Coordination of leaf structure and gas exchange along a height gradient in a tall conifer

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Summary The gravitational component of water potential and frictional resistance during transpiration lead to substantial reductions in leaf water potential (Ψ_1) near the tops of tall trees, which can influence both leaf growth and physiology. We examined the relationships between morphological features and gas exchange in foliage collected near the tops of Douglas-fir (Pseudotsuga menziesii (Mirb.) Franco) trees of different height classes ranging from 5 to 55 m. This sampling allowed us to investigate the effects of tree height on leaf structural characteristics in the absence of potentially confounding factors such as irradiance, temperature, relative humidity and branch length. The use of cut foliage for measurement of intrinsic gas-exchange characteristics allowed identification of height-related trends without the immediate influences of path length and gravity. Stomatal density, needle length, needle width and needle area declined with increasing tree height by $0.70 \text{ mm}^{-2} \text{ m}^{-1}$, 0.20 mm m^{-1} , $5.9 \times 10^{-3} \text{ mm m}^{-1}$ and $0.012 \text{ mm}^{2} \text{ m}^{-1}$, respectively. Needle thickness and mesophyll thickness increased with tree height by $4.8 \times 10^{-2} \text{ mm m}^{-1}$ and $0.74 \ \mu m \ m^{-1}$, respectively. Mesophyll conductance (g_m) and CO_2 assimilation in ambient $[CO_2]$ (A_{amb}) decreased by 1.1 mmol $m^{-2} s^{-1}$ per m and 0.082 µmol $m^{-2} s^{-1}$ per m increase in height, respectively. Mean reductions in g_m and A_{amb} of foliage from 5 to 55 m were 47% and 42%, respectively. The observed trend in A_{amb} was associated with $g_{\rm m}$ and several leaf anatomic characteristics that are likely to be determined by the prevailing vertical tension gradient during foliar development. A linear increase in foliar δ^{13} C values with height (0.042% m⁻¹) implied that relative stomatal and mesophyll limitations of photosynthesis in intact shoots increased with height. These data suggest that increasing height leads to both fixed structural constraints on leaf gas exchange and dynamic constraints related to prevailing stomatal behavior.

Keywords: growth limitation, leaf anatomy, mesophyll conductance, photosynthesis, Pseudotsuga menziesii.

Introduction

Several mechanisms have been studied as factors potentially responsible for reduced growth in trees as they age and increase in height, none of which are mutually exclusive. Proposed mechanisms include, but are not limited to, reduced leaf area as a result of reduced nutrient availability (Gower et al. 1996), genetics-related reductions in photosynthetic capacity induced by a limited capacity for repeated meristematic divisions (Haffner et al. 1991), reduced photosynthesis in response to increased hydraulic resistance with increased tree height (Mencuccini and Grace 1996, Ryan and Yoder 1997, McDowell et al. 2002), reduced foliar expansion due to lower turgor pressure (Koch et al. 2004, Woodruff et al. 2004), and reduced leaf hydraulic efficiency as a result of height-related restrictions on leaf expansion (Woodruff et al. 2008). Available evidence suggests that it is the size (height) as opposed to the age that is primarily responsible for ontogenetic patterns in growth (Koch et al. 2004, Woodruff et al. 2004, Bond et al. 2007, Mencuccini et al. 2007). Studies on how height affects gas exchange in trees have not yet parsed out the extrinsic effects of path length and gravity from the intrinsic effects of height-related trends in foliar structural characteristics on gas exchange.

Leaf structural characteristics can influence gas exchange through their effects on the efficiency or capacity of processes such as light absorption, carbon fixation and control of water loss. Leaf structural characteristics that influence gas exchange can vary in response to changes or gradients in environmental conditions (reviewed in Abrams 1994, Gutschick 1999), in response to increased age or height (Apple et al. 2002, England and Attiwill 2006) and in response to vertical water potential gradients (Koch et al. 2004, Woodruff et al. 2004, 2008). Mesophyll cells comprise the bulk of foliar tissue and represent the site of two types of resistance that can influence CO_2 assimilation (A): resistance to the transport of water from leaf veins and xylem conduits to stomata, and resistance to the movement of CO_2 from the intercellular air spaces to the sites of carboxylation inside the chloroplasts.

There are two predominant resistances to the diffusion of CO₂ as it moves from the atmosphere to the sites of carboxylation inside the leaf. Stomatal resistance occurs as CO2 diffuses through the stomata and into the sub-stomatal cavities. The less-studied type of foliar resistance to CO2 uptake is mesophyll resistance ($r_{\rm m}$, the inverse of mesophyll conductance, g_m) that involves the resistance to CO₂ diffusion through the mesophyll cell walls and membranes, and through the liquid phase inside the mesophyll cells. At the mesophyll cell surface, CO₂ enters the liquid phase and moves through the cell wall and through the chloroplast membrane to the site of carboxylation. Mesophyll resistance to CO₂ transfer can account for a substantial proportion of the total resistance to CO₂ transfer. Recent research has begun to reveal the importance of internal leaf conductance for gas exchange. Niinemets (1997) found a direct correlation between g_m and A in Picea abies (L.) H. Karst, both of which decreased with increasing tree age, and speculated that reductions in g_m may be a factor in size-related reductions in A for Picea abies and Pinus sylvestris L. (Niinemets 2002). Across a range of 50 species, internal leaf resistance accounted for about 40% of the decrease in CO2 concentration ([CO₂]) between the atmosphere and sites of carboxylation (Syvertsen et al. 1995, Hanba et al. 1999, Warren 2008 and references therein, Flexas et al. 2008). The determinants of internal leaf conductance are not yet fully understood, but are likely to include a combination of leaf anatomic and biochemical factors (Warren 2008). Leaf mesophyll structural characteristics that could affect the movement of either water or CO_2 are thus likely to have an effect on A.

Douglas-fir (*Pseudotsuga menziesii* (Mirb.) Franco) is one of the world's tallest tree species (Carder 1995) and can attain a height of > 100 m. The aim of this study was to determine the extent to which height imposes developmental constraints on Douglas-fir foliage that lead to intrinsic limitations for gas exchange. Two questions were investigated: (1) What effect does tree height have on the structural characteristics of Douglas-fir foliage? (2) Are there limitations on gas exchange associated with intrinsic leaf structural characteristics that can be attributed to path length and gravity during the time of foliar development? We hypothesized that height-related trends in leaf structure impose constraints on leaf gas exchange that are independent of the direct effects of vertical gradients in xylem tension on stomatal and photosynthetic physiology.

Materials and methods

Field site and sampling

Four stands, each containing Douglas-fir trees of a different height class, were located within 3.1 km of each other in the Wind River Basin of southwestern Washington, USA.

Access to tree tops in the 55-m sampling height class was facilitated by a 75-m-tall construction tower crane at the Wind River Canopy Crane Research Facility (WRCCRF). Periodic dieback of the tops of some of the trees within the WRCCRF stand suggested that these trees were close to their maximum height for this site. Tree tops in the two intermediate height classes were accessed by non-spur climbing and foliage from the lowest height class was accessed with a pole pruner. We collected 30-50-cm-long branch samples within 1-5 m of the tops of the trees at mean sampling heights of 5.0, 18.3, 33.5 and 55.0 m. All samples were obtained from fully sun-exposed locations during the summer and fall of 2007. Branches were collected from trees early in the morning before significant transpirational water loss and were placed in plastic bags with moist paper towels and stored in the dark at 5 °C.

The Pacific maritime climate of the region is characterized by wet winters and dry summers. Mean annual precipitation in the region is about 2.2 m, much of which falls as snow, with a dry season from June to September. Mean annual temperature is 8.7 °C with a mean of 0 °C in January and 17.5 °C in July. The soils are well drained and of volcanic origin (Shaw et al. 2004). Low precipitation between June and September (\sim 119 mm) typically leads to drought conditions in the upper portion of the soil profile. However, soil water remains accessible to Douglas-fir roots at depths greater than about 1 m throughout the summer dry period (Warren et al. 2005, Meinzer et al. 2007).

Leaf structural characteristics

We measured stomatal density (D_s) , length (L_n) , width (W_n) , thickness (T_n) , area (LA) and mesophyll thickness (T_m) of Douglas-fir needles. All needles were sampled exclusively from fully sun-exposed branches near the tops of trees of different height classes to rule out the potentially confounding influence of factors such as irradiance, relative humidity, and branch length upon height-related trends in leaf structural characteristics. Needle width, length and thickness were measured with digital calipers. Three needles were measured from n = 3 branches per tree. Leaf areas were measured on 10 needles per branch, on n = 3 branches per tree and three trees per height class. One-sided leaf areas were obtained using a scanner and ImageJ Version 1.27 image analysis software (Abramoff et al. 2004).

Mesophyll thickness ($T_{\rm m}$), which was used as a proxy for the distance necessary for water to move through mesophyll tissue inside the leaf, was estimated as the distance between the outside of the endodermis and the nearest inside edge of the leaf epidermis. Cross sections of needles were made by hand sectioning fresh tissue and mounting them on slides. Needle cross sections were analyzed for $T_{\rm m}$ with an image analysis system consisting of a compound microscope and a video camera. Three needles were analyzed from each of three branches, for a total of nine measurements per tree. Images were obtained using 10× objectives and a total magnification of 100×. Data were pooled per tree and subjected to regression analysis. Stomatal density was measured with a dissecting microscope and a linear scale in millimeter increments. Stomata were counted over a 1-mm length near the center of the needles. Number of stomata per millimeter of needle length was divided by needle width to generate number of stomata per mm² leaf area. Three needles were counted from each of n = 4-5 branches per tree.

Gas exchange and carbon isotope ratios

We evaluated several parameters related to A using photosynthetic CO₂ response curves, including A at ambient $[CO_2]$ (A_{amb}), maximum A (A_{max}), maximum carboxylation rate allowed by ribulose 1,5-bisphosphate carboxylase/ oxygenase (Rubisco) ($V_{\rm cmax}$), rate of photosynthetic electron transport (J), and mesophyll conductance (g_m) . These parameters were measured on branches sampled from the tops of trees of different height classes. We conducted measurements in the laboratory on detached shoots that had their bases immersed in water to eliminate the immediate effects of path length and gravity on gas exchange, thereby enabling us to isolate the influence of any height-related trends in foliar structure on the various gas exchange parameters. Before starting the gas-exchange measurement, we detached shoots of about 15-20 cm in length from the larger branch samples, taking care to submerge the shoot bases in degassed water as the cut was made. Our previous work on attached and detached Douglas-fir foliage showed that Douglas-fir shoots retain the same gas-exchange characteristics for about 4 days after detachment. In this study, mean A_{max} for samples taken at a tree height of 55 m at 0, 1 and 2 days following excision were 16.5, 14.6 and 19.7 µmol $m^{-2} s^{-1}$, respectively (P = 0.25 for regression analysis of $A_{\rm max}$ versus days since excision). This lack of a decrease in A_{max} suggests that the foliage was still fully active physiologically throughout the period of branch detachment while measurements were conducted.

Gas exchange was measured with a portable photosynthesis system equipped with a red or blue LED source and CO₂ injector (LI-6400, Li-Cor, Lincoln, NE). The instrument was zeroed and the chemicals were replaced prior to use each day. For determination of the dependence of CO₂ assimilation (A) on intercellular CO_2 concentration (A-C_i curves), photosynthetic photon flux was held at 1200 μ mol m⁻² s⁻¹, vapor pressure deficit at 1.0 kPa and leaf temperature at 25 °C. The cuvette [CO₂] was initially set near ambient, progressively lowered to 50 ppm, increased directly to ambient, and then progressively increased. The ambient [CO₂] were set at the following values and order: 400, 300, 200, 100, 50, 400, 400, 600, 800, 1000, 1200, and then increased at 200-ppm intervals until no further response of A was observed. Leaves were allowed to equilibrate for at least 2 min following the increase in [CO₂]. Branch samples were measured within 1-3 days of excision. Gas-exchange measurements were made on n = 2 branches per tree, and on three trees per height class.

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The $A-C_i$ curves were used to estimate g_m , V_{cmax} and J using a utility developed by Sharkey et al. (2007) based on an alternative $A-C_i$ curve fitting method (Ethier and Livingston 2004) that accounts for CO₂ transfer conductance through a non-rectangular hyperbola version of the model of Farquhar et al. (1980). In this model, the biochemical reactions of photosynthesis are considered to be limited either by the properties of Rubisco, the regeneration of the substrate ribulose bisphosphate (RuBP), or by triosephosphate use limitation. When A is Rubisco-limited, the response of A to [CO₂] can be described:

$$A = V_{\rm cmax} [C_{\rm C} - \Gamma^* / C_{\rm C} + K_{\rm C} (1 + O/K_{\rm O})] - R_{\rm d}, \quad (1)$$

where $C_{\rm C}$ and O are the partial pressures of $\rm CO_2$ and oxygen, respectively, at the sites of carboxylation, Γ^* is [CO₂] at which oxygenation proceeds at twice the rate of carboxylation causing photosynthetic uptake of CO₂ to be exactly compensated by photorespiratory CO₂ release, $K_{\rm C}$ is the Michaelis constant of Rubisco for CO₂, $K_{\rm O}$ is the inhibition constant of Rubisco for oxygen, and $R_{\rm d}$ is day respiration. When A is limited by RuBP regeneration:

$$A = J[C_{\rm C} - \Gamma^*/4C_{\rm C} + 8\Gamma^*] - R_{\rm d}, \qquad (2)$$

where J is rate of electron transport. In the $A-C_i$ utility used to analyze photosynthesis, $C_{\rm C}$ is replaced with $(C_i - A/g_m)$ and g_m is estimated from the observed data by nonlinear curve fitting and minimizing the sum of squared model deviations. Ideally a second method for determining $g_{\rm m}$, such as combined fluorescence and gas exchange, would be employed to verify the results from the curve analyses. However, the combined fluorescence and gas-exchange method is ill-suited for measurements of conifers (Ülo Niinemets, personal communication), and the previous research has shown a high correlation between the Ethier and Livingston (2004) method and the combined fluorescence and gas-exchange method (Niinemets et al. 2005). The accuracy of estimating $g_{\rm m}$ with the Sharkey utility can be enhanced by maximizing the number of values in the curvature region of the $A-C_i$ relationship (Ethier and Livingston 2004). The $A-C_{i}$ curves in this study were created and analyzed in a manner so as to minimize the sum of squares error in the models and maximize the number of values within the curvature region. The mean number of values in the curvature region of the curves was 6.8 (SE = 0.31), 5.8 (SE = 0.48), 8.8 (SE = 0.95) and 5.6 (SE = 0.37) for the foliage sampled from the trees at mean sampling heights of 5.0, 18.3, 33.5 and 55.0 m, respectively.

We determined the mean value of C_i at standard ambient [CO₂] (385 ppm) for all $A-C_i$ curves to characterize photosynthetic parameters at standard ambient [CO₂]. The mean leaf internal [CO₂] at 385 ppm sample [CO₂] was 235 ppm

Height (m)	$N_{\rm A}$ (g m ⁻²)	$\begin{array}{l} A_{\max} \\ (\mu \text{mol } \text{m}^{-2} \text{ s}^{-1}) \end{array}$	$A/g_{\rm s}$ (µmol mol ⁻¹)	$C_{\rm i}$ at $C_{\rm a}$ = 385 ppm (ppm)	$V_{\rm cmax} \\ (\mu {\rm mol} \ {\rm m}^{-2} \ {\rm s}^{-1})$	$\begin{array}{c}g_{\rm m}\\({\rm mmol}\ {\rm m}^{-2}\ {\rm s}^{-1})\end{array}$	$J \ (\mu mol m^{-2} s^{-1})$
5 (0)	1.9 (0.13)	21 (0.3)	94 (10)	224 (13)	95 (0.6)	106 (3.4)	104 (1.3)
18.3 (0.3)	1.8 (0.07)	20 (0.5)	96 (10)	219 (7)	105 (1.8)	84 (3.6)	112 (8.3)
33.5 (1.3)	1.8 (0.11)	21 (1.3)	72 (12)	248 (13)	139 (3.8)	43 (2.9)	134 (6.4)
55 (1.1)	1.5 (0.08)	17 (0.4)	62 (8)	252 (12)	77 (10)	56 (3.9)	104 (4.8)

Table 1. Characteristics of foliage sampled within 1–5 m of the tops of *P. menziesii* trees at mean sampling heights of 5.0, 18.3, 34.5 and 55.0 m (mean values with SE in parentheses). Internal [CO₂] at ambient [CO₂] (C_i at $C_a = 385$ ppm).

(SE = 6.25) (Table 1). Carbon dioxide assimilation at ambient [CO₂] (A_{amb}) was determined by first establishing a mean C_i value for ambient [CO₂] (385 ppm) and then determining A at that C_i value for each $A-C_i$ curve. Instantaneous water use efficiency (A/g_s) was determined by dividing A_{amb} by stomatal conductance (g_s) at $C_i = 235$ ppm.

Stable carbon isotope analysis was conducted on leaf tissue to determine integrated intrinsic water-use efficiency. Foliage on which gas exchange was measured plus foliage from one additional branch per tree was collected, dried and ground, and analyzed for δ^{13} C and nitrogen concentration ([N]) with a Carla Erba 1110 elemental analyzer coupled to a Thermo Finnigan Delta S isotope ratio mass spectrometer through an open split interface (Stable Isotope Ratio Facility for Environmental Research, University of Utah). Integrated water-use efficiency ($A/g_{s-integrated}$) was obtained as (Farquhar et al. 1982):

$$A/g_{\text{s-integrated}} = (C_{\text{a}} - C_{\text{i}})/1.6.$$
(3)

An analysis of the vertical trend in integrated, in situ photosynthesis ($A_{\text{integrated}}$) was performed by calculating Afrom needle δ^{13} C values. We determined discrimination (Δ) against 13 C in leaf tissue as (Farquhar et al. 1982):

$$\Delta = \delta \text{source} - \delta \text{product}/1 + (\delta \text{source}/1000), \quad (4)$$

where $\delta \text{source} = \delta^{13}\text{C}$ for atmospheric CO₂ (-8%), $\delta \text{product} = \delta$ and ¹³C for leaf tissue. We then used Eq. (5), also from Farquhar et al. (1982), to obtain C_i at standard atmospheric [CO₂] ($C_a = 385$ ppm):

$$\Delta = a + (b - a)C_{\rm i}/C_{\rm a},\tag{5}$$

where a = discrimination against ¹³C during diffusion (4.4%) and b = discrimination against ¹³C during the carboxylation reaction (27.0%). By rearranging Eq. (5) we were able to solve for C_i :

$$C_{\rm i} = C_{\rm a}(\Delta - a/b - a). \tag{6}$$

The intercellular [CO₂] is typically used in place of $C_{\rm C}$ in plant ecophysiological research because $C_{\rm C}$ cannot be estimated from gas-exchange measurements. Values of δ^{13} C reflect $C_{\rm C}$, not $C_{\rm i}$, and this analysis can lead to overestimates of $A/g_{\rm s-integrated}$. The degree to which $C_{\rm i}$ varies from $C_{\rm C}$ is dependent on $r_{\rm m}$, with higher values of $r_{\rm m}$ corresponding to greater overestimation of $A/g_{\rm s-integrated}$.

We used the three-parameter exponential equation $y = y_0 + a(1 - e^{-bx})$ describing the dependence of A on C_i obtained from gas-exchange measurements in the laboratory to estimate A from the C_i values obtained from foliar δ^{13} C. This methodology assumes that the non-stomatal, photosynthetic characteristics of the foliage are not substantially influenced by detachment. The value of $g_{s-integrated}$ is determined from estimates of $A/g_{s-integrated}$ and $A_{integrated}$. Stable isotope analyses were conducted on n = 3 branches per tree, and on three trees per height class.

Maximum leaf hydraulic conductance ($K_{\text{leaf-max}}$) was measured by methods adapted from Brodribb and Holbrook (2003) as described by Woodruff et al. (2008). The method involves use of Eq. (7) which is derived from the relationship between rehydrating a leaf and recharging a capacitor:

$$K_{\text{leaf}} = C \ln \left(\Psi_{\text{o}} \Psi_{\text{f}} \right) t, \tag{7}$$

where $K_{\text{leaf}} = \text{leaf}$ hydraulic conductance, C = capacitance, $\Psi_{\text{o}} = \text{leaf}$ water potential before rehydration, $\Psi_{\text{f}} = \text{leaf}$ water potential after rehydration, and t = duration of dehydration. We determined C on an individual tree basis from the slope of the relationship between relative water content and leaf water potential (Ψ_{l}) obtained from pressure-volume curves.

Results

Leaf structural characteristics

There were height-related trends in the examined leaf morphological characteristics (Figure 1). A 0.2 mm decline in needle length per meter increase in height was the single significant trend in needle morphology most (P < 0.0001). One-sided needle area showed the next most significant height-related trend, declining by 0.21 mm² for every meter increase in height (P = 0.0004). Needle width declined with increasing height by 5.9×10^{-3} mm m⁻¹ (P = 0.0025), and needle thickness increased with height by $8.9 \times 10^{-4} \text{ mm m}^{-1}$ (P = 0.0011). Stomatal density showed a significant linear decline with increasing height of -0.7 stomata mm⁻² m⁻¹ (P = 0.0032), but a threeparameter exponential decay function $(y = y_0 + ae^{-bx})$ yielded an improved regression fit with $r^2 = 0.77$ (data not shown) suggesting a possible limit to the reduction in



Figure 1. Leaf anatomic and structural characteristics in relation to tree height for foliage samples obtained within 1–5 m of the tops of *P. menziesii* trees at mean sampling heights of 5.0, 18.3, 34.5 and 55.0 m. Needle length (A), needle width (B), needle thickness (C), leaf area per needle (D), stomatal density (E) and mesophyll thickness (F). Bars denote \pm SE, n = 3 branches per tree for (A), (B), (C)s, (D) and (F); n = 4 branches for (E).

stomatal density with increased height. Mesophyll thickness showed a significant increase with increasing height of 0.74 μ m m⁻¹ (P = 0.0036).

Gas exchange

Gas-exchange measurements showed that, in the initial portion of the $A-C_i$ curves where A is heavily influenced by g_m , A declined with increasing height (Figure 2). Mean $(\pm SE)$ A_{amb} at 55 m was 5.49 ± 0.41 µmol m⁻² s⁻¹ and mean $A_{\rm amb}$ at 5 m was 9.52 \pm 0.42 μ mol m⁻² s⁻¹, representing a 42% reduction in assimilation at ambient [CO₂] over a 50-m increase in height (Figures 2 and 3). There were significant linear reductions in both A_{amb} and g_m with increasing height $(r^2 = 0.72, P = 0.00051; r^2 = 0.60, P = 0.0032,$ respectively, data not shown). A three-parameter exponential decay function $(y = y_0 + ae^{-bx})$ yielded improved regression fits with $r^2 = 0.83$ and 0.77 for A_{amb} and g_{m} plotted against height, respectively (Figure 3). Plotting Aamb against g_m revealed a more significant relationship than either variable plotted against height ($r^2 = 0.92$, P < 0.0001), consistent with g_m as an important determinant of A under non-saturating [CO₂] (Figure 3, inset). The correlation between A_{max} and height was also significant ($r^2 = 0.45$, P = 0.017; Table 1) although substantially less than the correlation between height and A_{amb} . Neither V_{cmax} nor J was significantly correlated with height $(r^2 = 0.04, P = 0.54; r^2 = 0.005, P = 0.83,$ respectively, Table 1). The correlation of nitrogen per unit leaf area $(N_{\rm A})$ with height was nearly significant $(r^2 = 0.32,$ P = 0.056; Table 1). Foliar δ_{13} C increased significantly with height by 0.042°_{00} m⁻¹ (*P* = 0.0002, data not shown). Integrated water-use efficiency increased significantly with height (P = 0.00019) by 0.45 µmol mol⁻¹ for every 1 m increase in height (Figure 4A). In contrast to $A/g_{s-integrated}$, there was a slight but significant (P = 0.011) decline in A/g. with increasing height (Figure 4B). Mesophyll thickness was negatively correlated with both K_{leaf-max} (P = 0.00008; Figure 5A) and A_{max} (P = 0.02; Figure 5B). Both A_{max} and $K_{\text{leaf-max}}$ were positively correlated with each other (P = 0.02, data not shown). Because $A_{\text{integrated}}$ was estimated from foliar δ^{13} C, it reflects the photosynthetic history of the foliage when the shoot was still attached to the tree. There was a linear decline in Aintegrated



Figure 2. CO_2 assimilation rate (*A*) versus intercellular CO_2 concentration (*A*-*C*_i curves) for foliage sampled within 1–5 m of the tops of *P. menziesii* trees at mean sampling heights of 5.0, 18.3, 34.5 and 55.0 m. The vertical dashed line represents 235 ppm, the mean value for *C*_i at standard ambient [CO₂] (385 ppm).

with increasing height ($r^2 = 0.89$, P < 0.0001), but the decline was steeper than that of A_{amb} with increasing height. A three-parameter exponential decay function yielded an improved regression fit with $r^2 = 0.99$ for the dependence of $A_{integrated}$ on height (Figure 6A). Mean integrated g_s ($g_{s-integrated}$) was also negatively correlated with height (P = 0.0002). A three-parameter exponential decay function yielded a regression fit with $r^2 = 0.98$ (Figure 6B).

Discussion

We were able to isolate gravity and path length as causal factors for trends in gas exchange by collecting our samples exclusively from the tops of Douglas-fir trees of different heights, within a localized geographic area, thereby eliminating appreciable differences in irradiance, humidity, temperature and branch length as confounding variables in the development of foliar characteristics that could influence gas exchange. These size-related trends in gas exchange of Douglas-fir can thus be attributed to intrinsic properties of the shoot that arise during tissue development.

Leaf structural characteristics

Leaf structural characteristics can strongly affect processes involved in gas exchange such as light absorption, carbon fixation and water loss. Needle length, width and area each showed significant negative trends with increasing tree height, supporting earlier observations of reduced leaf expansion along height gradients within the crowns of individual Douglas-fir trees (Woodruff et al. 2004, Meinzer et al. 2007). Although one might assume that smaller leaves have higher stomatal density because of the tighter packing of stomata among the less expanded epidermal cells as has

often been reported in sun versus shade leaves (Givnish 1988, Osborn and Taylor 1990), we found a 19% reduction in mean stomatal density from the lowest to the tallest height class studied. Previous work has shown that cell division as well as expansion are sensitive to turgor pressure (Boyer 1968, Kirkham et al. 1972, Hsiao et al. 1976, Gould and Measures 1977), which could have implications for the mechanisms that control the differentiation of epidermal cells to stomata. Beerling and Chaloner (1993) found a reduction in stomatal density in Quercus with increased temperature, and suggested that the observed response was a possible adaptation to reduce water loss. Although reduced stomatal density will tend to limit capacity for A, particularly in leaves of higher density in which lateral movement of CO₂ through internal air spaces is limited (Parkhurst 1986), its primary consequence may be to reduce water loss, particularly in cases in which stomata do not completely close (Caird et al. 2007).

Hydraulic resistance

The leaf mesophyll also presents a site of resistance to the transport of water from xylem conduits to the stomata. Water moves out of the leaf xylem and then through bundle sheath and mesophyll cells before evaporating in the intercellular air spaces. The resistance to water flow through living mesophyll tissue is substantially greater than through nonliving vessels and tracheids (Boyer et al. 1985), and thus mesophyll hydraulic resistance may represent a substantial limitation for gas exchange. Aasamaa et al. (2001) had found that the area of mesophyll and epidermal cells per unit length of leaf cross section was strongly and positively correlated with shoot hydraulic conductance across a range of deciduous trees. Brodribb et al. (2007) had found that across 43 species, A_{max} was correlated with the distance



Figure 3. CO₂ assimilation rate at an intercellular CO₂ concentration of 235 ppm (A_{amb}) versus height (A) and mesophyll conductance (g_m) versus height (B) in *P. menziesii*. Inset shows A_{amb} versus g_m . Bars denote \pm SE, n = 2 branches.

between veins and leaf surface and with K_{leaf} . They concluded that the relationship between vein location and photosynthetic rate was determined by the conductance of the leaf mesophyll to water flow. In our study, the highly significant correlation between T_{m} and $K_{\text{leaf-max}}$ (Figure 5A) suggests that leaf structural characteristics that influence r_{m} can have a substantial influence on leaf hydraulic architecture, and subsequently photosynthesis.

Resistance to transfer of CO₂

We found that the ratio of g_m from detached shoots to $g_{s-integrated}$ estimated from foliar $\delta_{13}C$ ranged from 0.47 to 0.71. Peña-Rojas et al. (2005) had found g_m in *Quercus* to be equal to about one-half of g_s . DeLucia et al. (2003)

had found g_m/g_s to be between 0.6 and 1.1 in conifers and between 1.0 and 4.2 in angiosperms, and in a survey of 50 species Warren (2008) had found g_m/g_s to be 1.3 in conifers and between 0.52 and 1.5 in angiosperms. We found a 47% reduction in mean g_m along a 50-m height gradient (Figure 3B), which was strongly correlated with variation in A_{amb} among height classes (Figure 3, inset). Mesophyll conductance is considered to be a key determinant of A at low values of C_i and thus the initial slope of $A-C_i$ curves (Farquhar et al. 1980). Consistent with this relationship, the most pronounced differences in A between height classes were observed under non-saturated [CO₂] (A_{amb} ; Figures 2 and 3), whereas the trend in A_{max} with increasing height was much less pronounced (Table 1). Given the importance of g_m as a determinant of A under



non-saturating [CO₂], the greater height-related decline in A_{amb} relative to that of A_{max} provides further evidence of g_{m} as a critical factor in the observed decline in A_{amb} with increasing height in Douglas-fir.

Water stress has been shown to lead to reductions in g_m , even in cases where the water stress is only moderate (O'Toole et al. 1976, Loreto et al. 1997, Peña-Rojas et al. 2005). A proposed mechanism for the water-stress-induced depression of g_m is through reduced leaf turgor and its influence on mesophyll surface area. The height-related decline in g_m in our study is thus likely to be associated with the effects of gravitational and transpirational xylem tension gradients on Ψ_1 (Bauerle et al. 1999) and turgor during leaf expansion (Woodruff et al. 2004). Warren et al. (2004) had used simultaneous gas-exchange and isotopic measurements to determine g_m for Douglas-fir seedlings under different hydration regimes and found 73% higher g_m in well-watered Douglas-fir seedlings than in water-stressed

Figure 4. Integrated water-use efficiency $(A/g_{s-integrated})$ versus height (A) and instantaneous water-use efficiency (A/g_s) versus height (B) in *P. menziesii*. Bars denote \pm SE, n = 3 branches for (A) and n = 2 branches for (B).

seedlings (0.076 and 0.044 mol m⁻² s⁻¹, respectively). In the first study to report g_m in a conifer, Warren et al. (2003) had found that the mean g_m was 0.16 mol m⁻² s⁻¹ in the lower canopy (17–20 whorls down from the top) and 0.20 mol m⁻² s⁻¹ in the upper canopy (4–8 whorls down from the top) within the same 34-m-tall Douglas-fir tree. Their observed height-related increase in g_m suggests that light exposure within a tree crown may mask intrinsic height-related trends in g_m .

Nitrogen- and water-use efficiency

Nitrogen concentration per unit leaf area (N_A) was 17% lower at 55 than at 5 m, and the correlation between height and N_A was nearly significant (P = 0.056). The correlation between A_{max} and N_A , however, was more significant (P = 0.043). The correlations of N_A with both A_{max} and A_{amb} (P = 0.052); and g_{m} with both A_{max} (P = 0.3) and



 $A_{\rm amb}$ (P < 0.0001) highlight their respective roles at different CO₂ concentrations, and in particular the influence of $g_{\rm m}$ on A at $C_{\rm i}$ values that correspond to natural ambient [CO₂]. Because our study involved sampling at four distinct stands, there exists the possibility that stand history and factors such as the former or current presence or absence of nitrogen-fixing plants such as Alnus spp. may confound analyses of nitrogen variation with height. Nevertheless, the lack of a more significant height-related decline in N_A , despite the highly significant height-related decline in $A_{\rm amb}$ (P = 0.00051) suggests a reduction in photosynthetic N-use efficiency with increasing height. Given that thicker needles are less likely to be able to maximize N use on an area basis, the observed height-related trend in N-use efficiency may to some extent be related to the observed height-related patterns in needle anatomy.

The contrast between the strong increase in $A/g_{s-integrated}$ and the slight decrease in A/g_s with increasing height (Figure 4) highlights the role of stomatal control of water loss and the resulting diffusional limitations on photosynthesis in attached shoots. The significant trend in $A/g_{s-integrated}$ with height (Figure 4A) is consistent with height-related trends in foliar δ^{13} C or isotope discrimination (Δ) obtained by others (Yoder et al. 1994, McDowell et al. 2002, Koch et al. 2004, McDowell et al. 2005). This height-related increase in intrinsic water-use efficiency is a key component

Figure 5. Maximum leaf hydraulic conductance ($K_{\text{leaf-max}}$) versus mesophyll thickness (T_{m}) (A) and maximum photosynthetic rate (A_{max}) versus T_{m} (B) in *P. menziesii*. Horizontal bars denote \pm SE, n = 3 branches for T_{m} (A and B). Vertical bars denote \pm SE, n = 2-8branches for $K_{\text{leaf-max}}$ (A), and n = 2branches for A_{max} (B). Three values in (A) represent trees with only one branch with a $K_{\text{leaf-max}}$ value and hence lack vertical bars.

of the hydraulic limitation hypothesis (Ryan and Yoder 1997). The lack of a height-related increase in A/g_s for shoots that were sampled along a height gradient, but which had their cut bases in water during the photosynthetic measurements, provides further evidence that path length and gravity are responsible for the height-related trend in $A/g_{s-integrated}$. The slight decline in A/g_s with increasing height for the cut shoots may reflect a heightrelated trend in concentrations of osmotic solutes in stomatal guard cells and other leaf cells. Osmotic adjustment that partially maintains leaf cell turgor along a height gradient has been observed in conifers (Koch et al. 2004, Woodruff et al. 2004). Stomatal opening is activated and maintained by a solute-mediated influx of water into the guard cells. Thus, when the impacts of gravity and path length on xylem tension are removed in detached shoots, they may exhibit greater g_s as a result of the height-related gradient in osmotic concentration.

Estimates of height-related variation of A in situ ($A_{integrated}$; Figure 6A) obtained from foliar δ^{13} C values and $A-C_i$ curves followed a pattern similar to that of A_{amb} (Figure 3), although $A_{integrated}$ declined more steeply than A_{amb} with increasing height. The reductions in mean $A_{integrated}$ and A_{amb} from the 5 to 55 m sampling heights were 5.64 and 4.03 µmol m⁻² s⁻¹, respectively. Integrated CO₂ assimilation incorporates the stomatal component of



Figure 6. Integrated CO₂ assimilation $(A_{\text{integrated}})$ versus height (A) and integrated stomatal conductance $(g_{s-\text{integrated}})$ versus height (B) in *P. menziesii*. Bars denote \pm SE, n = 3 branches.

limitation to A because it represents an estimate of integrated, in situ A, whereas A_{amb} excludes the stomatal component because it is derived from the dependence of A on substomatal [CO₂]. Mean integrated values of g_s ($g_{s-integrated}$) from the 5 to 55 m sampling heights declined by 2.5 mmol m⁻² s⁻¹ for every 1 m increase in height (Figure 6B). Although absolute values may not be directly comparable because of the uncertainties associated with estimating stomatal conductance from foliar δ^{13} C, the height-related trends in g_m and $g_{s-integrated}$ suggest that g_s decreases more rapidly than g_m with increasing tree height.

Our data support the hypothesis that height-related trends in leaf structure impose constraints on leaf gas exchange that are independent of the direct effects of vertical gradients in xylem tension on stomatal and photosynthetic physiology. Nevertheless, gravity and path length, through their effects on xylem tension during leaf expansion, were likely to have been the principle determinants of the trends in foliar characteristics that were correlated with the observed height-related trends in gas exchange.

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