical canopy gradients in light and nitrogen, the latter of which is

critical to the performance of  $J_{\max}$  and  $V_{\max}$  because of the presence of nitrogen in chlorophyll and the enzyme rubisco (Evans 1989; Smith, Vogelmann, DeLucia, Bell & Shepherd 1997; Evans & Poorter 2001), together with any variation across Alaskan needles and trees, has led to the observed negative relationships. However, strong relationships do exist between nitrogen and total pigment concentrations even though there is no apparent relationship between nitrogen and  $V_{\max}$ . Furthermore, we still find weak positive relationships between irradiance and both carotenoid concentrations and  $Chla:Chlb$  even with a narrow light gradient. Consequently, we conclude that a more mechanistic explanation must exist for the puzzling downregulation of  $V_{\max}$  and  $J_{\max}$ .

In this study, we focused on the latitudinal relationships between canopy light environments and foliar physiological traits, but intracanopy gradients in other environmental conditions may also impact  $V_{\max}$  and  $J_{\max}$  measurements (Martin, Hinckley, Meinzer & Sprugel 1999; Zweifel, Böhm & Häslér 2002; Bauerle, Bowden, Wang & Shahba 2009; Bachofen, D’Odorico & Buchmann 2020; Buckley 2021). Perhaps of particular relevance with respect to rubisco carboxylation is the reliance of this process on the enzyme rubisco, whose activity is well-known to be temperature dependent (Yamori & von Caemmerer 2009; Benomar *et al.* 2018). Work at alpine treelines has clearly documented the fact that as trees grow in stature, their foliage becomes increasingly coupled to atmospheric conditions such as temperature and wind stress (Körner 2012). Thus, low canopy foliage may derive protective benefits from temperature and wind extremes found to be detrimental to high canopy photosynthetic performance (Martin *et al.* 1999; Germino & Smith 1999; Johnson, Germino & Smith 2004; Maguire *et al.* 2019). As a result, it is possible that throughout the canopy of Alaskan trees, there may be a constant balancing act with regard to the resource allocation necessary to improve whole canopy photosynthetic carbon gain – namely between the need to devote resources to light capture in the high canopy, where irradiance is slightly more available, in order to take advantage of light energy when conditions are favorable, and the need to devote resources either to foliage protection in the high canopy or increased carbon gain in the low canopy when harsh conditions prevail. The result, ultimately, is what we find across the majority of physiological and biochemical traits, i.e., a convergence in photosynthetic carbon gain across the high and low canopy in high latitude trees. Without detailed information on temperature and wind profiles throughout the canopy, this hypothesis cannot be fully tested. Yet, it seems clear from the

linkages between irradiance, photosynthesis and tree structure across the latitudinal extremes that the environment is an integral driver of species' geographic range. We suggest that expanding to examine additional environmental conditions such as temperature and wind stress may be a compelling future direction, thereby adding to our overall understanding of the mechanisms controlling canopy carbon exchange across the massive forest tundra ecotone and more broadly across latitudinal gradients.

Accurately portraying canopy gradients is important to the precise prediction of carbon exchange and gross primary productivity (GPP) (Sands 1995; Bonan *et al.* 2012). However, the convergence in photosynthesis at the FTE suggests that complex canopy models used to account for canopy self-shading at mid-latitudes may not be necessary to model GPP in trees from high latitudes. The convergence in the photochemical reflectance index (PRI) across vertical canopy positions at the FTE similarly suggests that accounting for canopy self-shading may be less critical at northern latitudes as opposed to mid latitudes. Furthermore, the fact that PRI mirrors  $A_{1500}$  regardless of latitude suggests that this remotely sensed index may be a valuable future tool for accurately assessing gross primary productivity of other species with different structural properties and with large latitudinal distributions.

The detailed data on the physiology and biochemistry of white spruce not only allow for the examination of canopy differences across latitudinal range extremes, but also for the examination of locational differences in traits that may further illuminate possible drivers contributing to the range limits of this important species. Here, we observed important differences in physical and biochemical traits such as SLA, % N, C:N and carotenoid concentration between the southern and northern range extremes. Most relevant to the above discussion of light environment and its impacts on photosynthetic physiology is the finding of significantly higher carotenoid concentrations in needles of northern white spruce compared to southern white spruce. This indicates a greater investment in photoprotection at northern latitudes during the growing season. Elevated carotenoids at the northern range extreme may be in response to the 24 hour photoperiod at the Arctic tundra boreal forest ecotone. Past studies have found that very high levels of light protecting xanthophyll pigments can be a feature of Arctic plants subjected to continuous light (Magney *et al.* 2019). Lower overall SLA in Alaskan needles compared to BRF needles indicates that foliage at the northern edge of the distribution is thicker or denser. Although decreasing SLA is commonly found in response to increasing light

availability due to the added density of additional photosynthetic machinery (as seen throughout the BRF canopy), SLA is also commonly positively correlated to ambient growth temperatures (Atkin, Loveys, Atkinson & Pons 2006; Poorter, Niinemets, Poorter, Wright & Villar 2009). Thus, a low SLA at the FTE may indicate a greater investment in needle structure rather than photosynthetic machinery, a hypothesis that is corroborated by our finding of higher carbon to nitrogen ratios in Alaskan needles (González-Zurdo, Escudero, Babiano, García-Ciudad & Mediavilla 2016). Investment in structure is potentially adaptive in the harsh forest tundra ecotone where needles must survive multiple years of winter temperatures, snow or ice accumulation and windblown ice abrasion. Leaf nitrogen was also found to be slightly but significantly lower in needles from the northern range limit. This result confirms the findings of Griffin *et al.* (2022) and agrees with past studies that show negative correlations between nitrogen and both temperature and latitude (Körner 1989; Yin 1993; Reich & Oleksyn 2004). Nitrogen biogeochemistry is intimately linked to environmental and ecological factors. For example, harsh temperatures and unique hydrologic and permafrost dynamics inherent to the Arctic can slow decomposition rates and depress nitrogen fixation (Schimel & Stuart Chapin 1996; Schimel, Bilbrough & Welker 2004). The result is the limited availability of nitrogen in the harsh northern latitudes of the globe. The differences in biochemical and structural traits between our two study locations hint at environmental drivers, in addition to light availability, that should be considered in future studies examining latitudinal differences in canopy gradients.

Finally, in addition to biochemical differences between locations, metabolic differences are key in understanding the mechanisms controlling the range limits of this species. White spruce has been found to have extremely high rates of dark respiration at its northern compared to southern range limit, a finding that has led to the conclusion that respiratory carbon loss may be a crucial factor constraining the northern range limit of this species Griffin *et al.* (2022). We find similar patterns in the current study, with significantly higher rates of dark respiration (as estimated from the light response curves) in white spruce in Alaska than in New York. Furthermore, we find that these differences are apparent not only in dark respiration, but also in light respiration. Given that the northern range extreme of this species experiences 24 hours of sunlight during the majority of the growing season, the greater rates of light respiration in FTE compared to BRF trees provides important corroboration of the constancy of the extreme respiratory cost at this northern range limit and the role it may play in determining the location of

northern treeline. It is likely that these high respiratory fluxes to the atmosphere are related to high maintenance costs given the harsh environmental conditions of the region.

Northern white spruce are not only characterized by high carbon losses. The additional information on photosynthetic processes provided here further suggests that these carbon costs are not matched by similarly high carbon gains. We do not directly compare the ratios of photosynthetic carbon gain to respiratory carbon loss due to the use of different measurement temperatures chosen to reflect ambient conditions in each location, but the ability to normalize both  $V_{\text{cmax}}$  and  $J_{\text{max}}$  to 25°C allows us to assess the carbon balance of white spruce at its latitudinal range extremes. The results of this comparison are dramatic and demonstrate that northern trees function within a much narrower margin to maintain a positive carbon balance. It seems that it is not only high respiratory costs that result in slow growth rates of white spruce (Jensen et al., *pers. com.*) and constrain the latitudinal range limit of northern treeline. Instead, our study suggests that it is the combined effect of high respiration and low  $V_{\text{cmax}}$  and  $J_{\text{max}}$  that ultimately limit the northern range of this widely distributed and important boreal species.

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#### AUTHOR CONTRIBUTION

SCS, KLG, NTB, LAV and JUHE designed the research. They were assisted in data collection by SGB, LM, and JJ. SCS analyzed the data and wrote the first draft of the manuscript. All authors edited and approved the final version for submission.

#### DATA AVAILABILITY

Data will be archived with ORNL DAAC upon acceptance of the manuscript.

## REFERENCES

- Arguez A.I., Durre I., Applequist S., Squires M.F., Vose R.S., Yin X., ... Owen T. (2012) NOAA's 1981-2010 U.S. Climate Normals: An Overview. *Bulletin of the American Meteorological Society* **93**, 1687–1697.
- Atkin O.K., Loveys B.R., Atkinson L.J. & Pons T.L. (2006) Phenotypic plasticity and growth temperature: understanding interspecific variability. *Journal of Experimental Botany* **57**, 267–281.
- Ayub G., Smith R.A., Tissue D.T. & Atkin O.K. (2011) Impacts of drought on leaf respiration in darkness and light in *Eucalyptus saligna* exposed to industrial-age atmospheric CO<sub>2</sub> and growth temperature. *New Phytologist* **190**, 1003–1018.
- Bachofen C., D'Odorico P. & Buchmann N. (2020) Light and VPD gradients drive foliar nitrogen partitioning and photosynthesis in the canopy of European beech and silver fir. *Oecologia* **192**, 323–339.
- Bates D., Maechler M., Bolker B. & Walker S. (2015) Fitting Linear Mixed-Effects Models using lme4. *Journal of Statistical Software* **67**, 1–48.
- Bauerle W.L., Bowden J.D., Wang G.G. & Shahba M.A. (2009) Exploring the importance of within-canopy spatial temperature variation on transpiration predictions. *Journal of Experimental Botany* **60**, 3665–3676.
- Benomar L., Lamhamedi M.S., Pepin S., Rainville A., Lambert M.-C., Margolis H.A., ... Beaulieu J. (2018) Thermal acclimation of photosynthesis and respiration of southern and northern white spruce seed sources tested along a regional climatic gradient indicates limited potential to cope with temperature warming. *Annals of Botany* **121**, 443–457.
- Bolker B. & R Development Core Team (2020) bbmle: Tools for General Maximum Likelihood Estimation. R package version 1.0.23.1. <https://CRAN.R-project.org/package=bbmle>.
- Bonan G.B., Oleson K.W., Fisher R.A., Lasslop G. & Reichstein M. (2012) Reconciling leaf physiological traits and canopy flux data: Use of the TRY and FLUXNET databases in the Community Land Model version 4. *Journal of Geophysical Research: Biogeosciences* **117**, 1–19.
- Bond B.J., Farnsworth B.T., Coulombe R.A. & Winner W.E. (1999) Foliage physiology and biochemistry in response to light gradients in conifers with varying shade tolerance. *Oecologia* **120**, 183–192.

- Brown J.H., Stevens G.C. & Kaufmann D.M. (1996) The Geographic Range: Size, Shape, Boundaries, and Internal Structure. *Annual Review of Ecology and Systematics* **27**, 597–623.
- Buckley T.N. (2021) Optimal carbon partitioning helps reconcile the apparent divergence between optimal and observed canopy profiles of photosynthetic capacity. *New Phytologist* **230**, 2246–2260.
- Buckley T.N., Vice H. & Adams M.A. (2017) The Kok effect in *Vicia faba* cannot be explained solely by changes in chloroplastic CO<sub>2</sub> concentration. *New Phytologist* **216**, 1064–1071.
- Dale M.P. & Causton D.R. (1992) Use of the Chlorophyll a/b Ratio as a Bioassay for the Light Environment of a Plant. *Functional Ecology* **6**, 190–196.
- Demmig-Adams B. & Adams III W.W. (1992) Photoprotection and other responses of plants to high light stress. *Annu. Rev. Plant Physiol. Plant Mol. Biol.* **43**, 599–626.
- Duursma R.A. (2015) Plantecophys--An R Package for Analysing and Modelling Leaf Gas Exchange Data. *PloS One* **10**, e0143346.
- Eitel J.U.H., Griffin K.L., Boelman N.T., Maguire A.J., Meddens A.J.H., Jensen J., ... Jennewein J.S. (2020) Foliar remote sensing tracks radial tree growth dynamics. *Global Change Biology* **26**, 4068–4078.
- Eitel J.U.H., Maguire A.J., Boelman N., Vierling L.A., Griffin K.L., Jensen J., ... Sonnentag O. (2019) Proximal remote sensing of tree physiology at northern treeline: Do late-season changes in the photochemical reflectance index (PRI) respond to climate or photoperiod? *Remote Sensing of Environment* **221**, 340–350.
- Ellsworth D.S. & Reich P.B. (1993) Canopy Structure and Vertical Patterns of Photosynthesis and Related Leaf Traits in a Deciduous Forest. *Oecologia* **96**, 169–178.
- Evans J.R. (1989) Photosynthesis and nitrogen relationships in leaves of C<sub>3</sub> plants. *Oecologia* **78**, 9–19.
- Evans J.R. & Poorter H. (2001) Photosynthetic acclimation of plants to growth irradiance: the relative importance of specific leaf area and nitrogen partitioning in maximizing carbon gain. *Plant, Cell and Environment* **24**, 755–767.
- Farquhar G.D. & Busch F.A. (2017) Changes in the chloroplastic CO<sub>2</sub> concentration explain much of the observed Kok effect: a model. *New Phytologist* **214**, 570–584.
- Farquhar G.D., von Caemmerer S. & Berry J.A. (1980) A biochemical model of photosynthetic



- CO<sub>2</sub> assimilation in leaves of C<sub>3</sub> species. *Planta* **149**, 78–90.
- Field C. (1983) Allocating Leaf Nitrogen for the Maximization of Carbon Gain: Leaf Age as a Control on the Allocation Program. *Oecologia* **56**, 341–347.
- Gamon A., Penuelas J. & Field C.B. (1992) A Narrow-Waveband Spectral Index That Tracks Diurnal Changes in Photosynthetic Efficiency. *Remote Sensing of Environment* **41**, 35–44.
- Gamon J.A., Huemmrich K.F., Wong C.Y.S., Ensminger I., Garrity S., Hollinger D.Y., ... Penuelas J. (2016) A remotely sensed pigment index reveals photosynthetic phenology in evergreen conifers. *PNAS* **113**, 13087–13092.
- Gaston K.J. (1996) Species-range-size distributions: Patterns, mechanisms and implications. *Trends in Ecology and Evolution* **11**, 197–201.
- Germino M.J. & Smith W.K. (1999) Sky exposure, crown architecture, and low-temperature photoinhibition in conifer seedlings at alpine treeline. *Plant, Cell and Environment* **22**, 407–415.
- Givnish T.J. (1988) Adaptation to Sun and Shade: A Whole-plant Perspective. *Australian Journal of Plant Physiology* **15**, 63–92.
- González-Zurdo P., Escudero A., Babiano J., García-Ciudad A. & Mediavilla S. (2016) Costs of leaf reinforcement in response to winter cold in evergreen species. *Tree Physiology* **36**, 273–86.
- Griffin K.L., Griffin Z.M., Schmiede S.C., Bruner S., Boelman N.T., Vierling L.A. & Eitel J.U.H. (2022) Variation in white spruce needle respiration at the species range limits: a potential impediment to northern expansion. *Plant, Cell & Environment*, 1–15.
- Griffin K.L., Schmiede S.C., Bruner S.G., Boelman N.T., Vierling L.A. & Eitel J.U.H. (2021) High Leaf Respiration Rates May Limit the Success of White Spruce Saplings Growing in the Kampfzone at the Arctic Treeline. *Frontiers in Plant Science* **12**.
- Griffin K.L., Tissue D.T., Turnbull M.H., Schuster W. & Whitehead D. (2001) Leaf dark respiration as a function of canopy position in *Nothofagus fusca* trees grown at ambient and elevated CO<sub>2</sub> partial pressures for 5 years. *Functional Ecology* **15**, 497–505.
- Halekoh U. & Højsgaard S. (2014) A kenward-Roger approximation and parametric bootstrap methods for tests in linear mixed models-the R package pbkrtest. *Journal of Statistical Software* **59**, 1–32.
- Harley P.C. & Baldocchi D.D. (1995) Scaling carbon dioxide and water vapour exchange from

- leaf to canopy in a deciduous forest. I. Leaf model parametrization. *Plant, Cell and Environment* **18**, 1146–1156.
- Heskel M.A., Atkin O.K., Turnbull M.H. & Griffin K.L. (2013) Bringing the Kok effect to light: A review on the integration of daytime respiration and net ecosystem exchange. *Ecosphere* **4**, 1–14.
- Hikosaka K. & Terashima I. (1995) A model of the acclimation of photosynthesis in the leaves of C<sub>3</sub> plants to sun and shade with respect to nitrogen use. *Plant, Cell and Environment* **18**, 605–618.
- Hirose T. & Werger M.J.A. (1987) Maximizing daily canopy photosynthesis with respect to the leaf nitrogen allocation pattern in the canopy. *Oecologia* **72**, 520–526.
- Johnson D.M., Germino M.J. & Smith W.K. (2004) Abiotic factors limiting photosynthesis in *Abies lasiocarpa* and *Picea engelmannii* seedlings below and above the alpine timberline. *Tree Physiology* **24**, 377–386.
- Kirschbaum M.U. & Farquhar G.D. (1987) Investigation of the CO<sub>2</sub> Dependence of Quantum Yield and Respiration in *Eucalyptus pauciflora*. *Plant Physiology* **83**, 1032–1036.
- Kitajima K. & Hogan K.P. (2003) Increases of chlorophyll a/b ratios during acclimation of tropical woody seedlings to nitrogen limitation and high light. *Plant, Cell and Environment* **26**, 857–865.
- Kok B. (1948) A critical consideration of the quantum yield of *Chlorella*-photosynthesis. *Enzymologia* **13**, 1–56.
- Körner C. (1989) The nutritional status of plants from high altitudes - A worldwide comparison. *Oecologia* **81**, 379–391.
- Körner C. (2012) *Alpine treelines: Functional Ecology of the Global High Elevation Tree Limits*. Springer, Basel.
- Kuuluvainen T. (1992) Tree Architectures Adapted to Efficient Light Utilization: Is There a Basis for Latitudinal Gradients? *Nordic Society Oikos* **65**, 275–284.
- Kuznetsova A., Brockhoff P. & Christensen R. (2017) lmerTest Package: Tests in Linear Mixed Effects Models. *Journal of Statistical Software* **82**, 1–26.
- Lambers H. & Oliveira R.S. (2019) Physiological Ecology and the Distribution of Organisms. In *Plant Physiological Ecology*, 3rd Editio. pp. 2–5. Springer.
- Lenth R. (2020) emmeans: Estimated Marginal Means, aka Least-Squares Means. R package

- version 1.4.5. <https://CRAN.R-project.org/package=emmeans>.
- Lewis J.D., McKane R.B., Tingey D.T. & Beedlow P. A. (2000) Vertical gradients in photosynthetic light response within an old-growth Douglas-fir and western hemlock canopy. *Tree Physiology* **20**, 447–456.
- Lichtenthaler H.K. (1987) Chlorophylls and carotenoids: Pigments of photosynthetic biomembranes. *Methods in Enzymology* **148**, 350–382.
- Lowry E. & Lester S.E. (2006) The biogeography of plant reproduction: Potential determinants of species' range sizes. *Journal of Biogeography* **33**, 1975–1982.
- Magney T.S., Bowling D.R., Logan B.A., Grossmann K., Stutz J., Blanken P.D., ... Frankenberg C. (2019) Mechanistic evidence for tracking the seasonality of photosynthesis with solar-induced fluorescence. *Proceedings of the National Academy of Sciences of the United States of America* **116**, 11640–11645.
- Magney T.S., Eitel J.U.H., Griffin K.L., Boelman N.T., Greaves H.E., Prager C.M., ... Vierling L.A. (2016) LiDAR canopy radiation model reveals patterns of photosynthetic partitioning in an Arctic shrub. *Agricultural and Forest Meteorology* **221**, 78–93.
- Maguire A.J., Eitel J.U.H., Vierling L.A., Johnson D.M., Griffin K.L., Boelman N.T., ... Meddens A.J.H. (2019) Terrestrial lidar scanning reveals fine-scale linkages between microstructure and photosynthetic functioning of small-stature spruce trees at the forest-tundra ecotone. *Agricultural and Forest Meteorology* **269–270**, 157–168.
- Martin T.A., Hinckley T.M., Meinzer F.C. & Sprugel D.G. (1999) Boundary layer conductance, leaf temperature and transpiration of *Abies amabilis* branches. *Tree Physiology* **19**, 435–443.
- Mavi H.S. & Tupper G.J. (2004) Solar radiation and its role in plant growth. In *Agrometeorology: Principles and applications of climate studies in agriculture*. pp. 13–42. Food Products Press, Binghamton, NY.
- Niinemets Ü. (2007) Photosynthesis and resource distribution through plant canopies. *Plant, Cell & Environment* **30**, 1052–1071.
- Niinemets Ü., Keenan T.F. & Hallik L. (2015) A worldwide analysis of within-canopy variations in leaf structural, chemical and physiological traits across plant functional types. *New Phytologist* **205**, 973–993.
- Niinemets U., Kull O. & Tenhunen J.D. (1998) An analysis of light effects on foliar morphology,

- physiology and light interception in temperate deciduous woody species of contrasting shade tolerance. *Tree Physiology* **18**, 681–696.
- Nilsen J. (1983) Light Climate in Northern Areas. In *Plant Production in the North*. (eds A. Kaurin, O. Junttila & J. Nilsen), pp. 62–72. Norwegian University Press, Tromsø, Norway.
- Patterson A.E., Arkebauer R., Quallo C., Heskell M.A., Li X., Boelman N. & Griffin K.L. (2018) Temperature response of respiration and respiratory quotients of 16 co-occurring temperate tree species. *Tree Physiology* **00**, 1–14.
- Penuelas J., Filella I. & Gamon J.A. (1995) Assessment of photosynthetic radiation-use efficiency with spectral reflectance. *New Phytologist*, 291–296.
- Poorter H., Niinemets Ü., Poorter L., Wright I.J. & Villar R. (2009) Causes and consequences of variation in leaf mass per area (LMA): a meta-analysis. *New Phytologist* **182**, 565–588.
- de Pury D.G.G. & Farquhar G.D. (1997) Simple scaling of photosynthesis from leaves to canopies without the errors of big-leaf models. *Plant, Cell and Environment* **20**, 537–557.
- R Core Team (2021) R: A language and environment for statistical computing.
- Reich P.B. & Oleksyn J. (2004) Global patterns of plant leaf N and P in relation to temperature and latitude. *Proceedings of the National Academy of Sciences of the United States of America* **101**, 11001–11006.
- Reich P.B., Walters M.B., Ellsworth D.S., Vose J.M., Volin J.C., Gresham C. & Bowman W.D. (1998) Relationships of leaf dark respiration to leaf nitrogen, specific leaf area and leaf life-span: a test across biomes and functional groups. *Oecologia* **114**, 471–482.
- Rogers A., Medlyn B.E., Dukes J.S., Bonan G., von Caemmerer S., Dietze M.C., ... Zaehle S. (2017) A roadmap for improving the representation of photosynthesis in Earth system models. *New Phytologist* **213**, 22–42.
- Ruban A. V (2015) Evolution under the sun: optimizing light harvesting in photosynthesis. *Journal of Experimental Botany* **66**, 7–23.
- Sands P. (1995) Modelling Canopy Production. I. Optimal Distribution of Photosynthetic Resources. *Australian Journal of Plant Physiology* **22**, 593–601.
- Scartazza A., Di Baccio D., Bertolotto P., Gavrichkova O. & Matteucci G. (2016) Investigating the European beech (*Fagus sylvatica* L.) leaf characteristics along the vertical canopy profile: leaf structure, photosynthetic capacity, light energy dissipation and photoprotection mechanisms. *Tree Physiology* **36**, 1060–1076.

- Schimel J.P., Bilbrough C. & Welker J.M. (2004) Increased snow depth affects microbial activity and nitrogen mineralization in two Arctic tundra communities. *Soil Biology and Biochemistry* **36**, 217–227.
- Schimel J.P. & Stuart Chapin F. (1996) Tundra plant uptake of amino acid and NH<sub>4</sub><sup>+</sup> nitrogen in situ: Plants compete well for amino acid N. *Ecology* **77**, 2142–2147.
- Schnieder C.A., Rasband W.S. & Eliceiri K.W. (2012) NIH Image to ImageJ: 25 years of image analysis. *Nature Methods* **9**, 671–675.
- Schuster W.S.F., Griffin K.L., Roth H., Turnbull M.H., Whitehead D. & Tissue D. T (2008) Changes in composition, structure and aboveground biomass over seventy-six years (1930 - 2006) in Black Rock Forest, Hudson Highlands, southeastern New York State. *Tree Physiology* **28**, 537–549.
- Sellers P.J., Berry J.A., Collatz G.J., Field C.B. & Hall F.G. (1992) Canopy reflectance, photosynthesis, and transpiration. III. A reanalysis using improved leaf models and a new canopy integration scheme. *Remote Sensing of Environment* **42**, 187–216.
- Sharp R.E., Matthews M.A. & Boyer J.S. (1984) Kok effect and the quantum yield of photosynthesis: light partially inhibits dark respiration. *Plant Physiology* **75**, 95–101.
- Slaughter C.W. & Viereck L.A. (1986) Climatic Characteristics of the Taiga in Interior Alaska. In *Forest Ecosystems in the Alaskan Taiga: A synthesis of structure and function*. (eds K. Van Cleve, F.S. Chapin III, P.W. Flanagan, L.A. Viereck & C.T. Dyrness), pp. 9–21. Springer, New York.
- Smith W.K., Vogelmann T.C., DeLucia E.H., Bell D.T. & Shepherd K.A. (1997) Leaf Form and Photosynthesis: Do leaf structure and orientation interact to regulate internal light and carbon dioxide? *BioScience* **47**, 785–793.
- ter Steege H. (1993) *HEMIPHOT, a programme to analyze vegetation indices, light and light quality from hemispherical photographs*. Tropenbos Foundation, Wageningen.
- ter Steege H. (1997) *winphot, Version 5.0. A programme to analyse vegetation indices, light and light quality from hemispherical photographs*. Tropenbos-Guyana Programme.
- ter Steege H. (2018) Hemiphot.R: Free R scripts to analyse hemispherical photographs for canopy openness, leaf area index and photosynthetic active radiation under forest canopies.
- Survey U.S.G. (1999) Digital representation of “Atlas of United States Trees” by Elbert L. Little Jr. *U.S. Geological Survey Professional Paper 1650*.

- Tcherkez G., Gauthier P., Buckley T.N., Busch F.A., Barbour M.M., Bruhn D., ... Cornic G. (2017) Tracking the origins of the Kok effect, 70 years after its discovery. *New Phytologist* **214**, 506–510.
- Valladares F. & Niinemets U. (2008) Shade Tolerance, a Key Plant Feature of Complex Nature and Consequences. *Annual Review of Ecology, Evolution, and Systematics* **39**, 237–57.
- Viereck L.A., Van Cleve K. & Dyrness C.T. (1986) Forest Ecosystem Distribution in the Taiga Environment. In *Forest Ecosystems in the Alaskan Taiga: a synthesis of structure and function*. (eds K. Van Cleve, F.S. Chapin III, P.W. Flanagan, L.A. Viereck & C.T. Dyrness), pp. 22–43. Springer, New York.
- Wong C.Y.S. & Gamon J.A. (2015a) Three causes of variation in the photochemical reflectance index (PRI) in evergreen conifers. *New Phytologist* **206**, 187–195.
- Wong C.Y.S. & Gamon J.A. (2015b) The photochemical reflectance index provides an optical indicator of spring photosynthetic activation in evergreen conifers. *New Phytologist* **206**, 196–208.
- Yamori W. & von Caemmerer S. (2009) Effect of rubisco activase deficiency on the temperature response of CO<sub>2</sub> assimilation rate and rubisco activation state: Insights from transgenic tobacco with reduced amounts of rubisco activase. *Plant Physiology* **151**, 2073–2082.
- Yin X. (1993) Variation in foliar nitrogen concentration by forest type and climatic gradients in North America. *Canadian journal of forest research* **23**, 1587–1602.
- Zweifel R., Böhm J.P. & Häsler R. (2002) Midday stomatal closure in Norway spruce - Reactions in the upper and lower crown. *Tree Physiology* **22**, 1125–1136.

## TABLES

**Table 1.** Means  $\pm$  1 standard error for characteristics of trees from the Forest Tundra Ecotone (FTE), Alaska (n=18), and Black Rock Forest (BRF), New York (n = 6), including diameter at breast height (DBH; cm), tree height (m), dripline area (m<sup>2</sup>).

Location	DBH (cm)	Tree Height (m)	Dripline area (m <sup>2</sup> )
BRF	23.10 (1.99)	9.91 (0.73)	27.34 (2.00)
FTE	16.71 (0.94)	9.32 (0.57)	3.52 (0.37)



**Table 2.** Parameter estimates for the linear mixed effects models in which each physiological, leaf and spectral trait was modeled as a function of integrated average daily photochemical photon flux density (PPFD) and location (either Black Rock Forest (BRF), New York, or the Forest Tundra Ecotone (FTE), Alaska) with a random effect for each individual tree (Parameter ~ PPFD \* location + (1|tree)). Values in parentheses are SEs. Starred values are those that are significantly different from zero (\*\*\*P<0.001; \*\*P<0.01; \*P<0.05; +P<0.1). Significant differences between locations are indicated by different letters (P<0.05). Marginal and conditional coefficients of determination ( $r_m^2$  and  $r_c^2$ , respectively) are also reported.

Parameter	INTERCEPTS		SLOPES		$r_m^2$	$r_c^2$
	BRF	FTE	BRF	FTE		
<b>Photosynthesis</b>						
$A_{1500}$	9.282 (0.730)*** a	9.542 (1.389)*** a	0.066 (0.024)** a	-0.075 (0.054) <sup>+</sup> b	0.4	0.5
$\Phi$	0.055 (0.003)*** a	0.039 (0.004)*** b	-2e-04 (1e-04)** a	-3e-04 (2e-04) <sup>+</sup> a	0.67	0.82
LSP	366.132 (29.038)*** a	540.489 (51.805)*** b	6.912 (0.855)*** a	4.185 (2.012)* a	0.57	0.69
log(LCP)	2.744 (0.116)*** a	4.170 (0.223)*** b	0.038 (0.004)*** a	0.014 (0.009) b	0.75	0.79
$V_{cmax}$	43.123 (3.494)*** a	68.472 (6.867)*** b	0.316 (0.112)** a	-0.796 (0.270)** b	0.17	0.31
$J_{max}$	73.760 (5.499)*** a	108.241 (10.866)*** b	0.531 (0.178)** a	-1.092 (0.427)* b	0.17	0.31
<b>Respiration</b>						
$R_D$	0.72 (0.211)** a	2.321 (0.411)*** b	0.055 (0.007)*** a	0.006 (0.016) b	0.48	0.58
$R_L$	0.299 (0.228) a	1.726 (0.403)*** b	0.055 (0.007)*** a	0.020 (0.016) b	0.52	0.62
<b>Leaf traits</b>						
% Carbon	47.498 (0.514)*** a	49.003 (0.630)*** a	0.020 (0.007)** a	-0.015 (0.023) a	0.08	0.8
% Nitrogen	1.064 (0.052)*** a	0.972 (0.095)*** a	0.002 (0.001) a	4e-04 (0.004) a	0.17	0.53
$C_{area}$	702.603 (58.824)*** a	1481.729 (139.897)*** b	13.332 (2.136)*** a	2.254 (5.484) a	0.71	0.71
$N_{area}$	15.084 (1.684)*** a	26.773 (2.842)*** b	0.349 (0.037)*** a	0.176 (0.109) a	0.58	0.81
log(C:N)	3.822 (0.053)*** a	3.9303 (0.100)*** a	-0.002 (0.001) a	-9e-04 (0.004) a	0.18	0.54
SLA	65.803 (2.053)*** a	34.393 (4.748)*** b	-0.676 (0.072)*** a	-0.089 (0.186) b	0.76	0.77
<b>Pigments</b>						
$Chl_{area}$	77.180 (5.705)*** a	89.363 (11.060)*** a	0.325 (0.178) <sup>+</sup> a	0.288 (0.407) a	0.21	0.44
$Chl_{aarea}$	48.129 (3.576)*** a	54.266 (6.907)*** a	0.304 (0.111)* a	0.367 (0.254) a	0.3	0.51
$Chl_{barea}$	29.06 (2.353)*** a	35.186 (4.55)*** a	0.021 (0.073) a	-0.082 (0.167) a	0.09	0.36
$Car_{area}$	6.058 (0.905)*** a	10.722 (1.795)*** b	0.170 (0.030)*** a	0.160 (0.066)* a	0.67	0.74
Chl:Car <sub>area</sub>	11.583 (0.461)*** a	7.973 (0.901)*** b	-0.116 (0.015)*** a	-0.052 (0.033) a	0.69	0.77
Chl <sub>a</sub> :Chl <sub>b</sub> <sub>area</sub>	1.673 (0.070)*** a	1.510 (0.128)*** a	0.009 (0.002)*** a	0.017 (0.005)*** a	0.35	0.63
<b>Spectral Indices</b>						
PRI	0.051 (0.006)*** a	0.005 (0.015) b	-9e-04 (2e-04)*** a	8e-04 (6e-04) b	0.24	0.31

Abbreviations:  $A_{1500}$ , photosynthesis at 1500  $\mu\text{mol m}^{-2}\text{s}^{-1}$  ( $\mu\text{mol m}^{-2}\text{s}^{-1}$ );  $\Phi$ , apparent quantum yield; LSP, light saturation point ( $\mu\text{mol m}^{-2}\text{s}^{-1}$ ); LCP, light compensation point ( $\mu\text{mol m}^{-2}\text{s}^{-1}$ );  $V_{cmax}$ , maximum carboxylation rate ( $\mu\text{mol m}^{-2}\text{s}^{-1}$ );  $J_{max}$ , maximum electron transport ( $\mu\text{mol m}^{-2}\text{s}^{-1}$ );  $R_D$ , respiration in the dark ( $\mu\text{mol m}^{-2}\text{s}^{-1}$ );  $R_L$ , respiration in the light ( $\mu\text{mol m}^{-2}\text{s}^{-1}$ );  $C_{area}$ , carbon per leaf area ( $\text{mg C cm}^{-2}$ );  $N_{area}$ , nitrogen per leaf area ( $\text{mg N cm}^{-2}$ ); C:N, carbon to nitrogen ratio; SLA, specific leaf area ( $\text{m}^{-2}\text{g}$ );  $Chl_{area}$ , total chlorophyll on an area basis ( $\mu\text{g cm}^{-2}$ );

$Chl_{a_{area}}$ , area-based chlorophyll *a* content ( $\mu\text{g cm}^{-2}$ );  $Chl_{b_{area}}$ , area-based chlorophyll *b* content ( $\mu\text{g cm}^{-2}$ );  $Car_{area}$ , area-based carotenoid content ( $\mu\text{g cm}^{-2}$ );  $Chl:Car_{area}$ , ratio of total chlorophyll content to carotenoids;  $Chl_a:Chl_{b_{area}}$ , ratio of chlorophyll *a* to chlorophyll *b*; PRI, photochemical reflectance index.

**Table 3.** Summaries of the mixed effects models in which parameters were modeled as a function of interaction between canopy position and location with a random effect for each individual tree. P-values are estimated using the Kenward-Roger approximation of the degrees of freedom as suggested by (Halekoh & Højsgaard 2014). See table 2 for explanations of abbreviations and units. P<0.05 are bolded.

	Canopy		Location		Canopy x Location	
	<i>F</i> -value	<i>p</i> -value	<i>F</i> -value	<i>p</i> -value	<i>F</i> -value	<i>p</i> -value
$A_{1500}$	0.14	0.7106	26.14	<b>&lt;0.001</b>	13.28	<b>&lt;0.001</b>
AQY	15.57	<b>&lt;0.001</b>	60.94	<b>&lt;0.001</b>	0.68	0.4129
LSP <sub>1500</sub>	59.20	<b>&lt;0.001</b>	27.38	<b>&lt;0.001</b>	11.31	<b>0.0015</b>
log(LCP)	94.83	<b>&lt;0.001</b>	139.98	<b>&lt;0.001</b>	41.65	<b>&lt;0.001</b>
$V_{\text{cmax}}$	1.35	0.2506	0.23	0.6381	23.37	<b>&lt;0.001</b>
$J_{\text{max}}$	0.65	0.4233	1.00	0.3333	23.82	<b>&lt;0.001</b>
$R_D$	55.17	<b>&lt;0.001</b>	7.61	<b>0.0146</b>	46.38	<b>&lt;0.001</b>
$R_L$	49.13	<b>&lt;0.001</b>	10.09	<b>0.006</b>	25.33	<b>&lt;0.001</b>
% Carbon	1.55	0.2194	1.41	0.2482	12.79	<b>&lt;0.001</b>
% Nitrogen	0.09	0.7625	5.97	<b>0.0265</b>	1.88	0.1757
$C_{\text{area}}$	70.78	<b>&lt;0.001</b>	148.02	<b>&lt;0.001</b>	4.19	<b>0.0452</b>
$N_{\text{area}}$	77.16	<b>&lt;0.001</b>	25.95	<b>&lt;0.001</b>	19.83	<b>&lt;0.001</b>
log(C:N)	0.05	0.8181	6.25	<b>0.0236</b>	1.58	0.2146
SLA	90.17	<b>&lt;0.001</b>	134.52	<b>&lt;0.001</b>	31.35	<b>&lt;0.001</b>
$\text{Chl}_{\text{area}}$	2.68	0.1125	5.32	<b>0.0363</b>	0.02	0.8997
$\text{Chl}a_{\text{area}}$	8.12	<b>0.0081</b>	6.27	<b>0.0246</b>	0.05	0.825
$\text{Chl}b_{\text{area}}$	0.10	0.7494	3.09	0.0997	0.0004	0.9843
$\text{Car}_{\text{area}}$	49.82	<b>&lt;0.001</b>	22.34	<b>&lt;0.001</b>	3.56	0.0703
Chl:Car <sub>area</sub>	53.41	<b>&lt;0.001</b>	31.14	<b>&lt;0.001</b>	23.67	<b>&lt;0.001</b>
Chl $a$ :Chl $b$ <sub>area</sub>	34.49	<b>&lt;0.001</b>	0.37	0.5532	0.42	0.5213
PRI	7.37	<b>0.0094</b>	0.33	0.5751	20.94	<b>&lt;0.001</b>

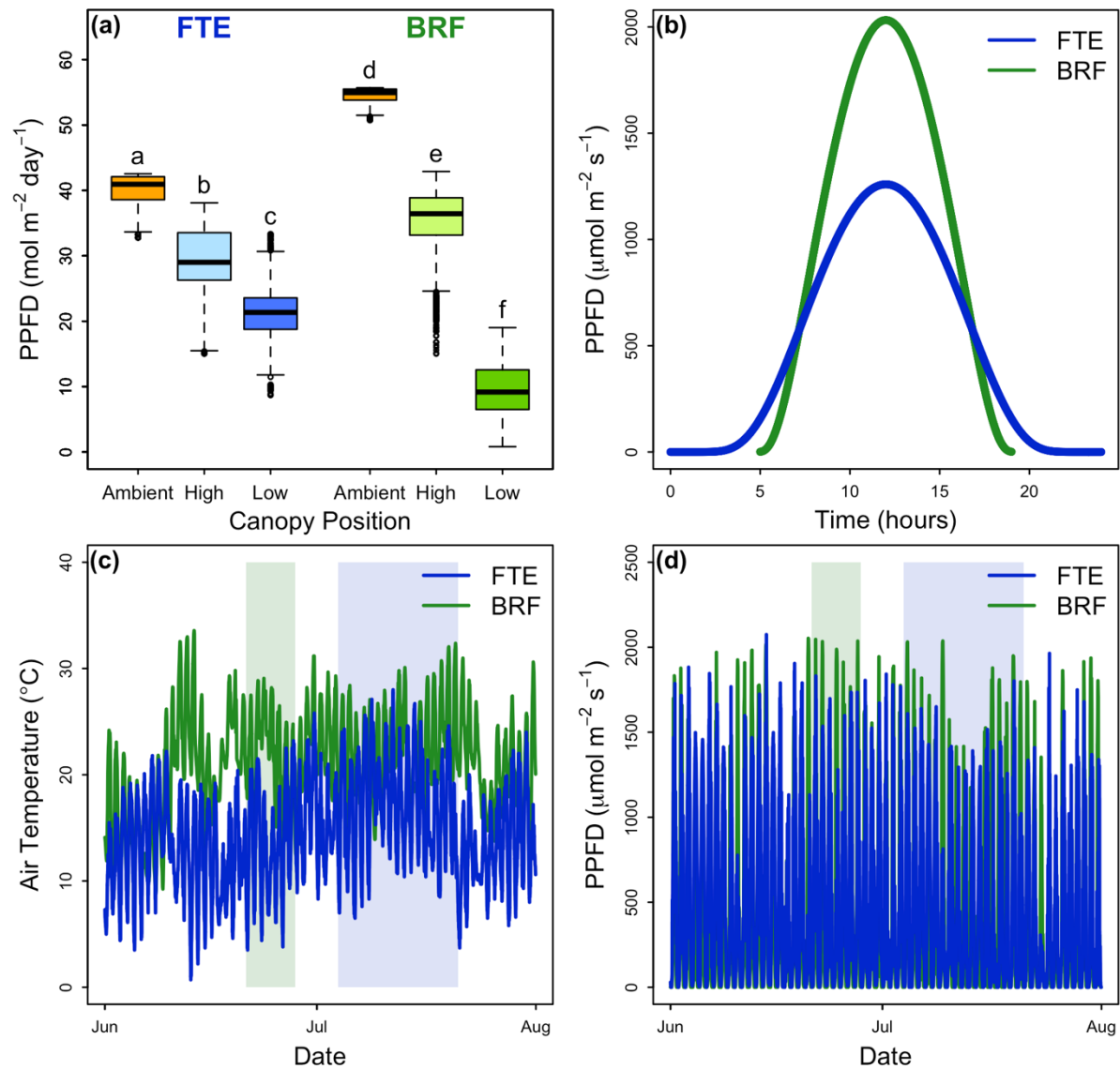
**Table 4.** Means  $\pm$  1 standard error for physiological, leaf and spectral traits. Several traits report back-transformed means including the light compensation point (LCP) and the ratio of carbon to nitrogen (C:N). Significant differences are marked by different letters ( $P < 0.05$ ) with differences between locations marked by capital letters and differences between location and canopy positions marked by lowercase letters. See table 2 for explanations of abbreviations and units. See table 2 for explanations of abbreviations and units.

	BRF	FTE	BRF low	BRF high	FTE low	FTE high
$A_{1500}$	10.8 (0.5) A	7.7 (0.4) B	9.8 (0.6) b	11.7 (0.6) a	8.3 (0.5) bc	7.0 (0.5) c
AQY	0.05 (0.002) A	0.03 (0.001) B	0.05 (0.002) a	0.05 (0.002) b	0.03 (0.002) c	0.03 (0.002) c
LSP <sub>1500</sub>	522.0 (18.6) A	643.9 (14.1) B	427.4 (22.5) a	616.7 (22.2) bc	605.6 (17.8) b	682.2 (18.4) c
LCP	36.8 (2.1) A	90.7 (4.5) B	20.7 (1.6) a	65.1 (5.0) b	79.7 (5.3) b	103.1 (7.2) c
$V_{cmax}$	50.3 (2.2) A	48.5 (1.7) A	46.2 (2.7) ab	54.5 (2.7) c	54.7 (2.2) bc	42.3 (2.35) a
$J_{max}$	85.9 (3.4) A	80.8 (2.7) A	78.7 (4.1) ab	93.1 (4.2) c	90.1 (3.5) bc	71.5 (3.7) a
$R_D$	1.98 (0.14) A	2.46 (0.11) B	1.14 (0.17) a	2.82 (0.17) b	2.41 (0.13) b	2.5 (0.14) b
$R_L$	1.58 (0.16) A	2.22 (0.12) B	0.77 (0.19) a	2.38 (0.19) b	2.05 (0.14) b	2.39 (0.14) b
% Carbon	47.96 (0.50) A	48.63 (0.30) A	47.64 (0.51) a	48.27 (0.51) b	48.81 (0.32) ab	48.45 (0.32) ab
% Nitrogen	1.1 (0.04) A	0.98 (0.03) B	1.09 (0.05) ab	1.12 (0.05) b	1.00 (0.03) ab	0.96 (0.04) a
$C_{area}$	1007.2 (33.5) A	1547.8 (29.5) B	813.4 (42.8) a	1201.1 (42.1) b	1430.0 (38.2) c	1665.5 (40.9) d
$N_{area}$	23.1 (1.3) A	31.2 (0.9) B	18.3 (1.4) a	27.9 (1.4) bc	29.7 (1.0) b	32.7 (1.1) c
C:N	43.9 (1.8) A	49.9 (1.5) B	44.4 (2.0) ab	43.3 (2.0) a	48.9 (1.7) ab	50.9 (1.8) b
SLA	50.4 (1.2) A	32.0 (1.1) B	60.2 (1.5) d	40.5 (1.5) c	34.5 (1.4) b	29.5 (1.5) a
$Chl_{area}$	84.6 (3.9) A	96.0 (3.1) B	81.4 (4.8) a	87.7 (4.5) ab	93.3 (4.6) ab	98.7 (3.4) b
$Chla_{area}$	55.0 (2.5) A	62.9 (2.0) B	51.5 (3.0) a	58.4 (2.9) ab	60.0 (2.9) ab	65.9 (2.2) b
$Chlb_{area}$	29.6 (1.6) A	33.1 (1.3) A	29.8 (1.9) a	29.3 (1.8) a	33.3 (1.9) a	32.9 (1.4) a
$Car_{area}$	9.9 (0.8) A	14.5 (0.5) B	7.6 (0.9) a	12.1 (0.8) b	13.2 (0.7) b	15.7 (0.6) c
$Chl:Car_{area}$	9.0 (0.3) A	6.7 (0.2) B	10.7 (0.4) b	7.3 (0.3) a	7.0 (0.3) a	6.4 (0.3) a
$Chla:Chlb_{area}$	1.87 (0.05) A	1.92 (0.04) A	1.75 (0.06) a	2.00 (0.05) bc	1.81 (0.05) ab	2.02 (0.04) c
PRI	0.03 (0.004) A	0.02 (0.004) A	0.04 (0.005) b	0.02 (0.005) a	0.02 (0.005) a	0.03 (0.004) ab

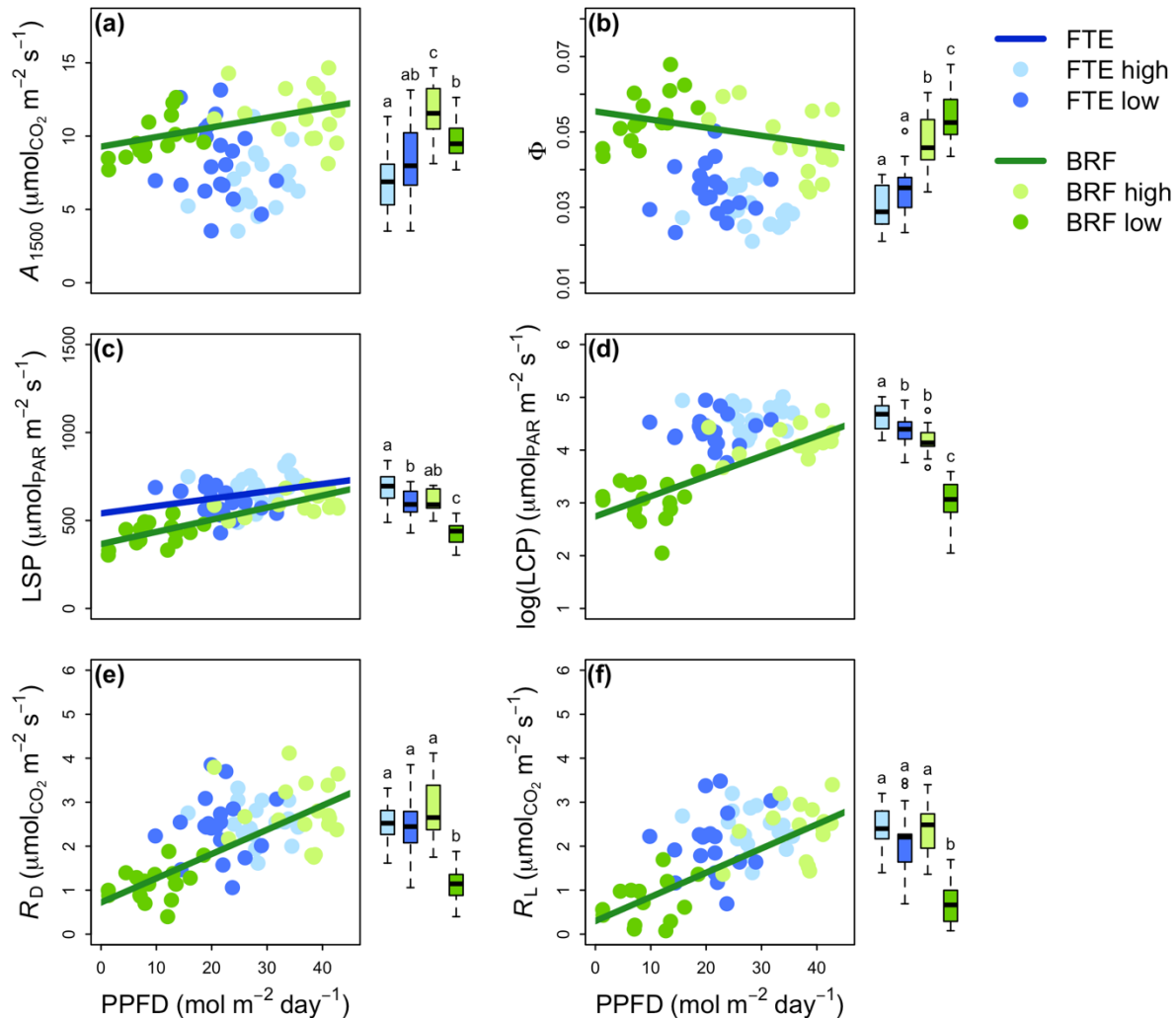
**Table 5.** Parameter estimates for the intercepts and slopes of linear mixed effects models with tree as a random effect examining relationships between photosynthesis at 1500  $\mu\text{mol m}^{-2}\text{s}^{-1}$  ( $A_{1500}$ ), maximum rate of carboxylation ( $V_{\text{cmax}}$ ), respiration in the dark ( $R_{\text{D}}$ ), respiration in the light ( $R_{\text{L}}$ ) or total pigment content ( $\text{Chl}+\text{Car}_{\text{area}}$ ) and nitrogen per leaf area ( $N_{\text{area}}$ ); as well as the relationship between photochemical reflectance index (PRI) and the chlorophyll to carotenoid ratio ( $\text{Chl}:\text{Car}_{\text{area}}$ ). In all models tree was included as a random effect. Values in parentheses are standard SEs. Starred values are those that are significantly different from zero (\*\*\* $P<0.001$ ; \*\* $P<0.01$ ; \* $P<0.05$ ; + $P<0.1$ ). Significant differences between locations are indicated by different letters ( $P<0.05$ ). Marginal and conditional coefficients of determination ( $r_{\text{m}}^2$  and  $r_{\text{c}}^2$ , respectively) are also reported.

Model	INTERCEPTS		SLOPES		$r_{\text{m}}^2$	$r_{\text{c}}^2$
	BRF	FTE	BRF	FTE		
$A_{1500} \sim N_{\text{area}} * \text{location}$	6.181 (1.250)*** a	8.130 (2.220)*** a	0.200 (0.049)*** a	-0.024 (0.071) b	0.5	0.64
$V_{\text{cmax}} \sim N_{\text{area}} * \text{location}$	28.828 (6.253)*** a	61.318 (12.106)*** b	0.934 (0.230)*** a	-0.440 (0.383) b	0.15	0.55
$R_{\text{D}} \sim N_{\text{area}} * \text{location}$	-0.761 (0.410)+ a	0.714 (0.620) a	0.118 (0.017)*** a	0.055 (0.020)** b	0.49	0.49
$R_{\text{L}} \sim N_{\text{area}} * \text{location}$	-0.908 (0.518)+ a	0.386 (0.723) a	0.113 (0.021)*** a	0.059 (0.023)* a	0.42	0.42
$(\text{Chl}+\text{Car}_{\text{area}}) \sim N_{\text{area}} * \text{location}$	62.309 (11.765)*** a	63.348 (22.613)** a	1.436 (0.501)** a	1.518 (0.712)* a	0.39	0.52
$\text{PRI} \sim \text{Chl}:\text{Car}_{\text{area}} * \text{location}$	-0.060 (0.015)*** a	0.033 (0.023) b	0.010 (0.002)*** a	-0.0008 (0.003) b	0.43	0.54

## FIGURES

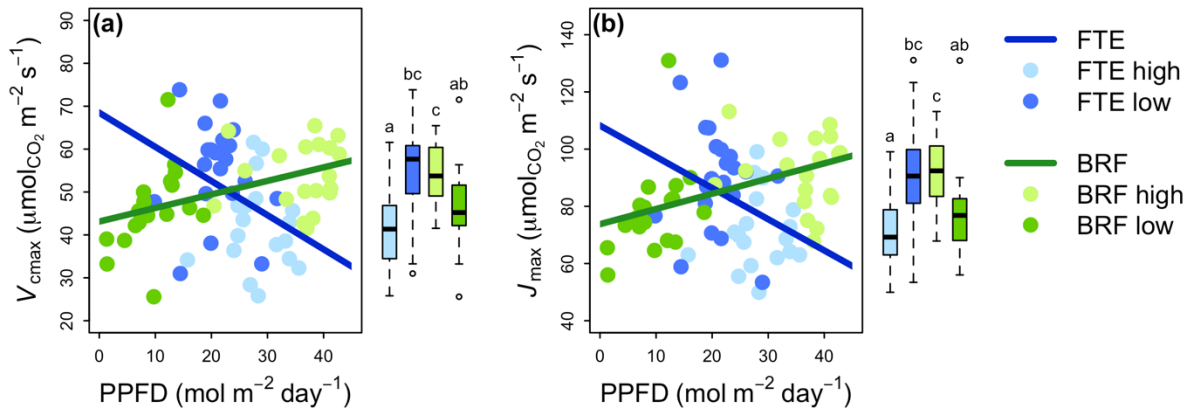


**Figure 1.** (a) Average daily PPFD calculated from canopy photos over the study period (June-July) at the Forest Tundra Ecotone (FTE), Alaska, and Black Rock Forest (BRF), New York. Ambient (above canopy) PPFD is shown in orange. Boxplots show the median and first and third quartiles. Whiskers display the range of groups with individual points representing outliers falling outside 1.5 times the interquartile range. Different letters represent significant differences between locations and canopy positions ( $P < 0.05$ ). (b) Ambient PPFD projected from canopy photos during one day (July 4, 2017) at both locations. (c) Air temperature (°C) measured at BRF and the FTE for June and July 2017. (d) Ambient PPFD measured from field instruments at BRF and the FTE for June and July 2017. Green and blue shaded areas in 1c and 1d denote the dates of the study campaigns at BRF and the FTE, respectively.

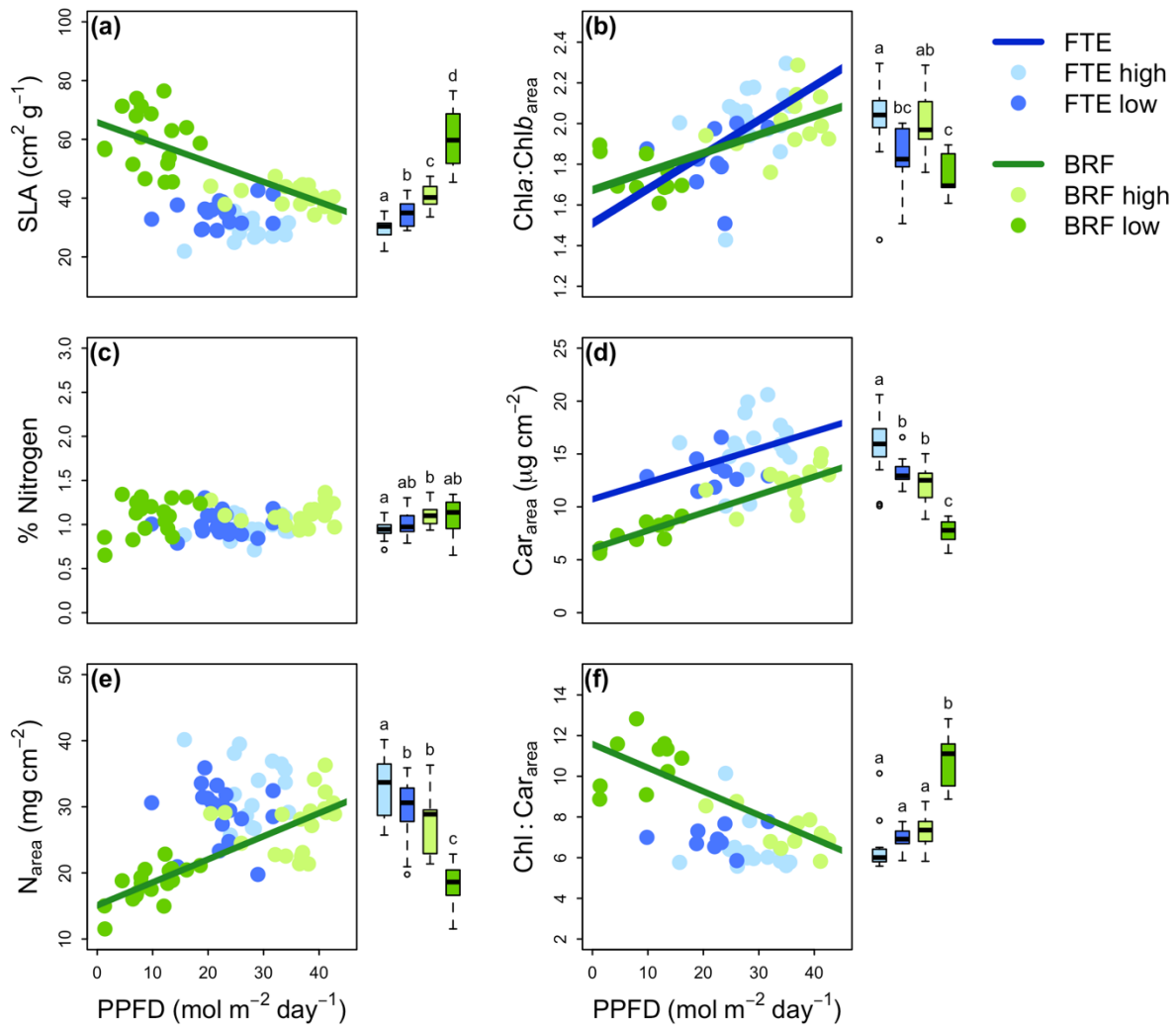


**Figure 2.** Linear relationships between average daily photosynthetic photon flux density (PPFD) over the measurement campaign and area-based foliar photosynthetic characteristics of white spruce from the Forest Tundra Ecotone (FTE; blue), Alaska, and Black Rock Forest (BRF; green), New York including photosynthesis at 1500  $\mu\text{mol m}^{-2}\text{s}^{-1}$  ( $A_{1500}$ ; 2a; FTE  $n = 38$ ; BRF  $n = 33$ ), apparent quantum yield ( $\Phi$ ; 2b; FTE  $n = 38$ ; BRF  $n = 33$ ), light saturation point (LSP; 2c; FTE  $n = 38$ ; BRF  $n = 33$ ), light compensation point (LCP; 2d; FTE  $n = 38$ ; BRF  $n = 33$ ), respiration in the dark ( $R_D$ ; 2e; FTE  $n = 38$ ; BRF  $n = 36$ ) and respiration in the light ( $R_L$ ; 2f; FTE  $n = 37$ ; BRF  $n = 29$ ). Light and dark colors at each location (BRF or FTE) represent high and low canopy positions, respectively. Parameter estimates of the linear mixed effects regression models, and statistical differences between slopes and intercepts are presented in Tables 2 & 3. Regression lines are only shown for significant relationships (slope  $P < 0.05$ ). Also included are boxplots by canopy position and location for each parameter. Different letters represent significant differences between locations and canopy positions ( $P < 0.05$ ).

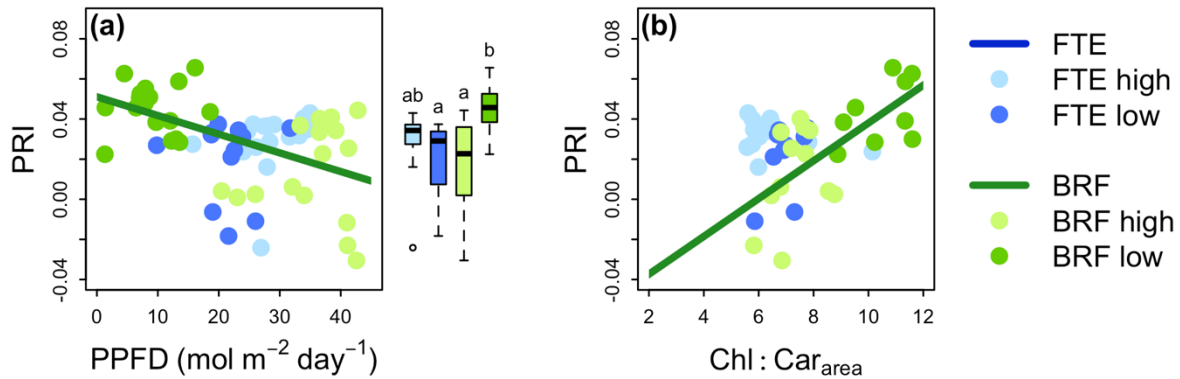




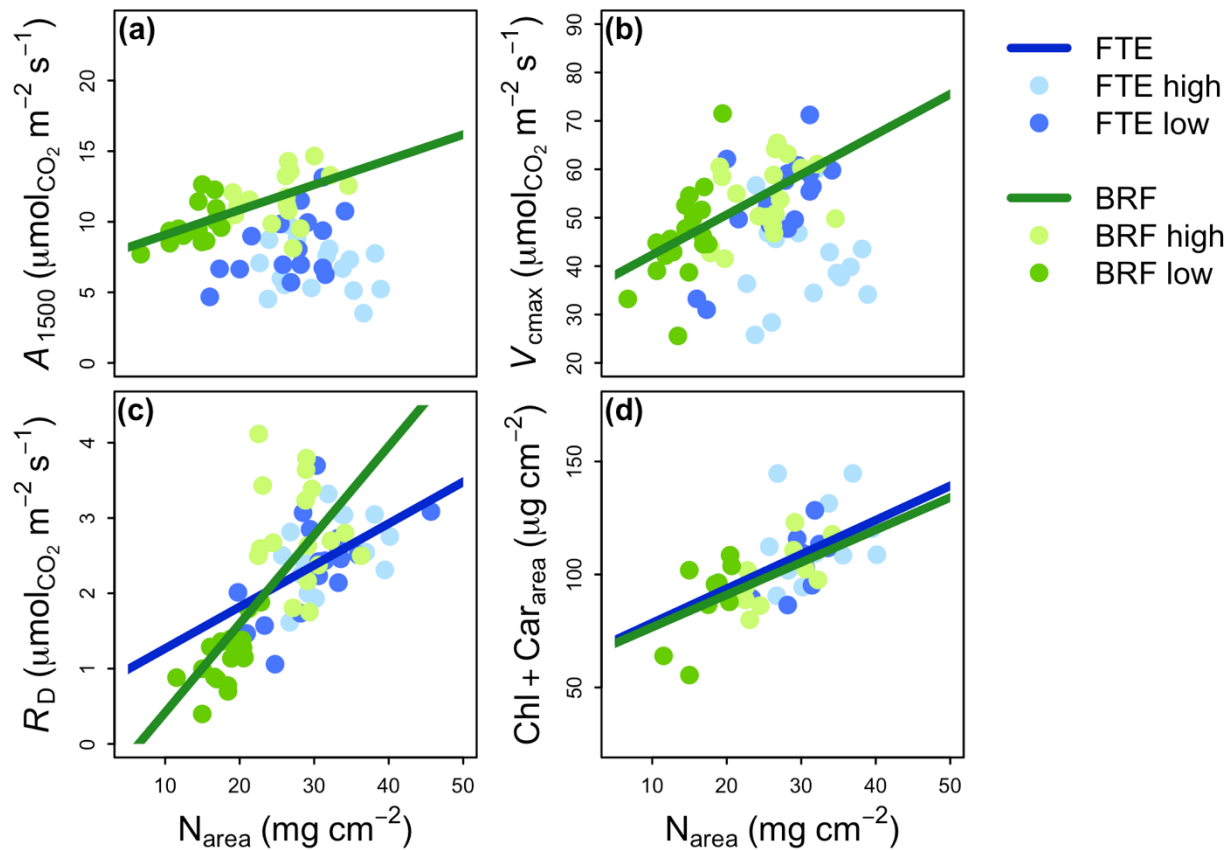
**Figure 3.** Linear relationships between average daily photosynthetic photon flux density (PPFD) and foliar respiratory characteristics on an area basis of white spruce from the Forest Tundra Ecotone (FTE; blue), Alaska, and Black Rock Forest (BRF; green), New York including the maximum rate of carboxylation ( $V_{\text{max}}$ ; 3a; FTE  $n = 39$ ; BRF  $n = 37$ ), and the maximum electron transport rate ( $J_{\text{max}}$ ; 3b; FTE  $n = 39$ ; BRF  $n = 37$ ). Light and dark colors at each location (BRF or FTE) represent high and low canopy positions, respectively. Parameter estimates of the linear mixed effects regression models, and statistical differences between slopes and intercepts are presented in Tables 2 & 3. Regression lines are only shown for significant relationships (slope  $P < 0.05$ ). Also included are boxplots by canopy position and location for each respiratory parameter. Different letters represent significant differences between locations and canopy positions ( $P < 0.05$ ).



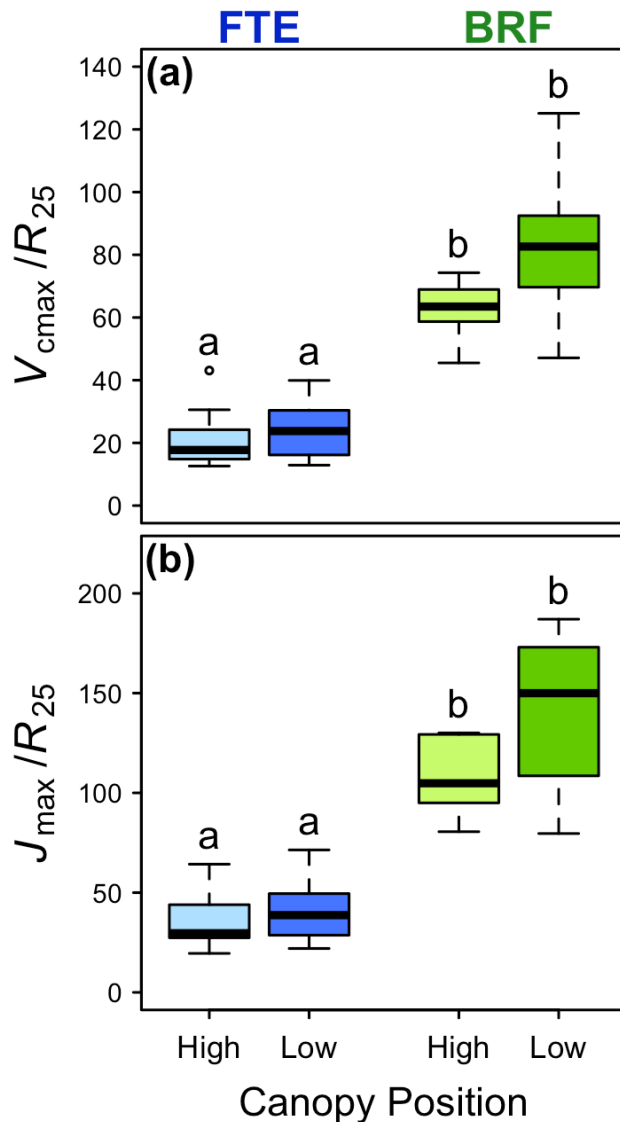
**Figure 4.** Linear relationships between average daily photosynthetic photon flux density (PPFD) and leaf traits of white spruce from the Forest Tundra Ecotone (FTE; blue), Alaska, and Black Rock Forest (BRF; green), New York including specific leaf area (SLA; 4a; FTE  $n = 37$ ; BRF  $n = 38$ ), the ratio of chlorophyll *a* to chlorophyll *b* (Chl $a$ :Chl $b$ <sub>area</sub>; 4b; FTE  $n = 25$ ; BRF  $n = 22$ ), % nitrogen (4c; FTE  $n = 37$ ; BRF  $n = 38$ ), carotenoids (Car<sub>area</sub>; 4d; FTE  $n = 25$ ; BRF  $n = 22$ ), nitrogen per leaf area (N<sub>area</sub>; 4e; FTE  $n = 37$ ; BRF  $n = 38$ ), and the ratio of chlorophylls *a* + *b* to carotenoids (Chl:Car<sub>area</sub>; 4f; FTE  $n = 25$ ; BRF  $n = 22$ ). Light and dark colors at each location (BRF or FTE) represent high and low canopy positions, respectively. Parameter estimates of the linear mixed effects models, and statistical differences between slopes and intercepts are presented in Tables 2 & 3. Regression lines are only shown for significant relationships (slope  $P < 0.05$ ). Also included are boxplots by canopy position and location for each leaf trait. Different letters represent significant differences between locations and canopy positions ( $P < 0.05$ ).



**Figure 5.** Linear relationships between photochemical reflectance index (PRI) and either average daily photosynthetic photon flux density (PPFD) (5a; FTE  $n = 30$ ; BRF  $n = 34$ ) or the ratio of chlorophylls  $a + b$  to carotenoids (Chl:Car<sub>area</sub>; 5b) of white spruce from the Forest Tundra Ecotone (FTE; blue), Alaska, and Black Rock Forest (BRF; green), New York. Light and dark colors at each location (BRF or FTE) represent high and low canopy positions, respectively. Parameter estimates of the linear mixed effects regression models, and statistical differences between slopes and intercepts are presented in Tables 2 & 3. Regression lines are only shown for significant relationships (slope  $P < 0.05$ ). Also included are boxplots by canopy position and location for PRI. Different letters represent significant differences between locations and canopy positions ( $P < 0.05$ ).



**Figure 6.** Linear relationships between photosynthesis at 1500  $\mu\text{mol m}^{-2}\text{s}^{-1}$  ( $A_{1500}$ ; 6a), maximum rate of carboxylation ( $V_{\text{cmax}}$ ; 6b), respiration in the dark ( $R_{\text{D}}$ ; 6c) and total pigment content ( $\text{Chl} + \text{Car}_{\text{area}}$ ; 6d) versus nitrogen per leaf area ( $N_{\text{area}}$ ) of white spruce from the Forest Tundra Ecotone (FTE; blue), Alaska and Black Rock Forest (BRF; green), New York. Light and dark colors at each location (BRF or FTE) represent high and low canopy positions, respectively. Parameter estimates of the linear mixed effects regression models, and statistical differences between slopes and intercepts are presented in Table 5. Regression lines are only shown for significant relationships (slope  $P < 0.05$ ).



**Figure 7.** Ratios of (a) maximum rate of carboxylation to respiration at 25°C from Griffin *et al.* (in revision) ( $V_{cmax}/R_{25}$ ) and (b) maximum electron transport rate to respiration at 25°C from Griffin *et al.* (in revision) ( $J_{max}/R_{25}$ ) of white spruce from high and low canopy positions at the Forest Tundra Ecotone (FTE, blue), Alaska, and Black Rock Forest (BRF, green), New York. Boxplots show the median and first and third quartiles. Whiskers display the range of groups with individual points representing outliers falling outside 1.5 times the interquartile range. Different letters represent significant differences between locations and canopy positions ( $P < 0.05$ ).