Canopy gradients in leaf functional traits for species that differ in growth strategies and shade tolerance

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In temperate deciduous forests, vertical gradients in leaf mass per area (LMA) and area-based leaf nitrogen (Narea) are strongly controlled by gradients in light availability. While there is evidence that hydrostatic constraints on leaf development may diminish LMA and Narea responses to light, inherent differences among tree species may also influence leaf developmental and morphological response to light. We investigated vertical gradients in LMA, Narea and leaf carbon isotope composition (δ¹³C) for three temperate deciduous species (Carpinus caroliniana Walter, Fagus grandifolia Ehrh., Liriodendron tulipifera L.) that differed in growth strategy (e.g., indeterminate and determinate growth), shade tolerance and leaf area to sapwood ratio (Al:As). Leaves were sampled across a broad range of light conditions within three vertical layers of tree crowns to maximize variation in light availability at each height and to minimize collinearity between light and height. All species displayed similar responses to light with respect to Narea and δ¹³C, but not for LMA. Light was more important for gradients in LMA for the shade-tolerant (C. caroliniana) and intol-erant (L. tulipifera) species with indeterminate growth, and height (e.g., hydrostatic gradients) and light were equally important for the shade-tolerant (F. grandifolia) species with determinate growth. Fagus grandifolia had a higher morphological plasticity in response to light, which may offer a competitive advantage in occupying a broader range of light conditions throughout the canopy. Differences in responses to light and height for the taller tree species, L. tulipifera and F. grandifolia, may be attributed to differences in growth strategy or Al:As, which may alter morphological and functional responses to light availability. While height was important in F. grandifolia, height was no more robust in predicting LMA than light in any of the species, confirming the strong role of light availability in determining LMA for temperate deciduous species.

Keywords: Carpinus caroliniana, Fagus grandifolia, leaf carbon isotope composition, leaf mass per area, leaf nitrogen, Liriodendron tulipifera.

Introduction

Within temperate deciduous forests, gradients in light availability are often implicated as a driving factor for two coordinated leaf traits, leaf mass per area (LMA) and leaf nitrogen per area (Narea) (Ellsworth and Reich 1993, Niinemets et al. 1998, 1999, Meir et al. 2002, Jones and Thomas 2007, Montpied et al. 2009, Coble and Cavalieri 2014, 2015, Scartazza et al. 2016). In temperate deciduous species, leaves growing in high light conditions with greater LMA often contain a thicker palisade layer that enhances light penetration to deeper layers of the mesophyll (Jackson 1967, Vogelmann and Martin 1993, Evans 1999, Oguchi et al. 2005) and with greater Narea partition more nitrogen into ribulose bisphosphate carboxylase (RuBisCO) leading to greater photosynthetic capacity (Evans 1989, Hikosaka and Terashima 1995). Due to the strong association among LMA, Narea and photosynthesis (Ellsworth and Reich 1993, Reich et al. 1999, Wilson et al. 2000, Kitaoka et al. 2002, Wright et al. 2004, Montpied et al. 2009), these easy-to-measure leaf traits are necessary for modeling canopy photosynthesis (Aber et al. 1996, Raulier et al. 1999,

Height has also been implicated as a strong driver of vertical gradients in LMA in very tall western conifer and tropical trees, and there is growing evidence that height may also influence LMA in shorter deciduous trees. Due to gravity alone, leaf water potential linearly decreases with height (−0.01 MPa m−1), in both conifer and temperate deciduous species (Scholander et al. 1965, Hellkvist et al. 1974, Bauerle et al. 1999, Woodruff et al. 2004, Coble and Cavaleri 2014). In western conifer trees such as the very tall Sequoia sempervirens (~97–113 m) (Koch et al. 2004, Ishii et al. 2008, Mullin et al. 2009, Oldham et al. 2010) and the shorter Pinus monticola (45 m), Pinus ponderosa (41 m) and Pseudotsuga menziesii (40 m) (Marshall and Monserud 2003) the gravitational potential gradient appears to limit leaf development at greater heights. Limited osmotic adjustments and leaf turgor at greater heights may restrict leaf expansion, resulting in leaves with high LMA (Woodruff et al. 2004, Meinzer et al. 2008). Vertical gradients in leaf carbon isotope composition (δ13C) in tall S. sempervirens trees provide further evidence for increasing water stress with height and more frequent stomatal closure of upper canopy leaves (Koch et al. 2004, Ishii et al. 2008). In tall tropical trees (~40 m), LMA was strongly and linearly correlated with height throughout the entire canopy, whereas light appeared to influence LMA only in lower light conditions (Cavaleri et al. 2010). In shorter temperate deciduous trees (~18 m), Zhang et al. (2011, 2012a, 2012b) found that midday leaf water potential values in Robinia pseudoacacia seedlings that corresponded with in situ measurements were sufficient to increase LMA through constraints on leaf expansion via reduced cell wall extensibility. Additional evidence for early season height effects on leaf development and LMA were observed during the leaf expansion phase in Acer saccharum (~21 m; Coble et al. 2016) and Quercus rubra trees (~18 m; Zwieniecki et al. 2004). A potential consequence of morphological adjustments to height is greater investment in carbon-rich compounds (e.g., lignin, cellulose), which improves mechanical strength and tolerance of lower water potential, while simultaneously diluting leaf nitrogen, which leads to suboptimal distributions in leaf nitrogen (Niinemets et al. 1999, Coble and Cavaleri 2015).

Inherent differences among temperate deciduous species, such as leaf growth strategies, could potentially influence leaf development and their ability to acclimate to the local light environment (Ishii and Ohsugi 2011). Woody plant species can be broadly classified as having indeterminate or determinate growth (Kozlowski 1964, Marks 1975). For indeterminate species, leaves and shoots are formed continuously during the season of growth, whereas leaves of many determinate species are initially formed in buds as leaf primordia in the year prior to leaf growth and experience one flush of leaves at the start of the growing season. Plants with indeterminate growth patterns are capable of acclimating to light through successive leaf emergence, which may contribute their greater responsiveness to changes in light availability (Naidu and DeLucia 1998, Ishii and Ohsugi 2011). For example, the indeterminate-growth species, Liriodendron tulipifera L., had a greater ability in morphological and photosynthetic acclimation to changing light availability compared with the determinate-growth and shade-tolerant species, A. saccharum (Naidu and DeLucia 1998). An advantage to indeterminate growth is that species can capitalize on short-term changes in light availability through adjustments in leaf morphology and function in newly formed leaves. In contrast, species with determinate growth patterns are somewhat constrained in acclimation because leaf anatomical structure is predetermined in the year prior to leaf flush and tends to show ‘carry-over’ effects from previous-year light conditions (Naidu and DeLucia 1998, Jones and Thomas 2007, Ishii and Ohsugi 2011). For example, Fagus sylvatica buds growing in full sunlight were found to contain leaf primordia with more cell layers compared with shaded buds, leading to two-layers of palisade cells for sun leaves and one-layer of palisade cells for shade leaves (Eschrich et al. 1989). These growth strategies (indeterminate and determinate) may potentially explain why tall, mature trees of different species may differ in their response to light availability within forest canopies.

The objective of this study was to identify any evidence for height constraints on leaf development for three tree species (Carpinus caroliniana Walter, L. tulipifera, Fagus grandifolia Ehrh.) that differ in growth strategies and shade tolerance. We used a sampling design that maximized the range of light availabilities within each vertical canopy layer to minimize collinearity between light and height when assessing their influence on leaf functional traits (LMA, Narea and δ13C). In this study, we also examine differences in leaf area to sapwood area ratio (A:L) because there is evidence that A:L plays in important role in whole-tree water relations, and reductions in A:L may be a mechanism for tolerating greater declines in leaf water potential and hydraulic constraints (McDowell et al. 2002). We tested the following hypotheses: (i) light is more important than height for determining LMA in temperate deciduous species; and (ii) LMA, Narea and δ13C are strongly coordinated in all species due to the coordination between leaf morphology and leaf function.

Materials and methods

Study site

The study was conducted in a mixed deciduous forest at the Smithsonian Environmental Research Center (SERC) in Edgewater, MD (38° 53′ N, 76° 33′ W). Mean local annual temperature and precipitation is 12.8 ± 0.7 °C and 1090 ± 172 mm, respectively (NOAA regional climatology 1895–2010). The stand is considered a ‘tulip poplar’ association (Brush et al. 1980) and originated around 1887 after a large pasture was abandoned. The diverse forest is dominated by L. tulipifera, which makes up
49% of total basal area (BA). In order of importance (sum of % density and % BA), other species include C. caroliniana, F. grandifolia, Cornus florida, Carya tomentosa, Liquidambar styraciflua, Quercus alba, Carya glabra, Quercus velutina and other species.

The target species for this study were C. caroliniana, L. tulipifera, and F. grandifolia. Carpinus caroliniana has indeterminate growth and is shade-tolerant. L. tulipifera has indeterminate growth and is shade-intolerant (Nobey and O’Neill 1991, Naidu and DeLucia 2015) and F. grandifolia has determinate growth and is shade-tolerant (Marks 1975, Goulet and Bellefeu 1985) (Table 1). Vertical stratification of foliage and phenology for the three species in a stand next to this study site is described in Parker et al. (1989) and Parker and Tibbs (2004). Briefly, C. caroliniana is primarily restricted to the lower canopy (<10 m), L. tulipifera occupies the mid- (10–25 m) and upper canopies (>25 m) and F. grandifolia occupies the mid- to lower canopies with a few large trees [diameter at breast height (dbh) >40 cm] that occupy portions of the upper canopy. The mean (± standard error) height of all C. caroliniana, L. tulipifera and F. grandifolia trees greater than 2.5 cm dbh in a 0.5 ha plot in the same stand in 1997 were 10.5 ± 0.01, 31.2 ± 0.03 and 25.1 ± 0.02 m, respectively (unpublished data).

**Study design**

Three trees per species were selected, and the crown of each tree was divided into three vertical layers based on crown height. Each layer was divided into five horizontal sections based on horizontal distance from the center of the crown. The division of crown layers into five horizontal sections was made in order to maximize the range of variation in light availability at each sample height for each of three species. The mean (± standard deviation) height of sample trees for C. caroliniana, L. tulipifera and F. grandifolia were 7.7 ± 2.2, 36.4 ± 1.8 and 24.5 ± 2.4 m, respectively (Table 1). Light and height measurements and leaf collections were conducted in August 1998 using a personnel hoist suspended from a mobile hydraulic crane.

**Light and height measurements**

At each leaf sampling location, hemispherical photographs were taken using a digital imaging system with a gimbaled 180° lens (CI-110 system with LLP Lens, CID, Vancouver, WA, USA). Images were taken in early morning or late evening. Digital hemispherical images were analyzed using Hemiphoto software to calculate direct site factor (DSF) (proportion of direct light relative to open conditions), indirect site factor (ISF) (relative illumination of diffuse light under a standard overcast sky (SOC) and canopy openness. All of these measures were highly correlated. Here we use the total site factor (TSF), a weighted average of ISF and DSF.

In the summer of 1994 in the same stand, we also measured the photosynthetic photon flux density (PPFD, 400–700 nm) along 29 vertical transects using a cosine-corrected quantum sensor (Model LI-190SB, Li-Cor, Lincoln, NE, USA). The sensor was attached to the top of a balloon and measurements were made at every meter up to 40 m in height. Photosynthetic photon flux density was also measured above the canopy during each transect to serve as the reference. Parker et al. (1996, 2002) provide a full description and application, respectively, of the balloon technique. Light transmittance values were made throughout the canopy, regardless of leaf locations, whereas measurements of TSF were made at the location of leaf samples. While the light transmittance measurements occurred 4 years prior to this study, these data are used to assess the extreme light conditions (5th and 95th percentile) of light transmittance in reference to the TSF values at each height. Furthermore, a tree census of a 0.5 ha plot that began in 1987 (Parker et al. 1989) and continued annually thereafter within the same stand indicated that the stand grew very little in height during the 1994–98 interval. Mean tree height in 1993 for C. caroliniana, L. tulipifera and F. grandifolia was 10.5, 31.2 and 25.1 m, respectively, which changed less than 0.1 m over 4 years. Height at each location and total tree height was measured by dropping a tape measure from the foliage sample to the ground.

**Leaf area to sapwood area ratio (A_L:A_S)**

Leaf area to sapwood area ratio (A_L:A_S, m² cm⁻²) was estimated for L. tulipifera and F. grandifolia by first calculating stem diameter at 1.37 m above the ground (dbh) using total tree height from this study and dbh–height allometric relationships reported by Anderson-Teixeira et al. (2015) for the same species at a site ~170 km from the current study location. Diameter at breast height and dbh–leaf area allometric relationships were used to estimate whole-tree leaf area. We estimated the allometric relationship between leaf area and dbh with a non-linear procedure (SAS proc NLIN) based on dataset of concurrent collections of stem diameter and litterfall from an age-series of local plots. Both stem and leaf were identified to the same species classes and leaf masses were converted to leaf area following the procedure described in Parker and Tibbs (2004). The plot total leaf

Table 1. Shade tolerance, growth strategy, mean tree height (m; ± standard deviation) and predicted mean leaf area to sapwood area ratio (m² cm⁻²) of sample trees. Different letters indicate significant differences between species (P < 0.05, Tukey’s HSD).

<table>
<thead>
<tr>
<th>Species</th>
<th>Shade tolerance</th>
<th>Growth strategy</th>
<th>Tree height (m)</th>
<th>Predicted A_L:A_S (m² cm⁻²)</th>
</tr>
</thead>
<tbody>
<tr>
<td>C. caroliniana</td>
<td>Tolerant</td>
<td>Indeterminate</td>
<td>7.7 ± 2.2</td>
<td>-</td>
</tr>
<tr>
<td>L. tulipifera</td>
<td>Intolerant</td>
<td>Indeterminate</td>
<td>36.4 ± 1.8</td>
<td>0.295 ± 0.006</td>
</tr>
<tr>
<td>F. grandifolia</td>
<td>Tolerant</td>
<td>Determinate</td>
<td>24.5 ± 2.4</td>
<td>0.339 ± 0.001</td>
</tr>
</tbody>
</table>
area and stem size distributions were used as constraints for fitting the allometric constants for each species. Sapwood area was estimated from dbh using dbh–sapwood area allometric relationships reported by Anderson-Teixeira et al. (2015). Given that sapwood area can change across soil types (Long and Smith 1988), we also estimated the leaf area to basal area ratio \( (m^2 \cdot m^{-2}) \) for these species growing within the same stand in order to validate our results. Across a number of temperate deciduous species, basal area and dbh are both strongly correlated with sapwood area (Gebauer et al. 2008). We estimated the leaf area to basal area ratio using stand basal area \( (m^2 \cdot m^{-2}) \) and LAI \( (m^2 \cdot m^{-2}) \) reported by Parker and Tibbs (2004). Stand-level leaf area per unit basal area was greater for C. caroliniana (5164 m\(^2\) m\(^{-2}\)) and F. grandifolia (3694 m\(^2\) m\(^{-2}\)) compared with L. tulipifera (487 m\(^2\) m\(^{-2}\)).

**Leaf mass per area, leaf nitrogen and carbon isotope composition**

At each location, we collected 10 leaves from C. caroliniana and 4–6 leaves from L. tulipifera and F. grandifolia at the ends of branch tips. Leaf collection of F. grandifolia and C. caroliniana (determinate growth) were from the initial flush of leaves in the early spring, whereas leaf collection of L. tulipifera (indeterminate growth) included a mix of older (e.g., from spring flush) and newer leaves. Leaves were temporarily stored in sealed plastic bags and refrigerated. Leaf area was measured using a benchtop leaf area meter (LI-3100 Area Meter, Li-Cor) to the nearest 0.1 cm\(^2\). Leaves were then dried at 60°C for 4 days and weighed to the nearest 0.1 mg. Leaf samples used to estimate leaf nitrogen and carbon isotope composition were processed at the SERC. Bulk leaf samples were ground to a fine powder. Leaf nitrogen on a mass basis \( (N_{mass}, \text{mg} \cdot g^{-1}) \) was converted to area-based nitrogen \( (N_{area}, \text{g} \cdot m^{-2}) \) using LMA \( (N_{area} = \text{LMA} \times N_{mass})/1000) \). Leaf δ\(^{13}\)C was determined using a Carlo Erba Instruments NC 2500 elemental analyzer coupled to a Thermo Scientific Delta V Plus IRMS. Leaf δ\(^{13}\)C was calculated as \( \frac{13C}{12C} = 1000 \times \left( R_{sample}/R_{standard} - 1 \right) \times 10^{-3} \), where \( R_{sample} \) is the \( ^{13}C/^{12}C \) ratio for the sample and \( R_{standard} \) is the ratio for a standard. We note that the leaf carbon isotope data (δ\(^{13}\)C) in this study was used in a previous publication, Graham et al. (2014). However, the primary focus of the data presented by Graham et al. (2014) was to assess canopy openness of ancient trees from leaf fossil isotopes, and the results in our study are displayed in a different context.

**Data analysis**

All statistical analyses were conducted using R software (R Development Core Team 2015). Regression analysis was used to identify the significance and strength of the relationship between height and TSF for each species. Total site factor was natural log-transformed because TSF displayed curved, asymptotic relationships with height and log-transformation of TSF satisfied regression assumptions. To address collinearity between TSF and height for each species, we calculated the variance inflation factor (VIF) between TSF and height (Ott and Longnecker 2001). The VIF quantifies the extent of inflation of the coefficient variance due to collinearity. A VIF of 1 indicates no collinearity, whereas a VIF of 10 indicates that collinearity is a problem. We used three approaches to separate the effects of TSF and height on LMA. First, we examined the residuals of LMA vs height plotted against TSF and the residuals of LMA vs TSF plotted against height, and compared their \( R^2 \) values. Second, we used pairwise partial correlations to separate the effects of TSF and height on LMA. The pairwise partial correlations allowed us to control for light and height when comparing LMA–height and LMA–TSF correlations, respectively. Third, we used partial \( R^2 \) analysis to compare the contribution of TSF and height to the full model \( (y = \beta_0 + \beta_1 \text{TSF} + \beta_2 \text{height}) \) for predicting LMA.

We used ANCOVA for species and covariate (height and TSF) effects on LMA, \( N_{area} \), and δ\(^{13}\)C, as well as the interaction effects (height × species, TSF × species). An initial analysis that included individual trees as a third factor in the ANCOVA model revealed a non-significant tree effect on LMA and \( N_{area} \). Thus, we pooled all data from the three study trees within species. We assessed plasticity for LMA and \( N_{area} \) by comparing slopes of the LMA–TSF, LMA–height, \( N_{area} \)-TSF and \( N_{area} \)-height linear regression models, as well as slopes of the LMA–\( N_{area} \), LMA–δ\(^{13}\)C and \( N_{area} \)-δ\(^{13}\)C models, among the three species. We tested for differences in slopes by first fitting the ANCOVA model (e.g., \( \text{LMA} = \beta_0 + \beta_1 \text{TSF} + \beta_2 \text{Species} + \beta_1x \text{TSF} \times \text{Species} \)) and assessing significance \( (P < 0.05) \) of the differences in slopes between the ‘reference’ species and the two remaining species in the summary output in R software. We repeated this analysis after changing the ‘reference’ species so that tests for differences in slopes could be made for all species. R software uses \( t \)-statistics to test whether slopes differ between individual regression models. We found that this approach generates identical results as those observed from the use of dummy variables in multiple regression for comparing slopes (Ott and Longnecker 2001).

**Results**

**Tree characteristics and light environment**

Liriodendron tulipifera had a smaller \( A_{II} \) compared with F. grandifolia (Table 1), which is consistent with our findings that L. tulipifera had a smaller stand-level leaf area to basal area ratio than F. grandifolia. Total site factor (TSF) as estimated with hemispherical photography was compared with a more extensive dataset of light transmittance values at the 5th and 95th percentile collected within the same stand (Figure 1). Our results confirm that leaf collections were made from a broad range of light conditions within each height for this stand. Total site factor was weakly correlated with height for all species.
where the $R^2$ values ranged from 0.20 to 0.29 (Table 2, Figure 1). The range of the VIF between height and (ln)TSF among all species was 1.25–1.41, which indicates that collinearity is not an issue in this study because the values were close to 1.

**Light versus height effects on LMA**

We found that TSF was more strongly correlated with residuals of the LMA vs height relationships for *C. caroliniana* ($R^2 = 0.22$) and *L. tulipifera* ($R^2 = 0.37$; Figure 2a) than between height and residuals of the LMA vs TSF relationships ($R^2 = 0.07$; Figure 2b). For *F. grandifolia*, the strength of the relationship between TSF and residuals of LMA vs height relationship ($R^2 = 0.55$; Figure 2a) was similar to the relationship between height and residuals of the LMA vs TSF relationship ($R^2 = 0.52$; Figure 2b). The partial $R^2$ analysis revealed that TSF contributed more to the full model (LMA = $\beta_0 + \beta_1 \text{TSF} + \beta_2 \text{Height}$) for *C. caroliniana* and *L. tulipifera*, while the contribution of height and TSF were similar for *F. grandifolia* (Table 3). Consistent with these findings, the pairwise partial correlation analysis showed that TSF was more strongly correlated with LMA when controlling for height for *C. caroliniana* and *L. tulipifera* compared with the correlation between LMA and height when controlling for TSF (Table 4). For *F. grandifolia*, the partial correlation also showed that the strengths of the LMA–TSF and LMA–height correlations were similar (Table 4).

**Species responses to light and height**

Among the three species, we observed differences in LMA responses to TSF and height, but all three species displayed similar changes in $N_{\text{area}}$ with increasing TSF and height. The ANCOVA indicated significant TSF ($P < 0.001$), species ($P < 0.001$) and TSF × species ($P = 0.003$) effects on LMA. Similarly, height ($P < 0.001$), species ($P < 0.001$) and height × species ($P = 0.002$) had significant effects on LMA. Leaf mass per area was linearly and positively correlated with TSF and height (Figure 3a and b). The slope of the LMA–TSF relationship for *F. grandifolia* was significantly greater than the slopes for *L. tulipifera* ($P = 0.004$) and *C. caroliniana* ($P = 0.017$), and no difference in slopes were detected between *L. tulipifera* and *C. caroliniana* ($P = 0.442$). The slope of the LMA–height relationship for *F. grandifolia* was significantly greater than the slope for *L. tulipifera* ($P < 0.001$), but not for *C. caroliniana* ($P = 0.264$).

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**Table 2. Model parameters for predicting height using total site factor (TSF) including the intercept, slopes, model $R^2$ values and variance inflation factor (VIF) for each species. Model: Height = $\beta_0 + \beta_1 (\ln)\text{TSF}$.**

<table>
<thead>
<tr>
<th>Species</th>
<th>$\beta_0$</th>
<th>$\beta_1$</th>
<th>Model $P$-value</th>
<th>$R^2$</th>
<th>VIF</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>C. caroliniana</em></td>
<td>-2.78</td>
<td>2.35</td>
<td>$P &lt; 0.01$</td>
<td>0.26</td>
<td>1.35</td>
</tr>
<tr>
<td><em>L. tulipifera</em></td>
<td>-3.27</td>
<td>7.53</td>
<td>$P &lt; 0.01$</td>
<td>0.29</td>
<td>1.41</td>
</tr>
<tr>
<td><em>F. grandifolia</em></td>
<td>-7.23</td>
<td>6.05</td>
<td>$P &lt; 0.05$</td>
<td>0.20</td>
<td>1.25</td>
</tr>
</tbody>
</table>

**Figure 1.** Relationships between height and light (total site factor, %) for *C. caroliniana*, *L. tulipifera* and *F. grandifolia*. The dashed and solid lines represent the 5th and 95th percentile, respectively, of light transmittance (%) values at 1 m height increments.

**Figure 2.** Residuals of LMA vs height (m) plotted against TSF (a) and residuals of LMA vs TSF plotted against height (b) for *C. caroliniana* (dotted lines), *L. tulipifera* (dashed lines) and *F. grandifolia* (solid lines). $R^2$ values: (a) *C. caroliniana*, 0.22; *L. tulipifera*, 0.37; *F. grandifolia*, 0.55; and (b) *C. caroliniana*, 0.07; *L. tulipifera*, 0.07; *F. grandifolia*, 0.52.

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Vertical leaf traits in a temperate deciduous forest

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Table 3. For each species, coefficients and $R^2$ values are displayed for regressions between light (TSF), height and LMA and for multiple regressions between light + height and LMA. Partial $R^2$ values are shown for adding light or height to the full model and provide an indication of the relative importance of each parameter (height or light) for predicting LMA.

<table>
<thead>
<tr>
<th>Response variable</th>
<th>Species</th>
<th>N</th>
<th>Light (TSF)</th>
<th>Height</th>
<th>Light and height</th>
<th>Partial $R^2$ for adding:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>$\beta_0$</td>
<td>$\beta_1$</td>
<td>$R^2$</td>
<td>$\beta_0$</td>
</tr>
<tr>
<td>LMA</td>
<td>C. caroliniana</td>
<td>27</td>
<td>23.40***</td>
<td>0.31***</td>
<td>0.48</td>
<td>25.30***</td>
</tr>
<tr>
<td></td>
<td>L. tulipifera</td>
<td>27</td>
<td>46.28***</td>
<td>0.46***</td>
<td>0.67</td>
<td>37.98***</td>
</tr>
<tr>
<td></td>
<td>F. grandifolia</td>
<td>25</td>
<td>24.81***</td>
<td>0.77***</td>
<td>0.71</td>
<td>18.77**</td>
</tr>
</tbody>
</table>

**$P < 0.01$, ***$P < 0.001$.**

Table 4. Pairwise partial correlations for LMA (response variable) and TSF while controlling for height, and partial correlations for LMA and height controlling for TSF.

<table>
<thead>
<tr>
<th>Species</th>
<th>Response variable</th>
<th>Independent variable</th>
<th>Controlled variable</th>
<th>N</th>
<th>Coefficient</th>
<th>$P$-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>C. caroliniana</td>
<td>LMA</td>
<td>TSF</td>
<td>Height</td>
<td>27</td>
<td>0.557</td>
<td>0.001</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>TSF</td>
<td>27</td>
<td>0.308</td>
<td>0.113</td>
</tr>
<tr>
<td>L. tulipifera</td>
<td>LMA</td>
<td>TSF</td>
<td>Height</td>
<td>27</td>
<td>0.734</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>TSF</td>
<td>27</td>
<td>0.309</td>
<td>0.112</td>
</tr>
<tr>
<td>F. grandifolia</td>
<td>LMA</td>
<td>TSF</td>
<td>Height</td>
<td>25</td>
<td>0.881</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>TSF</td>
<td>25</td>
<td>0.888</td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>

Fagus grandifolia also had a greater range in LMA than the other species (Figure 3a and b).

For $N_{area}$, we observed significant height ($P < 0.001$), species ($P = 0.037$) and TSF ($P < 0.001$) effects, and the interaction height x species ($P = 0.383$) and TSF x species ($P = 0.746$) had no effect on $N_{area}$. $N_{area}$ was linearly and positively correlated with TSF and height (Figure 3c and d). There were no significant differences in the $N_{area}$-TSF and $N_{area}$-height slopes among the three species ($P > 0.05$).

For $\delta^{13}$C, we observed a significant height ($P < 0.001$), species ($P = 0.005$) and TSF ($P < 0.001$) effect, and the interaction height x species ($P = 0.061$) and TSF x species ($P = 0.750$) had no effect on $\delta^{13}$C. $\delta^{13}$C was linearly and positively correlated with TSF and height (Figure 3e and f). There were no significant differences in the $\delta^{13}$C-TSF slopes among the three species ($P > 0.05$). Despite the non-significant height x species effect on $\delta^{13}$C, the slope of the $\delta^{13}$C-height relationship for C. caroliniana was greater than the slopes for L. tulipifera ($P = 0.024$) and F. grandifolia ($P = 0.019$). No differences in $\delta^{13}$C-height slopes were detected between L. tulipifera and F. grandifolia ($P = 0.860$).

Coordination in leaf traits among the three species

Relationships between the coordinated leaf traits LMA, $N_{area}$ and $\delta^{13}$C were investigated to identify potential explanations for differences in leaf function and morphology among the three species (Figure 4). We found that LMA was positively correlated with $N_{area}$ and $\delta^{13}$C in all three species—the relation was linear in C. caroliniana and L. tulipifera, but non-linearly (quadratic in shape) correlated with $N_{area}$ and $\delta^{13}$C for F. grandifolia (Figure 4a and b). The interaction (LMA x Species) had a significant effect on both $N_{area}$ ($F = 5.76, P = 0.005$) and $\delta^{13}$C ($F = 3.17, P = 0.048$), indicating that the slopes were significantly different among species. In contrast, $N_{area}$ and $\delta^{13}$C were linearly correlated among all three species (Figure 4c), and the slopes of these relationships were similar among the three species ($F = 0.20, P = 0.818$).

Discussion

Light and height effects on LMA

Consistent with our first hypothesis, we found that light was more important than height in determining LMA for the indeterminate species, L. tulipifera and C. caroliniana. Light was also important in predicting LMA in F. grandifolia, but height was equally important. For L. tulipifera, F. grandifolia and other temperate deciduous species, both leaf thickness and LMA increase with increasing light availability (Goulet and Bellefleur 1985, Niinemets and Fleck 2002), suggesting that changes in LMA along vertical gradients in L. tulipifera and F. grandifolia are likely influenced by leaf thickness. Thicker leaves in temperate deciduous species tend to have thicker palisade layers (Eschrich et al. 1989, Oguchi et al. 2005), which facilitate greater light penetration to deeper layers of the mesophyll (Vogelmann and Martin 1993, Evans 1999). Leaf thicknesses of Fagus crenata, Fagus japonica and F. sylvatica leaves are determined during bud development in the summer of the previous year and during leaf expansion (Eschrich et al. 1989, Kimura et al. 1998, Uemura et al. 2000), which may explain why light was not considerably more important than height in regulating LMA for F. grandifolia.
Over the lifespan of deciduous leaves, the light environment can change considerably, particularly during leaf expansion when total crown leaf area rapidly increases from zero to a near-maximum value for the growing season (Parker and Tibbs 2004).

*Fagus grandifolia* is also more shade tolerant and has a greater $A_r/A_s$ than *L. tulipifera*, which is consistent with other studies that observed greater $A_r/A_s$ in *F. grandifolia* than other species (Matheny et al. 2014) and greater $A_r/A_s$ in more shade-tolerant species (Waring et al. 1982). There is strong evidence that $A_r/A_s$ plays an important role in whole-tree water relations (Whitehead et al. 1984, McDowell et al. 2002, Simonin et al. 2006). Within species, lower $A_r/A_s$ at drier sites (Mencuccini and Grace 1994) and in taller trees (McDowell et al. 2002) may act to compensate high evaporative demand and reduce gradients in leaf water potential. Similarly, the lower $A_r/A_s$ of the taller *L. tulipifera* may reduce gradients in leaf water potential thereby reducing height.

Figure 3. Relationships between LMA and total site factor (TSF) (a), LMA and height (b), $N_{area}$ and TSF (c), $N_{area}$ and height (d), $\delta^{13}$C and TSF (e), and $\delta^{13}$C and height for *C. caroliniana* (dotted lines), *L. tulipifera* (dashed lines) and *F. grandifolia* (solid lines).
effects on leaf development and LMA, whereas the high $A_iA_e$ in *F. grandifolia* may experience larger gradients in leaf water potential and height effects on LMA. Shoot $A_iA_e$ of *L. tulipifera* decreases with increasing height and light availability within tree crowns (Sack et al. 2006), which may also reduce gradients in leaf water potential brought about by height or greater exposure. Furthermore, greater investment in vascular tissue in leaves of *L. tulipifera* growing at high light may enhance water supply to leaves (Niinemets and Fleck 2002), potentially minimizing effects of height on LMA. It is worth noting that $A_iA_e$ may also compensate for changes in other factors such as leaf specific hydraulic conductivity without compensating for changes in gradients in leaf water potential. Furthermore, leaf water potential was not measured in this study. Thus, further work in identifying consequences of lower $A_iA_e$ on gradients in leaf water potential and leaf morphology among temperate deciduous species is required.

Regardless of the mechanistic explanations of LMA gradients for *F. grandifolia*, this species displayed a greater morphological plasticity in response to light and greater range of LMA compared with *L. tulipifera* and *C. caroliniana*. The high plasticity of *F. grandifolia* may partially explain why this species is successful in persisting in the understory of dense canopies for a long period of time (45–50 years), while occupying high-light, upper canopy positions at maturity (Canham 1990).

Vertical gradients in $\delta^{13}C$

Within forest canopies, the pattern of increasing leaf $\delta^{13}C$ with height has been attributed to both reduced stomatal conductance with height as a result of gradients in leaf water potential (e.g., decline in CO$_2$ supply; Koch et al. 2004, Ishii et al. 2008, Coble and Cavaleri 2015) and greater photosynthetic capacity with increasing light availability (e.g., increase in photosynthetic demand of CO$_2$ and greater RuBisCO; Berry et al. 1997, Duursma and Marshall 2006). Gradients of $\delta^{13}C$ of canopy air have been observed near the soil surface, but $\delta^{13}C$ of air has been found to be constant above 3 m (Berry et al. 1997) and contributes little to the vertical change in leaf $\delta^{13}C$ (Buchmann et al. 1997). In this study, there appears to be more evidence to support the hypothesis that gradients in $\delta^{13}C$ are associated with increasing CO$_2$ demand as a result of greater photosynthetic capacity in the upper canopy. $N_{\text{area}}$ is strongly correlated with photosynthetic capacity in many temperate deciduous tree species (Ellsworth and Reich 1993, Wilson et al. 2000), including the European beech (*F. sylvatica*; Scartazza et al. 2016), a congener of one of our species. $N_{\text{area}}$ and $\delta^{13}C$ were linearly and strongly correlated, providing evidence that greater $\delta^{13}C$ in the upper canopy was primarily a result of higher photosynthetic demand of CO$_2$ (e.g., higher RuBisCO activity) in the photosynthetic tissue. Similar to the results found in this study, increasing photosynthetic capacity (high CO$_2$ demand) was primarily
attributed to increasing δ\textsuperscript{13}C with canopy height in a conifer-dominated stand that displayed strong correlations between δ\textsuperscript{13}C and N\textsubscript{area} (Duursma and Marshall 2006).

**Coordination of LMA, N\textsubscript{area} and δ\textsuperscript{13}C**

Our results support our second hypothesis that LMA, N\textsubscript{area} and δ\textsuperscript{13}C are inter-related due to the link between leaf morphology and function. While N\textsubscript{area} and δ\textsuperscript{13}C displayed a similar response to light among the three species (e.g., similar slopes), we found that LMA responded differently among the three species. These results are unexpected for LMA and N\textsubscript{area}. Due to the use of LMA in calculating N\textsubscript{area} (N\textsubscript{area} = LMA × N\textsubscript{mass}) and the fact that N\textsubscript{mass} tends to be constant throughout the canopy, changes in N\textsubscript{area} typically follow changes in LMA. While N\textsubscript{area} generally followed changes LMA in this study, we also showed that for *F. grandifolia*, N\textsubscript{area} and LMA were non-linearly correlated, particularly at higher LMA where increases in LMA corresponded with diminishing increases in N\textsubscript{area}. This is likely due to lower N\textsubscript{mass} in the upper canopy for *F. grandifolia* (data not shown), leading to a non-linear relationship between N\textsubscript{area} and LMA. From a physiological viewpoint, this may be due to greater investment in nitrogen-poor compounds to leaves such as cell walls or cuticular waxes leading to a dilution effect on leaf nitrogen. In another tall, shade-tolerant tree (*Tilia cordata*), there was evidence that high LMA and denser leaves contained greater lignin concentrations and lower nitrogen concentrations, which likely improved foliar mechanical strength and tolerance to low leaf water potential while constraining photosynthetic capacity (Niinemets et al. 1999). In this study, structural adjustments to tolerate saturating light conditions and higher evaporative demand at greater heights for *F. grandifolia* appeared to have functional consequences. More specifically, N\textsubscript{area} (a proxy for photosynthetic capacity) may have been constrained due to greater investment in carbon-rich compounds to offset low water potential and increase mechanical strength for exposed leaves growing at greater heights. This may explain why both height and light were equally important for explaining LMA for *F. grandifolia*. While height appeared to be important in explaining LMA gradients and constraints on leaf function in *F. grandifolia*, leaf area above 20 m for *F. grandifolia* is relatively low compared with lower portions of the canopy and with other species at the same height in the same stand (Parker et al. 1989, Parker and Tibbs 2004).

**Conclusions**

This study compared differences in LMA, N\textsubscript{area} and δ\textsuperscript{13}C along canopy gradients in light availability among tree species with very different growth strategies and shade tolerance. Our study highlights the important finding that three species displayed differing LMA responses to light, but the N\textsubscript{area} response to light was consistent among all three species. Leaf mass per area of *C. caroliniana* and *L. tulipifera*, both indeterminate growth species, were finely tuned with light availability along the canopy profile, while that of *F. grandifolia*, a determinate growth species, was affected by both light and height. *Fagus grandifolia* also displayed the greatest morphological plasticity in response to light. The equal importance of light and height in *F. grandifolia* may be associated with the determinate growth strategy and/or a greater A\textsubscript{max}/A\textsubscript{N} compared with the other tall tree, *L. tulipifera*. There also appeared to be consequences of maintaining a higher LMA in the upper canopy for *F. grandifolia*, such as constraints on N\textsubscript{area} compared with *L. tulipifera*. While there was evidence for height effects in *F. grandifolia*, light appeared to be most important for explaining vertical gradients in LMA when considering all three species.

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**Conflict of interest**

None declared.

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