Ecophysiological responses to different forest patch type of two codominant tree seedlings

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Introduction
Tree species have varying leaf life spans, nitrogen concentrations, respiration rates, carboxylation rates, and other traits that are collectively described species’ ecophysiological traits (Kitao et al. 2000; Walters and Reich 2000; Miller et al. 2004; Niinemets 2010; Hitsuma et al. 2012). Ecophysiological traits are considered to influence where species can appear, which species can coexist, and how species respond to light conditions such as shading and full sun (Yoshimura 2010; Wyka et al. 2012). Studies on species distribution and coexistence often demonstrate that some environment factors can induce the plastic response of important ecophysiological traits (Miller et al. 2004; Niinemets 2010; Hitsuma et al. 2012). Vegetation is likely to experience heterogeneous patch habitats throughout the gap-phase dynamics. According to a theory of gap-phase dynamics proposed by Whitmore (1989), cycle progress is initiated by a natural disturbance. Four distinct patches may be recognized: a gap patch (G), building patch (B), mature patch (M), and degeneration patch (D) (Whitmore 1989). The forest community is composed of different patches, and the heterogeneous light conditions in the different patches can lead to changes in patch development and dynamics (Whitmore 1989).

Species-specific responses to heterogeneous environments reflect a number of adjustment mechanisms that enable plants to optimize gas exchange and resource investment strategy. Vegetation foliage modifies its ecophysiological and morphological trait to optimize ambient light utilization as light conditions change (Miller et al. 2004; Niinemets 2010; Hitsuma et al. 2012). Many studies have identified important species-specific light-harvesting factors (e.g., Yoshimura 2010; Wyka et al. 2012), but few have simultaneously researched the suite of other important ecophysiological traits in different patches.

Abstract
According to gap-phase dynamics theory, forests can be divided into four distinct patch types: gap patch (G), building patch (B), mature patch (M), and degeneration patch (D). Varying light conditions across patch types are one of the most important factors affecting the coexistence of vegetation. Mechanisms of coexistence can be understood through detailed knowledge of ecophysiological responses of codominant tree seedlings to patch types. The following study was conducted to determine ecophysiological responses of Cyclobalanopsis glauca (an evergreen broad-leaved species) and Bothrocaryum controversum (a deciduous broad-leaved species) to four different patch types. During the gap-phase dynamics, light intensity and the magnitude of change in the four different patches followed the order of: G > B > D > M. Both species had the greatest photosynthetic capacity in the G patch. Dry leaf mass per area (LMA), Chlorophyll a + b concentration (Chl), carotenoids (Car), and nitrogen content per area (Na) all responded to changes in light across patch type, but B. controversum showed greater sensitivity and changes than C. glauca. From G to M patch, the maximal quantum efficiency of PSII (Fv/Fm) had a larger variation magnitude for B. controversum than for C. glauca. From G to M patch, B. controversum showed significant changes in gas exchange, while C. glauca showed only small changes. Ecophysiological trait partitioning of response to light in different patches provides a possible explanation of a coexistence mechanism.

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Species-specific ecophysiological traits may be very important to their coexistence along G-B-M-D gradients with gap-phase dynamics, but ecophysiological traits responsible for plant functional types are not fully understood.

Leaves of evergreen species are thicker and longer-lived than leaves of deciduous species (Reich et al. 1998). In particular, it is unclear how plants of different functional groups coexist in terms of ecophysiological trait partitioning. *Cyclobalanopsis glauca* and *Bothrocaryum controversum* are common codominant tree species in the evergreen and deciduous broad-leaved forests of subtropical China. We hypothesize that ecophysiological traits of the two codominant species response to light environments along gap-phase dynamics (G-B-M-D) are similar, but that *B. controversum* (a deciduous species) has higher plasticity than *C. glauca* (an evergreen species) during the gap-phase dynamics, because deciduous plants are more sensitive to light changes than evergreen plants (Cao 2001; Miyazawa and Kikuzawa 2006; Böhne and Bruehlheide 2013). Here, we used ecophysiological traits of *C. glauca* and *B. controversum* for identifying their niche partitioning within the gap-phase dynamics.

**Materials and Methods**

**Study site and species**

An evergreen and deciduous broadleaf mixed forest was selected on the southern slope of the Dabieshan Mountain (31°02′35″ N, 116°06′47″ E, and 680 m a.s.l) in Anhui, eastern China. According to data from Yaoluoping climate station (30°59′07″ N, 116°04′56″ E, 1060 m a.s.l.), the mean annual air temperature is 12.3°C, and the mean annual precipitation is 2032.6 mm, falling mainly in the summer (44.2%). The codominant tree species in the experimental site (100 × 100 m) are *B. controversum* and *C. glauca*. The stand age is more than 150 years old by the tree ring analysis of the highest tree. The main subdominant tree species include evergreen plants, such as *Ilex pedunculosa*, *Eurya muricata*, and *Cyclobalanopsis gracilis*, and deciduous plants, such as *Rhododendron sinssi*, *Platycaea strobilacea*, and *Lindera glauca*. Common herbs are *Cardamine flexuosa*, *Salvia farinacea*, *Arthroxon hispidus*, and *Polygonum bungeanum*, while common vines included *Ficus sarmentosa* and *Paederia scandens*.

*Cyclobalanopsis glauca* and *B. controversum* seedlings were raised from seeds in the greenhouse of Anqing Normal University. All seeds of the same species were collected from the same tree in autumn 2009. In spring 2010, seeds were sown in 30 × 40 cm trays filled with a mixed substrate of sand and potting soil (1:2, v/v). All seeds were exposed to a 24/19°C day/night cycle and 14 h daytime. Three weeks after germination, they were transplanted to pots (1.6 L) filled with the same substrate. Plants were weeded periodically and watered to saturation to avoid water stress. In April 2010, 60 seedlings per species were selected and transplanted into 3.0-L pots filled with a mixed substrate of compost, vermiculite, and top forest soil from our field site (1:1:3, v/v/v).

**Patch chosen**

The experimental site was divided into 400 quadrats (5 × 5 m) (see Fig. 1). Height and DBH (diameter at breast height (cm)) of all trees were measured. Average tree height (H (m)) and the average height of the canopy trees (HC) were 6.9 m and 10.2 m, respectively. Using Daubenmire’s cover class method (Daubenmire 1959), we measured the percent cover of each quadrant (5 × 5 m).

Combining our observations and field work, we identified the following four development patches (Table 1). Gap patch (G): an open area with the brightest light conditions in the experimental site and identifiable gap makers and/or decomposing dead logs. The average tree height of regenerating layer is less than half that of canopy layer. Building patch (B): average tree height exceeds half of the average tree height of canopy layer. Some trees are close to the height of the main canopy layer. Mature patch (M): consists of mainly canopy trees with heights greater than the average tree height of the surrounding canopy. Degenerate patch (D): some large emergent trees with hollow trunks, sparse foliage, bald necrotic holes, and/or spots in the stems. The height of emergent trees is the same or exceeds the height of main canopy layer.

**Experimental design**

In March 2011, *C. glauca* and *B. controversum* seedlings that had been fostered in a greenhouse were transplanted to 20 randomly selected natural patch types (five replicates of G, M, B, and D patches). There were three *C. glauca* seedlings and three *B. controversum* seedlings in each patch. Each seedling was planted at random. Some herb and vine plants grew around our studied seedlings. Especially, parts of the stems and leaves may grow toward or twine around the trunk of the seedlings of *C. glauca* and/or *B. controversum*. To avoid their influence on the growth and survival of the seedlings, we removed the herb and vine plants around our studied seedlings. Throughout the experiment, seedlings were watered to saturation more than twice weekly to avoid water stress.

**Light intensity**

To reflect daily variation of light intensity in different patches, we used a scout™ light meter (3415FSE Dual
To compare the difference of light in four patch types, we randomly selected one patch type in 20 patches to measure the light intensity of every patch at a height of 1.60 m above the ground. This standard height was chosen as seedling height varied across the patch types. The light intensity in all four selected patches was measured every 2 h from 6:00 to 18:00 on each cloudless day (5–11 July 2012). The 20 values of light intensity in less than 1 min were recorded exactly at the same time by four researchers (one researcher in a patch) in each patch type.

Leaf gas exchanges were measured from 8:30 to 10:30 on sunny days (July 12–22, 2012) in the mid-to-upper parts of leaves (the region of the individual crown) of the selected plants. Only current year, fully developed leaves were selected to reduce the possible effect of leaf age. We used a portable photosynthesis measurement system (LI-6400, LI-COR Inc. Lincoln, NE) to determine the light and CO₂ photosynthetic response curves.

Table 1. The characteristics and classification standards of mosaics patch in an evergreen and deciduous broadleaf mixed forest.

| Patches | Characteristics | Classification standards |
|---------|-----------------|--------------------------|
| G       | 4.01 ± 0.34     | 1.01 ± 0.47              | 2.13 ± 0.62 | <6.9 | <6.9 | <6.9 | Yes | No | 0–10 |
| B       | 8.03 ± 2.12     | 4.74 ± 1.56              | 7.64 ± 3.11 | <6.9 | 6.9–10.2 | <10.2 | No | No | 10–30 |
| M       | 16.27 ± 2.98    | 10.65 ± 3.12             | 13.61 ± 4.34 | >6.9 | >10.2 | >10.2 | No | No | 60–100 |
| D       | 14.32 ± 2.45    | 8.12 ± 2.45              | 18.64 ± 5.17 | >6.9 | >10.2 | >10.2 | No | Yes | 40–60 |

Data are presented as mean ± standard error (SE). DBHq: average DBH. Hmax: maximum tree height. Hq: average tree height in the quadrate (5 m × 5 m). HCq: average tree height of canopy layer in the quadrate (5 m × 5 m). Emergent trees: tree height exceeds the main canopy layer in the sample. They have sparse leaves, broken branches, and their stems have some bald necrotic spots or holes.

Gas exchange measurement

Figure 1. The mosaics patch in an evergreen and deciduous broadleaf mixed forest (100 m × 100 m). The 20 patches with blue box are the selected patches.
Light response curves for both species were generated using 12 photosynthetic photon flux density (PPFD: 2000, 1500, 1000, 800, 600, 400, 200, 100, 50, 25, 15, and 0 μmol photons m$^{-2}$s$^{-1}$) with a stable CO$_2$ concentration (400 μmol·mol$^{-1}$). All measurements were obtained at a leaf temperature of 25°C, and the leaf-to-air vapor pressure difference (LVPD) was < 1.0 kPa. The maximal net photosynthetic rate ($P_{\text{max}}$) and the dark respiration rate ($R_d$) were derived from the light response curves using Photosyn Assistant Ver. 1.1.2 software (Dundee Scientific, Dundee, UK). At their saturating PPFD, CO$_2$ photosynthetic response curves were measured with similar leaf temperature (25°C) and LVPD (< 1.0 kPa). The leaves were exposed to 400 μmol mol$^{-1}$ CO$_2$ and 1000 μmol·m$^{-2}$·sec$^{-1}$ PPFD until their gas exchange was steady (at least 15 min). After steady-state photosynthesis was induced, we measured the responses of the photosynthetic rate ($P_n$) to changes in intercellular CO$_2$ concentration ($C_i$). The CO$_2$ concentration ($C_a$) was set to 400 μmol·mol$^{-1}$, and then it was sequentially lowered to 300, 200, 150, 100, and 50 μmol·mol$^{-1}$. Following this, $C_a$ was returned to 400 μmol·mol$^{-1}$, and then we measured CO$_2$ assimilation rate at four levels $C_a$: 800, 1000, 1200, and 1500 μmol·mol$^{-1}$. The $P_n$-$C_i$ curves were constructed from the measured values of $P_n$ and $C_i$ at each given $C_a$. The electron transport used in regeneration of RuBP ($J_{\text{max}}$), the maximal rates of carboxylation by Rubisco ($V_{\text{max}}$), and photosynthetic energy transformation efficiency ($\delta$) were calculated as described by Farquhar and Sharkey (1982), Loustau et al. (1999) and Schreiber et al. (1986), respectively.

**Chlorophyll fluorescence measurement**

After gas exchange measurements, we estimated the chlorophyll fluorescence parameters on the same leaves used for the gas exchange measurements using a portable pulse-modulated fluorometer (PAM-2100; Walz, Effeltrich, Germany) connected to a computer running control software. One B. controversum seedling and one C. glauca seedling were randomly selected in each patch. Total eight seedlings (four B. controversum seedlings and four C. glauca seedlings) were measured in four selected patches (the same four patches as those measured in light intensity).

For consecutive measurements of chlorophyll fluorescence parameters, we marked a fixed position on each sampled leaf. To measure the minimal fluorescence ($F_0$) and maximal fluorescence ($F_m$) parameters, the sample was first dark-adapted for at least 20 min using a leaf clip. The maximal photochemical efficiency of PSII ($F_{\text{m}}/F_m$) and apparent electron transport rate (ETR) were measured at 2-h intervals from 6:00 to 18:00 on 23–29 July 2012.

**Leaf morphology**

Following the measurements of chlorophyll fluorescence and gas exchange, we harvested the leaves, extracted leaf disks (1 cm diameter), and then separated these leaf disks into two parts. One part was used to determinate the chlorophyll a (Chla), chlorophyll b (Chlb), carotenoid (Car), and total chlorophyll content per unit area (Chl) using UV-vis spectrophotometry after extraction with dimethylformamide. The other part was dried at 60°C for 48 h. We calculated the LMA according to the dry weight per area and then ground these dried leaves to fine powder for nitrogen analysis by the Kjeldahl method. Finally, we calculated photosynthetic nitrogen use efficiency ($PNUE = P_{\text{max}}/N_a$), the ratio of Chl to N$_a$ (Chl/N$_a$), and the ratio of Chla to b (Chla/b), where N$_a$ is the area-based leaf nitrogen content. According to the method proposed by Niinemets and Tenhunen (1997), we assessed the leaf nitrogen allocated to bioenergetics ($P_{\text{b}}$), Rubisco ($P_{\text{R}}$), and light-harvesting components ($P_{\text{l}}$).

**Seedling survival, height growth, and dry weight**

When the experiment ended (July 2012), seedling survival rates per species and patch were recorded. Seedling height was measured, and relative growth rate (RGR) was calculated as $RGR = (\ln w_2 - \ln w_1)/\tau$, where $w_2$ and $w_1$ represent the final and initial height at time $\tau$ and $t_1$, respectively. Four seedlings per species and treatment ($n = 32$) were harvested, and they were washed carefully to avoid losing fine roots. They were oven-dried at 70°C for 72 h, and the dry weight of roots, shoots, and leaves was measured. The root mass ratio (RMR), shoot mass ratio (SMR), and leaf mass ratio (LMR) were calculated as ratio of dry weight of each fraction to the total biomass.

**Statistical analysis**

Statistica v8.0 (StatSoft Inc. Tulsa, Oklahoma, USA) was used to perform all analyses. Each value of ecophysiological characteristics was presented as the mean and standard error (SE).

Spatial autocorrelation of the selected patched was calculated as Moran’s I (Fortin and Dale 2005), and we did not observe that there lied a spatial autocorrelation. The survival of seedlings was analyzed with the log-likelihood
ratio test (G-test). Differences between species and treatments were analyzed by repeated measures ANOVA using Tukey’s test. Before ANOVA, data were log-transformed to meet assumptions of homogeneity of variance and normality when necessary.

Results

Light intensity

From 06:00 h to 18:00 h, light intensity followed a diurnal pattern increasing to midday and then decreasing. The greatest value appeared at 14:00 h (Fig. 2). Light intensity and the change magnitude (the difference of the max value and the min value from 6:00 to 18:00) in the four different patches followed the order of: G > B > D > M (Fig. 2).

Photosynthetic traits

For *B. controversum*, the maximum net photosynthetic rate (P_{\text{max}}), the photosynthetic nitrogen use efficiency (PNUE), the photosynthetic energy transformation efficiency (\(\delta\)), the maximum carboxylation rate (V_{\text{cmax}}), and the maximum electronic transfer rate (J_{\text{max}}) differed significantly among the four different patches. The highest values were in the G patch, and the lowest values were in the M patch (Table 2). All measured gas exchange parameters for *C. glauca* were not significantly different among four patches. The dark respiration rate (R_{\text{d}}) for *C. glauca* was significantly different (Table 2). In the G patch, the gas exchange parameters (P_{\text{max}}, PNUE, \(\delta\), V_{\text{cmax}} and J_{\text{max}}) of *B. controversum* were higher than those of *C. glauca* (Table 2), but in the M patch, these

![Figure 2](image_url). The diurnal variation of the light intensity in four selected patches in the gap-phase dynamics (G, B, M and D) at a height of 1.60 m on seven cloudless days (5–11 July 2012).

| Species | P_{\text{max}} (\text{mol m}^{-2} \text{sec}^{-1}) | PNUE | \(\delta\) | V_{\text{cmax}} (\text{mol m}^{-2} \text{sec}^{-1}) | J_{\text{max}} (\text{mol m}^{-2} \text{sec}^{-1}) |
|---------|----------------------------------|------|---------|----------------------------------|----------------------------------|
| G       | 10.45 ± 1.89 a                   | 0.26 ± 0.03 a | 0.23 ± 0.03 a | 11.25 ± 1.53 a | 3.02 ± 0.03 a |
| B       | 6.54 ± 0.96 b                    | 0.25 ± 0.03 b | 0.21 ± 0.03 b | 10.73 ± 2.09 b | 2.90 ± 0.40 b |
| M       | 3.27 ± 0.07 b                    | 0.26 ± 0.03 b | 0.26 ± 0.04 a | 19.78 ± 2.14 b | 5.89 ± 0.56 b |
| D       | 8.18 ± 0.80 b                    | 0.24 ± 0.06 b | 0.24 ± 0.04 a | 28.58 ± 4.24 b | 5.97 ± 0.89 b |
| G       | 4.88 ± 0.39 b                    | 0.25 ± 0.03 b | 0.19 ± 0.02 b | 26.36 ± 4.24 b | 4.46 ± 0.31 b |
| B       | 4.35 ± 0.39 b                    | 0.24 ± 0.02 b | 0.16 ± 0.02 b | 27.41 ± 1.23 b | 6.52 ± 0.37 b |
| M       | 4.11 ± 0.14 b                    | 0.14 ± 0.03 b | 0.17 ± 0.02 b | 26.46 ± 4.18 b | 5.35 ± 0.55 b |
| D       | 4.41 ± 0.51 b                    | 0.14 ± 0.03 b | 0.16 ± 0.04 a | 28.02 ± 4.12 b | 5.53 ± 0.64 a |

Data are presented as mean ± standard error (SE). Means with different letters differ significantly in the same species and among different patches (a > b > c > d). PNUE is the photosynthetic nitrogen use efficiency (\(\mu\text{mol m}^{-2} \text{sec}^{-1}\)), \(\delta\) is the photosynthetic energy transformation efficiency (electrons quanta\(^{-1}\)), V_{\text{cmax}} is the maximum carboxylation rate (\(\mu\text{mol m}^{-2} \text{sec}^{-1}\)), and J_{\text{max}} is the maximum electronic transfer rate (\(\mu\text{mol m}^{-2} \text{sec}^{-1}\)). P_{\text{max}} and R_{\text{d}} are the leaf nitrogen allocated to Rubisco, bioenergetics, and light-harvesting components, respectively.
parameters of B. controversum were lower than those of C. glauca (Table 2). Compared to the other three patches, B. controversum and C. glauca in the M patch both had the lowest values of the leaf nitrogen allocated to Rubisco ($P_b$) and bioenergetics ($P_L$), as well as the highest leaf nitrogen allocated to light-harvesting components ($P_L$) (Table 2). Only $P_L$ showed a significant difference among the four patches for both tree species.

**Chlorophyll fluorescence measurements**

In the G patch, the maximal quantum efficiency of PSII ($F_v/F_m$) values of B. controversum and C. glauca was the highest (around 0.8). The two species both had significant differences among the four different patches in the PSII ($F_v/F_m$) (Fig. 3). From G to M patch, the $F_v/F_m$ values of B. controversum and C. glauca decreased by approximately 44.4% and 22.8%, respectively (Fig. 3). At G patch, the values of the photochemical quenching ($q_N$), the actual photochemical efficiency of PSII ($\Phi_{PSII}$), and electron transport rate (ETR) measured late morning/early afternoon were higher for B. controversum than for C. glauca (Fig. 4). At 14:00 h in the G patch, there was some photoinhibition (a decreasing photosynthetic efficiency with high light at noon) for the two species, as nonphotochemical quenching ($q_N$) was more than 0.6, and $q_P$, $\Phi_{PSII}$, and ETR were less than 0.5, 0.4, and 60 µmol electrons m$^{-2}$.sec$^{-1}$, respectively (Fig. 4). In the G patch, $q_N$, $\Phi_{PSII}$, and ETR have a greater diurnal variation (the difference of the max value and the min value from 6:00 to 18:00) for B. controversum than for C. glauca (0.44 vs. 0.39 of $q_N$, 0.539 vs. 0.301 of $\Phi_{PSII}$, 92 vs. 41 of ETR), while $q_P$ has no significant difference (Fig. 4). In the M patch, $q_P$ and $\Phi_{PSII}$ have a greater diurnal variation for B. controversum than for C. glauca (0.47 vs. 0.37 of $q_P$, 0.105 vs. 0.037 of $\Phi_{PSII}$), while $q_N$ and ETR have no significant difference (Fig. 4).

**Leaf morphology**

Bothrocaryum controversum had a lower leaf dry mass per area (LMA), but higher nitrogen content per area ($N_a$), Chl a + b concentration per area (Chl), and carotenoid (Car) in all patches compared to C. glauca (Fig. 5A–D). Bothrocaryum controversum had a significantly lower LMA, $N_a$, and Car in the M patch compared to the other three patches, while no significant difference was observed for C. glauca in all patches (Fig. 5A, B, and D). The two species both had significantly higher Chl and Chl$b$ in the M patch than in the other three patches (Fig. 5C, E), and no significant differences in Chl$a/b$ in all patches (Fig. 5F).

**Seedling survival and growth**

The survival of B. controversum showed a significant change from G to M (92% to 36%), while C. glauca showed only a small change (86–70%). Relative growth rate (RGR) differed significantly between patches for both species with the greatest values appearing in the G patch and the smallest values in the M patch (Table 3). For B. controversum, root mass ratio (RMR) and shoot mass ratio (SMR) decreased significantly and leaf mass ratio (LMR) increased significantly from the G to M patch as shading increased (Table 3). Conversely, RMR and LMR for C. glauca for the same conditions showed the opposite changes though changes were small and not significant (Table 3).

**Discussion**

Light is very important for the survival, growth, and development of higher plants (Valladares et al. 1997; Walters and Reich 2000; Durand and Goldstein 2001; Hitsumta et al. 2012). During the gap-phase dynamics, the light environment varies diurnally and by patch type. Individual plants are in reality fine-tuning the expression of physiological traits (within the range possible given their genetic makeup) to optimize their carbon gain at whatever environmental condition they are experiencing. The leaf is a good representation of the ability of a plant to respond to light changes (Poorter and Bongers 2006).
Ecophysiological responses to G patch (high light)

Gap is an important forest growth stage driving productivity and plant regeneration (Houter and Pons 2005; Kitaoka et al. 2009; Kuptz et al. 2010). In the G patch, B. controversum and C. glauca seedlings both had the highest photosynthetic capacity growth ratio, and survival, indicating that a forest gap is beneficial to growth and regeneration. Our results are in agreement with previous gap dynamics research (Gómez-Aparicio et al. 2006; Oguchi et al. 2008; Kitaoka et al. 2009; Kuptz et al. 2010). Photoinhibition was observed midday in the G patch due to high-intensity light conditions. Seedlings need to develop protective mechanisms to reduce light absorbance and avoid photooxidation in high light. These mechanisms include increasing the carotenoid concentration and nitrogen content per area (Gómez-Aparicio et al.
decreasing the ratio of Chl to Na, and the chlorophyll content per area. In addition, to prevent high-intensity light damage, species tend to invest more nitrogen in protecting leaf structure (e.g., the epidermis) and invest little nitrogen in photosynthetic organs (Oguchi et al. 2008; Kitaoka et al. 2009; Kuptz et al. 2010). These protective mechanisms were observed in the two codominant tree seedlings in this study.

**Ecophysiological responses to M patch (low light)**

Plants with different functional types develop acclimation mechanisms to optimize ambient light utilization under low light conditions (Miller et al. 2004; Yoshimura 2010; Hitsuma et al. 2012; Wyka et al. 2012). Natural growth and regeneration in lower light conditions are related to
the photosynthetic capacity in combination with morphological and physiological changes (Gommers et al. 2013). Changes induced by competition for light reflect plant shade avoidance or shade-tolerance ability (Gommers et al. 2013). Under low light in the M patch, shade-avoidance species, like B. controversum, invested relatively higher mass ratios to leaf (higher LMR) at the expense of root biomass (lower RMR), while shade-tolerance species, like C. glauca, showed little change. In addition, plants have a trade-off in leaf N allocation to Rubisco, bioenergetics, and light-harvesting components. Higher leaf N allocated to light-harvesting organs under low light is associated with lower investment in Rubisco, which induces lower PNUE (Xiang et al. 2013). In our study, the evergreen species, C. glauca, showed increased Chl:N ratios, Pn, and PNUE in the M patch as a result of the increase in chlorophyll, which indicates that the C. glauca performs well under low light. Bothrocaryum controversum had a poor shade acclimation, indicating a trade-off between high light (in the G patch) and shade adaptation (in the M patch). Different responses in morphology and physiology to low light can induce mortality difference (Gómez-Aparicio et al. 2006; Feijó et al. 2009; Gommers et al. 2013). We observed that the mortality of B. controversum was approximately two times higher than that of C. glauca in the M patch. The different mortality is a good indicator of their interspecific differences in low light tolerance.

Heterogeneous ecophysiological trait partitioning to gap-phase dynamics

Environmental resource segregation and trade-offs in ecophysiological traits are commonly used to study whether trait partitioning is beneficial to species coexistence for different functional groups (Gómez-Aparicio et al. 2006; Niinemets 2010). Codominant species have different resource acquisition methods by ecophysiological trait partitioning (Gómez-Aparicio et al. 2006). During the gap-phase dynamics, B. controversum exhibited higher degree of change, which was in accordance with its shorter leaf life span and higher potential photosynthetic rates. The greater changes within the gap-phase dynamics were inherently associated with the higher flexibility in utilizing available resources in different patches. C. glauca exhibited small changes with slow growth and little variation in ecophysiological traits during the gap-phase dynamics, for evergreen species have a stable physiological performance due to its sclerophyllous and long-living leaves (Böhneke and Bruelheide 2013). The various interspecific responses to the four different patches provide new insights into the coexistence mechanism for B. controversum and C. glauca with the gap-phase dynamics.

Light interception and utilization efficiency along light gradients changes for plants of different functional groups (Niinemets 2010). A suite of traits, such as the biomass allocation to leaves, leaf chlorophyll content, leaf dry mass per unit area, and leaf nitrogen allocation, at various scales and plasticity determine plant light-harvesting efficiency (Feijó et al. 2009; Niinemets 2010). Mechanisms of light interception by seedlings of different functional groups have been addressed frequently as one of the most important causes of species coexistence and biodiversity conservation in tropical and subtropical forests (Feijó et al. 2009; Niinemets 2010; Gommers et al. 2013). Our results indicate the importance of temporal and spatial variation of light during the gap-phase dynamics and fluctuating ecophysiological trait partitioning for stable coexistence of B. controversum and C. glauca.

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Conflict of Interest

The authors declare that there is no conflict of interest with any financial organization mentioned in the manuscript.

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