## Abstract

Plants exhibit higher leaf-to-root ratios (L/R) and lower leaf nitrogen content \( (N_{\text{area}}) \) in low-light than in high-light environments, but an ecological significance of this trait has not been explained from a whole-plant perspective. This study aimed to theoretically and experimentally demonstrate whether these observed L/R and \( N_{\text{area}} \) are explained as optimal biomass allocation that maximize whole-plant relative growth rate (RGR). We developed a model which predicts optimal L/R and \( N_{\text{area}} \) in response to nitrogen and light availability. In the model, net assimilation rate (NAR) was determined by light-photosynthesis curve, light availability measured during experiments, and leaf temperature affecting the photosynthesis and leaf dark respiration rate in high and low-light environments. Two pioneer trees, *Morus bombycis* and *Acer buergerianum*, were grown in various light and nitrogen availabilities in an experimental garden and used for parameterizing and testing the model predictions. They were grouped into four treatment groups (relative photosynthetic photon flux density, RPPFD 100% or 10% × nitrogen-rich or nitrogen-poor conditions) and grown in an experimental garden for 60 to 100 days. The model predicted that optimal L/R is higher and \( N_{\text{area}} \) is lower in low-light than high-light environments when compared in the same soil nitrogen availability. Observed L/R and \( N_{\text{area}} \) of the two pioneer trees were close to the predicted optimums. From the model predictions and pot experiments, we conclude that the pioneer trees, *M. bombycis* and *A. buergerianum*, regulated L/R and \( N_{\text{area}} \) to maximize RGR in response to nitrogen and light availability.

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* E-mail: dsugiura@biol.s.u-tokyo.ac.jp

## Introduction

Plants have the ability to alter their phenotype to maximize fitness according to the external environment. For example, they often change leaf properties and biomass allocation pattern in accordance with light and nutrients conditions [1,2]. Criteria for determining leaf to root ratio (L/R) have been investigated by many researchers because it could be a major factor dictating plant growth rate and fitness [3]. Thus, elucidating the L/R will be helpful for understanding plant growth strategies in natural ecosystems.

Until now, many researchers have worked with the subject, and proposed the balanced growth hypothesis where plants allocate more biomass to the organ capturing the most limiting resources, such as light and nutrients [4–6]. According to this hypothesis, for example, producing more leaves at the sacrifice of root growth is favoured in low-light environments to capture more light to enhance growth rate. However, this hypothesis is only an intuitive explanation and can’t propose a quantitative estimation of L/R. Since leaf and root functions are closely interrelated, producing excessive leaves may decrease growth rate due to decreased root functions, such as nitrogen uptake capacity. This lead to an idea that there will be an equilibrium between leaves and roots for optimal biomass allocation that maximizes whole-plant growth rate [1,7].

Theoretical analyses and experimental confirmation of the hypothesis have been performed for plants in high-light environments. Such studies revealed that the L/R and leaf nitrogen content were mainly optimized to maximize RGR with soil nitrogen availability [7,8]. In contrast, plants growing in low-light environments generally have higher L/R than those growing in high-light environment regardless of functional groups [1,9–13]. In these studies, however, the high L/R were only explained from the balanced growth hypothesis as mentioned above, and theoretical studies accounting for this biomass allocation pattern are still lacking. Thus, it has not been quantitatively determined whether the high L/R in low-light environments is as a result of maximization of relative growth rate (RGR) to maximize.

We noticed that plants growing in low-light environments have lower leaf nitrogen content per leaf area \( (N_{\text{area}}) \) and associated lower maximum photosynthetic and dark respiration rate [1,9,14]. Nitrogen is almost thoroughly absorbed by root and considered to be a primary mineral which dictate amount of photosynthesize and growth [15] through the balance between photosynthetic and respiration rate and light availability. For example, higher \( N_{\text{area}} \) realize higher maximum photosynthetic rate, but if light availability is low, the amount of photosynthesize rather decreases because dark respiration rate is also higher. Therefore, it is hypothesized that the higher L/R might be due to lower nitrogen demand for maximizing growth rate than in high-light environments, not due to capturing more light by increasing leaf area at the sacrifice of root growth in low-light environments. Since \( N_{\text{area}} \) is determined by L/R, leaf mass per unit area, and root nitrogen uptake capacity, we are able to estimate optimal L/R and \( N_{\text{area}} \).
which maximize whole plant growth rate by considering above-mentioned plant traits. In this study we developed a biomass allocation model based on that of Osone and Tateno (2005) to demonstrate whether the L/R and $N_{area}$ in a low-light environment are optimized to maximize relative growth rate (RGR). Leaf (leaf mass per area and photosynthesis) and root (nitrogen absorption) properties were incorporated into the model. We also estimated the leaf net assimilation rate (NAR; g m$^{-2}$ d$^{-1}$) in various light environments to clarify relationship between light availability and nitrogen demand. There, $N_{area}$ and photosynthetic parameters were associated with actual meteorological data measured throughout the growth period. Using the $N_{area}$ - NAR relationship, we could predict the optimal L/R and $N_{area}$ in various light environments. Two pioneer trees, *Morus bombycis* and *Acer buergerianum*, were used for testing the model predictions. Finally, we discuss the biomass allocation strategy in a low-light environment from a whole-plant perspective.

**Materials and Methods**

**Plant materials and experimental design**

Experiments were conducted at the Nikko Botanical Gardens of the University of Tokyo (139°360’E, 36°450’N, 650 m a.s.l.). The mean air temperature was 12°C, and the annual precipitation was 2100 mm.

We used 1-year-old seedlings of mulberry tree (*Morus bombycis* Koidz.) and Trident maple tree (*Acer buergerianum* Miq.). These are typical pioneer deciduous trees in East Asia, which change their morphological and physiological traits largely. Seedlings grow fast because leaves flush sequentially and root growth continues throughout the growing season.

*Morus bombycis* seeds were collected from a wild *M. bombycis* tree in Nikko city in 2007. The seedlings were grown in plastic pots in an open field in 2007 and used for experiments from April to August 2008. One-year-old *A. buergerianum* seedlings which were grown in natural open environments were purchased from a nursery (Kairyoen, Saitama, Japan). They used for experiments from July to September 2009. Initial pot size was about 3 liter and seedlings were further transplanted carefully to 10 liter pots according to root size. Until just before the experimental period, those seedlings were placed in shade houses which were made of greenhouse frames and shade cloths. Relative photosynthetic flux density (RPPFD) in the shade houses was about 10% (measured by two quantum sensors, LI-1000, Li-Cor, Lincoln, NE, USA).

At the beginning of the experimental period, the main stem of each seedling was cut, and only one shoot was allowed to grow. Then, half of the seedlings were placed in the open field and the rest were in the shade houses, respectively. Pots were placed separately to avoid mutual shading. They were also grouped into two nutrient conditions with different nitrogen concentrations. Other than N, these solutions contained the following: 3 mM K$_2$HPO$_4$, 1 mM MgSO$_4$·7H$_2$O, 3 mM CaCl$_2$, 25 mM H$_2$BO$_3$, 2 μM MnSO$_4$·5H$_2$O, 2 μM ZnSO$_4$·7H$_2$O, 0.5 μM CuSO$_4$·5H$_2$O, 0.5 μM Na$_2$MoO$_4$·2H$_2$O, and 20 μM Fe-EDTA [16]. NH$_4$NO$_3$ was added to this solution and adjusted to 20 or 50 μM to avoid mutual shading. They were also grouped into two quantum sensors, LI-1000, Li-Cor, Lincoln, NE, USA).

Pot seedlings from all four treatments were used for the measurements. The measurement conditions were as follows: CO$_2$ concentration, 400 µmol mol$^{-1}$; leaf temperature, 25°C; and relative humidity, 50%. The maximum photosynthetic rate was measured at 1000 µmol m$^{-2}$ s$^{-1}$ for the sun-exposed leaves (100%RPPFD) and at 200 µmol m$^{-2}$ s$^{-1}$ for the shade leaves, so as not to cause photoinhibition. We also measured temperature dependency of photosynthetic rate by changing leaf temperature and irradiance variably. After the measurements, total nitrogen content of the leaves was measured for evaluating $N_{area}$ by a carbon-nitrogen (CN) analyzer (Vario EL, Elemental Analyse-systeme GmbH, Hanau, Germany).

**Sampling**

*Morus bombycis* were harvested in mid-April and mid-August of 2008, and *A. buergerianum* were harvested in early July and early September. Final biomass of the seedlings became much larger than initial biomass. Seedlings seemed not to be self-shaded because they had only one shoot per individual. At each harvest, four to ten seedlings per treatment group were sampled and divided into leaves, stems, and roots. After measuring the leaf area, each part of the seedlings was oven-dried at 80°C for more than 4 days. The samples were then weighed, and nitrogen content was measured with the CN analyzer.

**Calculation**

Nitrogen absorption rates per unit root dry mass (SAR; g N g$^{-1}$ d$^{-1}$) were calculated considering the difference in total nitrogen content and root dry mass between the two harvests following Osone & Tateno (2005). Changes in root dry mass were assumed to be exponential between harvests. The leaf mass per area (LMA; g m$^{-2}$), L/R, and leaf mass per shoot mass ($P_{leaf}$; g g$^{-1}$) were also determined for each treatment group and applied for model prediction.

**The models**

First, we developed an optimal growth model that predicts the optimal biomass allocation ratio and leaf nitrogen content under various irradiance levels. The structure of the model was fundamentally based on that described by Osone and Tateno (2005).

In our model, the $N_{area}$ - NAR relationship was used as the plant growth indicator. NAR was estimated using an actual PPFD and photosynthetic light-response curve in which the temperature dependency of the photosynthesis and dark respiration rate were considered.

Net photosynthetic rate at certain PPFD ($I$) and leaf temperature ($T_L$), $A_o(I,T_L)$, was expressed as follows:

$$A_o(I,T_L) = A_s(I,T_L) - R_d(T_L)$$

(1)
where $A(I, T_L)$ is the gross photosynthetic rate at $I$ and $T_L$ and $R_d(T_L)$ is the leaf dark respiration rate at $T_L$ (°C). $A(I, 25)$ is measured and expressed as the photosynthetic light-response curve which is a non-rectangular hyperbola:

$$A(I, 25) = \frac{\psi I + A_{\text{max}} - \left( (\psi I + A_{\text{max}})^2 - 4\psi I A_{\text{max}} \right)^{1/2}}{2\psi}$$

(2)

where $A_{\text{max}}$ is the light-saturated rate of gross photosynthesis, $\psi$ is the initial slope of the light-response curve, $\theta$ is the convexity of the light-response curve, $A_{\text{max}}$ and $R_d(25)$ are expressed as a function of $N_{\text{area}}$ and $\theta$ and $\psi$ are assumed to be constant.

The temperature dependency of gross photosynthetic rate was incorporated as a function of $T_L$. $A(I, T_L)$ was expressed as an quadric approximation formula where $A(I, 25)$ was relativized to 1 as a standard value:

$$A(I, T_L) = A(I, 25)(a_1 T_L^2 + a_2 T_L + a_3)$$

(3)

where $a_1$, $a_2$ and $a_3$ were constant values and obtained from the photosynthesis measurements for each species.

The temperature dependency of $R_d(T_L)$ is described as [17]:

$$R_d(T_L) = R_d(25) \exp \left[ \frac{\Delta H_s(T_L - 25)}{298 R (T_L + 273)} \right]$$

(4)

where $R_d(T_L)$ and $R_d(25)$ are values of $R_d$ at $T_L$ (°C) and 25°C, respectively. $R$ is the gas constant (0.0083 J K$^{-1}$ mol$^{-1}$) and $\Delta H_s$ is the activation energy of $R_d$ (66.405 kJ mol$^{-1}$) [18]. We also considered the Kok effect by which the dark respiration rate decreases when leaves are exposed to sunlight [19,20]:

$$R_d(T_L) = 0.4 R_d(T_L) \quad \text{(when } I > 0 \mu \text{mol m}^{-2} \text{s}^{-1})$$

(5)

$$R_d(T_L) = R_d(T_L) \quad \text{(when } I = 0 \mu \text{mol m}^{-2} \text{s}^{-1})$$

(6)

where eqn. 5 and eqn. 6 represent the dark respiration rate during the day and night, respectively.

The leaf photosynthesis parameters ($A_{\text{max}}$, $R_d$, $\theta$, and $\psi$) at 25°C were described following Hikosaka et al. (1999). Relationship between $A_{\text{max}}$–$N_{\text{area}}$ relationship and $R_d$–$N_{\text{area}}$ relationship were expressed as:

$$A_{\text{max}} = b_1 \frac{N_{\text{area}} - b_2}{N_{\text{area}} - b_2 + b_3}$$

(6)

$$R_d = b_4 N_{\text{area}} + b_5$$

(7)

where $b_1$, $b_2$, and $b_3$ were maximum rate of $A_{\text{max}}$, x-intercept of the curve, and a constant that determines initial slope of the $A_{\text{max}}$ – $N_{\text{area}}$ relationship, respectively, and $b_4$ and $b_5$ were the slope and y-intercept of the $R_d$ – $N_{\text{area}}$ relationship. These parameters were obtained from the photosynthesis measurements for each species.

For a given $N_{\text{area}}$, NAR was calculated by substituting the light dataset into above equations (eqn. 1 to 6), integrating $A(I)$, converting CO$_2$ to carbohydrate (1/6C$_6$H$_{12}$O$_6$), multiplying a transform coefficient of assimilated carbohydrate to the structural carbohydrate, and dividing the integrated $A(I)$ by the growth period (day). The transform coefficient was found to be about 0.4, in which both construction and maintenance costs of leaves, stems, and roots were considered [21–23]. We also estimated NAR in low-light environments using datasets with PPFD reduced to 10% and repeated the above processes.

We determined optimal plant property values using the $N_{\text{area}}$ – NAR relationship and an optimal biomass allocation model based on that of Osone and Tateno (2003). The model plant consisted of three parts: the leaf, stem, and root. The whole plant biomass ($H$) is expressed as:

$$W = W_L + W_S + W_R$$

(M1)

where $W_L$, $W_S$, and $W_R$ are the leaf, stem, and root biomass, respectively. Leaf area ($L\alpha$) is expressed as:

$$L\alpha = \frac{W_L}{LMA}$$

(M2)

where LMA is the leaf mass per area (g m$^{-2}$), a constant determined in each light environment.

Leaf nitrogen content per biomass ($N_L$) is different from stem and root nitrogen content per biomass ($N_S$ and $N_R$) and they are highly correlated. Because these relationships affect the prediction of optimal biomass allocation (Osone & Tateno 2003), we defined $N_S$ and $N_R$ as functions of $N_L$ as follows:

$$N_S = c_1 N_L + c_2$$

(M3)

$$N_R = c_3 N_L + c_4$$

(M4)

where $c_1$, $c_2$, $c_3$, and $c_4$ are constant values. By introducing these relationships, absorbed nitrogen is partitioned into leaf, stem, and root correctly.

Leaf nitrogen content per leaf area ($N_{\text{area}}$) is expressed as:

$$N_{\text{area}} = \frac{N_L}{L\alpha}$$

(M5)

Plant biomass production per day is a product of net assimilation rate (NAR) and $L\alpha$:

$$\frac{dW}{dt} = NAR \cdot L\alpha$$

(M6)

Newly produced biomass was first divided between shoot and root following Hilbert (1990) using the allocation coefficient $P_{\text{Shoot}}$, which is shoot biomass per total biomass. Then, newly shoot biomass is further partitioned into shoot leaf and stem according to $P_{\text{Leaf}}$ which is leaf biomass per shoot biomass, following Osone & Tateno (2003). Because there was almost no variation in $P_{\text{Leaf}}$ during the growth period for each species growing in each light environment, we only have to estimate the effect of $P_{\text{Shoot}}$ in the model simulation. Thus, the new biomass increment for each organ is expressed as:

$$\frac{dW_L}{dt} = P_{\text{Shoot}} \cdot P_{\text{Leaf}} \cdot \frac{dW}{dt}$$

(M7)

$$\frac{dW_S}{dt} = P_{\text{Shoot}} \cdot (1 - P_{\text{Leaf}}) \cdot \frac{dW}{dt}$$

(M8)
Optimal Biomass Allocation and Leaf Nitrogen

where 0<P_{Shoot}<1 and 0<P_{Leaf}<1.

Nitrogen uptake rate is proportional to the root biomass:

\[
\frac{dW_R}{dt} = (1 - P_{Shoot}) \frac{dW}{dt}
\]

(M9)

where N is total nitrogen content and SAR is the specific absorption rate, which represents both the plant nitrogen uptake capacity of the roots and soil nitrogen availability [24]. Then, the RGR is calculated as:

\[
RGR = \frac{dW}{dt} \frac{1}{W}
\]

(M10)

For the given plant growth parameters (N_{area} - NAR relationship, P_{Shoot}, P_{Leaf} and SAR), steady-state L/R, N_{area} and RGR are obtained by repeating the model processes numerically by Euler’s method. By changing P_{Shoot} the optimal biomass allocation rate and N_{area} which maximizes RGR for various RPPFD and nitrogen availabilities, can be calculated.

As the N_{area} - NAR relationship differed in each light environment, we obtained the optimal L/R and N_{area} under different light and nutrient conditions.

Results

Model parameters

Parameter values obtained from the measurements, determination coefficients, and corresponding equations were listed in Table 1, which showed good correlation. Only the relationships between nitrogen content of each organ were determined for each light environment (eqn. M3, M4) using c1, c2, c3, and c4.

As for the leaf photosynthesis parameters (A_{max}, R_{k}, \theta, and \phi) at 25°C, parameters of M. bombycis were shown in Fig. 1(A-D) as representative. A_{max} and R_{k} were highly correlated with N_{area} (Fig. 1A, D). Averaged values were used for \phi and \theta for both species because these values were almost constant irrespective of the N_{area} and light environments (Fig. 1B, C).

During the growth periods in 2008 and 2009, mean diurnal air temperatures were 20.5 and 23°C, and the average daily PPFDs were 22.5 and 22.1 (mol m^{-2} d^{-1}), respectively. A T_{L} estimating equation was developed by multi-regression analysis using the recorded PPFD and T_{a} values measured in August 2009. For PPFD, the accumulated value of the last 3 minutes, P_{min} (mol m^{-2}) was used for the multi-regression analysis because it showed the highest correlation. The equation is expressed as:

\[
T_{L} = 0.875T_{a} + 9.61 \times 10^{-4}P_{3min} + 2.94 \ (r^2 = 0.96)
\]

T_{L} at night was set to night T_{a} because these values were nearly the same.

NAR was calculated by substituting the observed environmental data into the N_{area} - photosynthesis relationship and expressed as a function of N_{area} for both species and light environments (Fig. 2). NAR increased with N_{area} for 100% RPPFD, whereas it reached a maximum value at low N_{area} for 10% RPPFD for both species.

Table 2 also shows values determined from the pot experiments, which represented typical morphological and physiological traits in response to light and nitrogen availabilities. Values of LMA, P_{Leaf}, SAR and the the N_{area} - NAR relationship were used for following model predictions.

Model predictions

We simulated general trends of the effects of light availability and soil N availability on optimal L/R and N_{area} using the model. We used parameter values of M. bombycis (Table 1, 2) for the simulation because it becomes the basically same result even if the parameters of either species were used. SAR was changed within a realistic range, from 0.0005 to 0.005 gN g^{-1} d^{-1}. As described in the model description, we can simulate L/R and corresponding N_{area} and RGR uniquely by changing P_{Shoot} for the given parameters. Figure 3 shows the relationship between N_{area} and relative growth rate (RGR; g g^{-1} d^{-1}). For a given SAR, the optimal N_{area} that maximized RGR was obtained for both high- (Fig. 3A) and low- (Fig. 3B) light environments. Optimal N_{area}, and the associated maximum RGR was higher for 100% RPPFD than 10% RPPFD when compared with the same nitrogen availability, SAR.

The relationships between L/R, RGR and N_{area} are shown in Figure 4. Smaller L/R (larger root fraction) increased N_{area}, but too high N_{area} which was due to lower amount of photosynthetic organs (leaves) lead a decrease in NAR (Fig. 2) because increase in

| Parameters | M. bombycis | A. buergerianum |
|------------|-------------|----------------|
| a, a1, a2 | -0.0014 | -0.0013 |
| b, b1, b2 | 0.083 | 0.1045 |
| b, b1, b2 | 0.17 | 0.169 |
| c1, c2 | 0.71 | 0.51 |
| c1, c2 | 0.0083 | 0.0169 |
| c1, c2 | 0.375 | 0.0145 |
| c1, c2 | 0.51 | 0.65 |
| c1, c2 | 0.64 | 0.65 |
| c1, c2 | 0.153 | 0.62 |
| c1, c2 | 0.79 | 0.83 |
| c1, c2 | 0.73 | 0.83 |
| c1, c2 | 0.45 | 0.83 |
| c1, c2 | 0.0002 | 0.76 |
| c1, c2 | 0.73 | 0.76 |
| c1, c2 | 0.597 | 0.76 |
| c1, c2 | 0.0006 | 0.51 |
| c1, c2 | 0.85 | 0.51 |
| c1, c2 | 1.229 | 0.51 |
| c1, c2 | 0.019 | 0.51 |
| c1, c2 | 0.03 | 0.51 |
| c1, c2 | 0.091 | 0.51 |

Values of c1, c2, c3, and c4 were determined in each light environment. \( r^2 \) values represent determination coefficients of each parameter set.

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A max with N area is saturated, whereas increase in respiration rate (RD) with N area is linear (Fig. 1A,B). Consequently RGR also reduces when N area is too high. However, a larger L/R (smaller root fraction) also decreased RGR by decreasing N area and NAR (Fig. 2). Thus, optimal L/R and associated N area are determined by these balances, which change depending on light and nitrogen availability.

The effects of SAR on optimal L/R and N area differed between high- and low-light environments (Fig. 4 A,B). In a low-light environment, optimal L/R was higher and optimal N area was lower than those in the high-light environment, which can be interpreted as follows. Under a high-light environment, more biomass allocated to the roots increased nitrogen absorption resulting in high N area, NAR, and RGR. In contrast, in a low-light environment, due to saturation of NAR at low N area, the plant favoured a smaller fraction of root biomass, a large L/R, and a low N area to achieve maximum RGR.

Figure 5 shows the effects of changing SAR on optimal L/R and N area which give maximum RGR. Optimal L/R was always higher and optimal N area was always lower for 10% RPPFD for all ranges of SAR (Fig. 5). Optimal N area increased sharply with SAR for 100% RPPFD, whereas it was almost saturated at a low value for 10% RPPFD indicating difference in nitrogen demand between high- and low-light environments.

Comparison of actual biomass allocation with model predictions

The above predictions were tested using two deciduous pioneer tree species.
The pot experiment parameters showed morphological and physiological plasticity corresponding to the light environment and nitrogen availability (Table 2). LMA was higher for HR and HP (100% RPPFD) than for SR and SP (10% RPPFD), and SAR was higher for HR and SR (nitrogen-rich) than for HP and SP (nitrogen-poor) for both species. Constants for stem and root N concentration (eqn. M3, M4) differed between species and light environments, but each set of constants ($c_1$ and $c_2$, $c_3$ and $c_4$) showed high determination coefficients, as reported by Osone and Tateno (2003). The SAR of A. buergerianum was higher than that of M. bombycis, indicating different intrinsic capacities for nitrogen uptake [25,26].

Using these parameter values observed for each species (Table 1, 2) and the optimal biomass allocation model, we calculated optimal L/R and $N_{area}$ and compared these results with actual L/R and $N_{area}$ for each species. We set ranges of values for L/R and $N_{area}$ which cover 98% of the maximum RGR because the RGR curves against L/R and $N_{area}$ were gradual and maintained high RGR around optima, as shown in Figs. 3 and 4.

Measured L/R were higher and $N_{area}$ were lower in low-light than high-light environments, and all measured L/R and $N_{area}$ almost fell within the estimated ranges which cover 98% of the maximum RGR for both species (Fig. 6). The $N_{area}$ values for SR and SP were particularly close to the optimums for both species (Fig. 6 B, D). Thus, L/R and $N_{area}$ almost satisfied the model-predicted optimums for these pioneer trees growing both in high- and low-light environments.

**Discussion**

From the model predictions and pot experiments (Fig. 6), we could demonstrate that L/R and leaf $N_{area}$ of the pioneer tree

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**Table 2. Morphological and physiological parameters for material species.**

| Species         | Parameter | Definition                  | Units                  | HR       | HP       | SR       | SP       |
|-----------------|-----------|-----------------------------|------------------------|----------|----------|----------|----------|
| M. bombycis     | LMA       | leaf mass per area          | g m$^{-2}$             | 60.2     | 56.9     | 16.8     | 18.4     |
|                 | $N_{max}$ | leaf nitrogen content per mass | g N g$^{-1}$          | 0.026    | 0.017    | 0.046    | 0.029    |
|                 | $P_{leaf}$| fraction of leaf biomass in shoot biomass | -                     | 0.44     | 0.44     | 0.39     | 0.33     |
|                 | SAR       | specific absorption rate    | gN g$^{-1}$ d$^{-1}$   | 0.00104  | 0.00023  | 0.00142  | 0.00024  |
|                 | $A_{max}$ | maximum photosynthetic rate | $\mu$ mol m$^{-2}$ s$^{-1}$ | 7.86     | 5.86     | 5.21     | 4.33     |
| A. buergerianum | LMA       | leaf mass per area          | g m$^{-2}$             | 42.9     | 39.8     | 23.4     | 23.1     |
|                 | $N_{max}$ | leaf nitrogen content per mass | g N g$^{-1}$          | 0.038    | 0.025    | 0.034    | 0.028    |
|                 | $P_{leaf}$| fraction of leaf biomass in shoot biomass | -                     | 0.53     | 0.45     | 0.51     | 0.45     |
|                 | SAR       | specific absorption rate    | gN g$^{-1}$ d$^{-1}$   | 0.00342  | 0.00107  | 0.00124  | 0.00077  |
|                 | $A_{max}$ | maximum photosynthetic rate | $\mu$ mol m$^{-2}$ s$^{-1}$ | 7.45     | 3.93     | 3.66     | 3.31     |
|                 | RGR       | relative growth rate        | g g$^{-1}$ d$^{-1}$    | 0.0261   | 0.0143   | 0.01495  | 0.0103   |

Values of LMA, $P_{leaf}$, and SAR, were used for the model simulations. Values are shown for each species and treatment group.
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![Figure 3. Changes in the relative growth rate (RGR) with increasing leaf nitrogen content ($N_{area}$) when SAR was changed.](https://doi.org/10.1371/journal.pone.0022236.g003)
species, *M. bombycis* and *A. buergerianum*, were explained as optimal biomass allocation to maximize RGR in both high and low-light environments. Thus, we conclude that the balance between leaves and roots are regulated by nitrogen demand and nitrogen availability, each of which was dependent on light and soil nitrogen availability respectively. This is interpreted as follows: Plants need less nitrogen and fewer roots which absorb nitrogen to maximize growth rate when light availability was lower, because too high $N_{area}$ rather decrease NAR (Fig. 2) and consequently RGR (Fig. 3B, 4B). Conversely, plants need more nitrogen and more roots when light availability is high, because NAR plateau at higher $N_{area}$ (Fig. 3A, 4A). In addition to nitrogen demand, nitrogen availability expressed as SAR also plays an important role to determine optimal L/R and $N_{area}$ (Fig. 5) as demonstrated in classical studies [7,27]. It is especially notable that light availability determines the biomass allocation pattern through the change in nitrogen demand (Fig. 5).

The results obtained in the high-light environment were consistent with previous research [7,27], whereas this is the first study to theoretically and experimentally explain optimal biomass allocation of plants in high and low-light environments systematically (Figs. 5 and 6). Higher L/R and lower $N_{area}$ were also observed in many previous studies both in experimental fields and natural conditions [27,28]. Thus, these observations would be also understood as the optimal biomass allocation to maximize RGR.

In our model, we used morphological and physiological properties such as LMA, $P_{leaf}$, $N_{area}$ - NAR relationship, and SAR which were actually observed in response to light and nitrogen availabilities in the present study. This enables more quantitative predictions of optimal biomass allocation compared with the previous models [29–31]. Furthermore, our model can predict the optimal L/R and leaf $N_{area}$ in various light environments. This is mainly because we used light intensity measured throughout the growth period and incorporated the temperature dependency of photosynthetic rate and the dark respiration rate to estimate leaf productivity (Fig. 2). These characteristics of our model must improve the accuracy of the predictions compared with previous studies where leaf or canopy productivity was estimated by simpler models (e.g. [31–34]). In these simple models, leaf or canopy productivity was estimated by simplifying light availability as a sine square curve and leaf temperature was not also considered. However, actual light availability varies depending on weather conditions, forest structure, and seasonality [35]. Climate condition also affects leaf temperature expressed as functions of PPFD and air temperature, which mainly affects dark respiration rate exponentially (eqn. 4). In fact, NAR estimated by the simple model using the sine square curve was rather higher than that of the present study in which actual climate condition is considered. Therefore, optimal values predicted by using these NAR were deviated from those predicted by SAR of the present study especially in low-light environments. Therefore, these parameters should be considered to estimate accurate long-term leaf productivity [20] and to predict the optimal L/R and $N_{area}$ in our study.

Another important finding was that the extent of nitrogen limitation was smaller in the low-light environment than the high-light environment. Specifically, differences in L/R, $N_{area}$ and RGR between SR and SP were smaller than those between HR

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**Figure 4.** Effects of the leaf-to-root ratio (L/R) on relative growth rate (RGR) and leaf nitrogen content ($N_{area}$). (A) 100% photosynthetic photon flux density (100%RPPFD). (B) 10%RPPFD. Solid lines represent RGR and dashed lines represent $N_{area}$ respectively. Thick lines represent nitrogen absorption rates per unit root mass (SAR) = 0.005 and thin lines represent SAR = 0.0005. Black circles represent the maximum relative growth rate (RGR) and white circles represent the associated $N_{area}$. Parameter values obtained from *Morus Bombycis* were used (Table 2).

**Figure 5.** Optimal leaf-to-root ratio (L/R) and optimal leaf nitrogen content ($N_{area}$). Thick lines represent 100% photosynthetic photon flux density (100%RPPFD) and thin lines represent 10%RPPFD. Solid lines represent optimal L/R and dashed lines represent optimal $N_{area}$ respectively.

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Table 2. Parameter values obtained from *Morus Bombycis*.
and HP (Fig. 6, Table 2). This was probably due to less nitrogen demand of the plants to achieve maximum NAR and RGR in the low-light environment (Figs. 2, 3, and 5). Although the present study only focused on the pioneer trees, other shade-tolerant tree species also have low leaf \( N_{\text{area}} \) with lower L/R \([9,10,36-38]\). Thus, the extent of growth limitation caused by low nitrogen availability would be smaller in understory vegetation than in gap sites for any plants because of the difference in nitrogen demand. Furthermore, these facts indicate that not only pioneer trees with indeterminate growth but also shade-tolerant trees with determinate growth might achieve the optimal L/R. Our present model is not suitable for evaluating those shade-tolerant species because these species tend to produce leaves once a year and tougher stems and roots throughout a year. These organs live longer and reserve more carbohydrate and nitrogen for next year in contrast to pioneer species, and these traits are considered to be related to their survival \([9,39]\). Thus, elucidating optimality of shade-tolerant species is remaining as an interesting topic future research.

To understand plant biomass allocation strategy from a more realistic and versatile perspective, two major remaining subjects should be investigated. One is water absorption capacity of root, which contributes to growth through transpiration during photosynthesis and to survival by preventing from drying. Thus, there must be minimum requirements of root mass to leaves or whole-plant biomass, and it will change according to light environments because transpiration rate generally change in response to light intensity. From this viewpoint, it is predicted that plants with too high L/R would suffer from water stress and accordingly their RGR would rather decrease. Consequently, the ranges covering 98% of maximum RGR (Fig. 4) would be confined to narrower regions.

The other is stem mechanical constraint which causes increase in stem mass fraction. Especially in a low-light environment, many researchers have reported that the stem mass ratio of pioneer trees increases due to stem elongation and increase in specific stem length \([2,37,40,41]\). Although we could not find clear differences in \( P_{\text{Leaf}} \) between treatment groups in either species in the pot experiments, this would partly due to the shorter growth period (about 60 to 100 days) than in previous studies. In the year following this study, \( P_{\text{Leaf}} \) was lower in the low-light than high-light environment for \( M. \) bombycis (data not shown). Since the amount of biomass allocation to the stem should increase to maintain mechanical stability as plants grow higher \([42,43]\), stem elongation would become more and more costly especially for pioneer tree species in a shaded condition due to lower leaf productivity (Fig. 2).

Developing a model considering above-mentioned constraints combined with our present biomass allocation model would be helpful for further understanding of the plant biomass allocation strategy in various light environments.

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**References**

1. Givnish TJ (1988) Adaptation to sun and shade: a whole-plant perspective. Australian Journal of Plant Physiology 15: 63–92.

2. Poser R, Nagel O, Anderson JM, van Bel AJE, Knoblauch M, et al. (2000) The role of biomass allocation in the growth response of plants to different levels of light. CO. Aust. J. Plant Physiol 27: 595–607.

3. Iwasa Y, Roughgarden J (1984) Shoot/root balance of plants: Optimal growth of a system with many vegetative organs. Theoretical Population Biology 25: 78–105.

4. Garmier E (1991) Resource capture, biomass allocation and growth in herbaceous plants. Trends in Ecology & Evolution 6: 126–131.

5. McCarthy MC, Enquist BJ (2007) Consistency between an allometric approach and optimal partitioning theory in global patterns of plant biomass allocation. Ecology 21: 713–720.

6. Shipley B, Meziane D (2002) The balanced-growth hypothesis and the allometry of leaf and root biomass allocation. Functional Ecology 16: 326–331.

7. Osone Y, Tateno M (2003) Effects of stem fraction on the optimization of biomass allocation and maximum photosynthetic capacity. Functional Ecology 17: 627–636.

8. Hilbert DW (1990) Optimization of plant root: shoot ratios and internal nitrogen concentration. Annals of Botany 66: 91–99.

9. Kitajima K (1994) Relative importance of photosynthetic traits and allocation patterns as correlates of seedling shade tolerance of 13 tropical trees. Oecologia 98: 419–428.

10. Walters MB, Kruger EL, Reich PB (1993b) Growth, biomass distribution and CO 2 exchange of northern hardwood seedlings in high and low light: relationships with successional status and shade tolerance. Oecologia 94: 7–16.

11. Walters MB, Reich PB (2000) Seed size, nitrogen supply, and growth rate affect tree seedling survival in deep shade. Ecology 81: 1807–1901.

12. Valladares F, Wright SJ, Lasso E, Kitajima K, Pearcy RW (2000) Plastic phenotypic response to light of 16 congeneric shrubs from a Panamanian rainforest. Ecology 81: 1923–1936.

13. Sack L, Grubb BJ (2002) The combined impacts of deep shade and drought on the growth and biomass allocation of shade-tolerant woody seedlings. Oecologia 131: 175–185.

14. Walters MB, Kruger EL, Reich PB (1993a) Relative growth rate in relation to physiological and morphological traits for northern hardwood tree seedlings: species, light environment and ontogenetic considerations. Oecologia 96: 219–231.

15. Field C, Mooney HA (1986) The photosynthesis-nitrogen relationship in wild plants. In: Givnish TJ, ed. On the economy of plant form and function. Harvard Forest: Cambridge University Press. pp 28–49.

16. Hirose T, Kitajima K (1996) Nitrogen uptake and plant growth I. Effect of nitrogen removal on growth of Polygonum cuspidatum. Annals of Botany 59: 479–496.

17. Hikosaka K, Ishikawa K, Borjigidai A, Muller O, Onoda Y (2006) Temperature acclimation of photosynthesis: mechanisms involved in the changes in temperature dependence of photosynthetic rate. Journal of experimental botany 57: 291–302.

18. Farquhar GD, Caemmerer S, Berry JA (1980) A biochemical model of photosynthetic CO 2 assimilation in leaves of C 3 species. Planta 149: 78–90.

19. Arkin OK, Evans JR, Ball MC, Lamberts H, Pons TL (2000) Leaf respiration of snow gum in the light and dark. Interactions between temperature and irradiance. Plant Physiology 122: 915–924.

20. Oguchi R, Hikosaka K, Hira T, Hirose T (2008) Costs and benefits of photosynthetic light acclimation by tree seedlings in response to gap formation. Oecologia 155: 665–675.

21. Sato M, Ohyanma K (1976) Studies on photosynthesis and translocation of photosynthetic in mulberry tree, 3: Utilization of reserve substance in the process of regrowth after shoot pruning in a growing season. Proceedings of the Crop Science Society of Japan 45: 51–56.

22. Murakami T (1983) Studies on the reserve substrate and photosynthesis related to the early growth in cutting propagation of mulberry plant. Bulletin Sericultural Experiment Station Ministry of Agriculture, Forestry and Fisheries 29: 1–164.

23. Tsutsumi T (1989) Forest ecology. Asakura System: Tokyo. 68 p.

24. Chapin III FS (1980) The mineral nutrition of wild plants. Annual review of ecology and systematics 11: 253–260.

25. Reich PB, Walters MR, Tjoelker MG, Vanderklein D, Buschena C (1998b) Photosynthesis and respiration rates depend on leaf and root morphology and nitrogen concentration in nine boreal tree species differing in relative growth rate. Functional Ecology 12: 393–403.

26. Osone Y, Tateno M (2003) Nitrogen absorption by roots as a cause of interspecific variations in leaf nitrogen concentration and photosynthetic capacity. Functional Ecology 19: 460–470.

27. Schreng LS, Koh R, Walters MB (2005) Tree seedling growth, survival, and morphology in response to landscape-level variation in soil resource availability in northern Michigan. Canadian Journal of Forest Research 35: 263–273.

28. Seiwa K (2007) Trade-offs between seedling growth and survival in deciduous broadleaved trees in a temperate forest. Annals of Botany 99: 537–544.

29. Aşen GI, Ingestad T (1987) Root: shoot ratio as a balance between nitrogen productivity and photosynthesis. Plant Cell Environment 10: 579–586.

30. Gleeson SK (1993) Optimization of tissue nitrogen and root-shoot allocation. Annals of Botany 71: 23–31.

31. Hilbert DW, Reynolds JE (1991) A model allocating growth among leaf proteins, shoot structure, and root biomass to produce balanced activity. Annals of Botany 68: 417–425.

32. Hirose T, Weger MJA (1987) Maximizing daily canopy photosynthesis with respect to the leaf nitrogen allocation pattern in the canopy. Oecologia 72: 520–526.

33. Hikosaka K, Terashima I (1993) A model of the acclimation of photosynthesis in the leaves of C 3 plants to sun and shade with respect to nitrogen use. Plant Cell and Environment 16: 605–605.

34. Hikosaka K, Sudo S, Hirose T (1999) Light acquisition and use by individuals competing in a dense stand of an annual herb, Xanthium canadense. Oecologia 119: 388–396.

35. Ida TY, Kudo G (2008) Timing of Canopy Closure Influences Carbon Translocation and Seed Production of an Understorey Herb, Trillium apetalon (Trilliaceae). Annals of Botany 101: 435–446.

36. Ellsworth DS, Reich PB (1992) Leaf mass per area, nitrogen content and photosynthetic carbon gain in Acer saccharum seedlings in contrasting forest light environments. Functional Ecology 6: 423–435.

37. Walters MB, Reich PB (1996) Are shade tolerance, survival, and growth linked? Low light and nitrogen effects on hardwood seedlings. Ecology 77: 841–853.

38. Niinemets U (1997) Role of foliar nitrogen in light harvesting and shade tolerance of four temperate deciduous woody species. Journal of Ecology 11: 510–531.

39. Myers JA, Kitajima K (2007) Carbohydrate storage enhances seedling shade and stress tolerance in a neotropical forest. Ecology 95: 383–395.

40. Poorter R, Nagel O, Anderson JM, van Bel AJE, Knoblauch M, et al. (2000) The role of biomass allocation in the growth response of plants to different levels of light. CO. Aust. J. Plant Physiol 27: 595–607.

41. Hikosaka K, Ishikawa K, Borjigidai A, Muller O, Onoda Y (2006) Temperature acclimation of photosynthesis: mechanisms involved in the changes in temperature dependence of photosynthetic rate. Journal of experimental botany 57: 291–302.

42. Farquhar GD, Caemmerer S, Berry JA (1980) A biochemical model of photosynthetic CO 2 assimilation in leaves of C 3 species. Planta 149: 78–90.

43. Arkin OK, Evans JR, Ball MC, Lamberts H, Pons TL (2008) Leaf respiration of snow gum in the light and dark. Interactions between temperature and irradiance. Plant Physiology 122: 915–924.

44. Oguchi R, Hikosaka K, Hira T, Hirose T (2008) Costs and benefits of photosynthetic light acclimation by tree seedlings in response to gap formation. Oecologia 155: 665–675.

45. Satoh M, Ohyanma K (1976) Studies on photosynthesis and translocation of photosynthetic in mulberry tree, 3: Utilization of reserve substance in the process of regrowth after shoot pruning in a growing season. Proceedings of the Crop Science Society of Japan 45: 51–56.

**Author Contributions**

Conceived and designed the experiments: DS MT. Performed the experiments: DS MT. Analyzed the data: DS MT. Contributed reagents/materials/analysis tools: DS MT. Wrote the paper: DS.