Predictability of leaf traits with climate and elevation: a case study in Gongga Mountain, China

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Leaf mass per area (\(M_a\)), nitrogen content per unit leaf area (\(N_{area}\)), maximum carboxylation capacity (\(V_{cmax}\)) and the ratio of leaf-internal to ambient CO\(_2\) partial pressure (\(\chi\)) are important traits related to photosynthetic function, and they show systematic variation along climatic and elevational gradients. Separating the effects of air pressure and climate along elevational gradients is challenging due to the covariation of elevation, pressure and climate. However, recently developed models based on optimality theory offer an independent way to predict leaf traits and thus to separate the contributions of different controls. We apply optimality theory to predict variation in leaf traits across 18 sites in the Gongga Mountain region. We show that the models explain 59% of trait variability on average, without site- or region-specific calibration. Temperature, photosynthetically active radiation, vapor pressure deficit, soil moisture and growing season length are all necessary to explain the observed patterns. The direct effect of air pressure is shown to have a relatively minor impact. These findings contribute to a growing body of research indicating that leaf-level traits vary with the physical environment in predictable ways, suggesting a promising direction for the improvement of terrestrial ecosystem models.

**Keywords:** deciduous LMA prediction, elevation gradients, leaf functional traits, leaf nitrogen prediction, optimality-based models, trait–climate relationships.

**Introduction**

A number of leaf traits are diagnostic of photosynthetic processes. The ratio of leaf-internal to external CO\(_2\) (\(\chi\)) reflects the stomatal regulation of CO\(_2\) uptake, which has to be balanced against water loss (Wang et al. 2017b). The maintenance of transpiration involves a carbon cost, in the form of respiration by living parenchyma cells, to maintain active water transport tissues. The maximum capacity of carboxylation at a standard temperature of 25 \(^\circ\)C, \(V_{cmax25}\), is a measure of the control of photosynthesis by the amount of the enzyme (Rubisco) responsible for carbon fixation (Wang et al. 2020). The maintenance of photosynthetic capacity also incurs a substantial carbon cost in the form of leaf respiration to support protein synthesis. Leaf mass per unit area (\(M_a\)) determines the total carbon cost of leaf construction (Wright et al. 2004). Nitrogen is required for both metabolic processes and leaf construction (Lambers and Poorter 1992, Onoda et al. 2004). Leaf nitrogen content per unit area (\(N_{area}\)) thus provides a combined measure of the metabolic and structural costs.

Empirical analyses of large trait data sets have shown that variation in each of these traits is related to climate, and indeed specific climate variables can be shown to influence individual...
impacts of climate and elevation on photosynthesis. Optimality theory is predicated on the idea that through evolutionary processes (including selection for plasticity as well as environmental filtering of lineages) plants are adapted to the environmental conditions under which they live. The values of photosynthetic parameters are then predicted as the result of trade-offs between competing requirements, such as the need to balance CO₂ uptake against water loss. The balance between maintaining carboxylation capacity and transpiration capacity can be described in terms of the least-cost hypothesis (Wright et al. 2003, Prentice et al. 2014), which states that plants minimize the combined costs of maintaining these capacities. This hypothesis allows us to predict $\chi$. The coordination hypothesis (Chen and Reynolds 1997, Maire et al. 2012, Wang et al. 2017b) indicates that carbon gain is maximized through balancing light and Rubisco limitations on photosynthesis. This hypothesis allows us to predict $V_{\text{cmax25}}$ (Smith et al. 2019). The need to allocate nitrogen to structural and metabolic processes allows us to predict $N_{\text{area}}$ as a function of $V_{\text{cmax25}}$ and $M_a$ (Dong et al. 2017). According to the optimal leaf longevity (LL) hypothesis (Kikuzawa 1991), plants maximize the time-averaged net carbon gain of leaves, taking into account the construction costs (amortized over the leaf lifetime) and the decline in photosynthetic capacity with increasing age. This hypothesis allows $M_a$ to be predicted from LL. The LL of deciduous species is constrained by growing season length (gsl); thus, $M_a$ of deciduous species should be predictable from gsl.

In this study, we draw on these theoretical developments to predict trait variability in response to climate and elevation gradients in the Gongga Mountain region, China. We develop a new optimality model to predict $M_a$ of deciduous species and a simplified optimality approach to predict $N_{\text{area}}$. These optimality models were developed independently of the observations used in this study and require no calibration. We show that these models capture observed variations in photosynthetic traits at sites in the Gongga Mountain region. We then use these models to quantify the relative contribution of different factors to the observed changes in trait values at these sites.

**Materials and methods**

**Study sites**

We collected photosynthetic trait data from 18 sites in the Gongga Mountain region of Sichuan Province, China (Figure 1a and b). The study area extends from 29° 22’ to 29° 55’ N and from 101° 1’ to 102° 9’ E. The sampled sites span an elevation gradient from 1143 to 4361 m, and as a result, there is a considerable gradient in growing season temperature (see Table S1 available as Supplementary data at Tree Physiology online). Sites from the western part of the Gongga Mountain region tend to be drier than the sites at a corresponding elevation in the eastern part, and thus, our data set also
samples a large moisture gradient (see Table S1 available as Supplementary data at Tree Physiology online). The vegetation at lower elevations is deciduous broad-leaved forest dominated by Betulaceae, Urticaceae, Caprifoliaceae and Rosaceae, and it is replaced by evergreen needle-leaved forest and subsequently by deciduous shrubland dominated by Pinaceae and/or Rosaceae and Ericaceae (see Table S2 available as Supplementary data at Tree Physiology online) with increasing elevation. Although evergreen woody species are present at all of the sites (see Table S2 available as Supplementary data at Tree Physiology online), and trait measurements were made on these species, our subsequent analyses of photosynthetic traits focused entirely on the deciduous species because of the difficulty of obtaining reliable estimates of leaf age based on a single sampling of a site.

Sample collection and analysis
Trait data were measured in late July 2018 and early August 2019 during the active growing season in the Gongga Mountain region. We used a stratified sampling strategy at each site in order to sample the dominant species in each canopy stratum. In forest sites, we sampled a minimum of five tree, five shrub and five forb species at each site and also sampled graminoids, lianas and climbers, and pteridophytes when present (see Table S2 available as Supplementary data at Tree Physiology online). At the highest elevation sites (3794, 3943, 4081 and 4361 m), where shrubs form the upper canopy, we only sampled shrubs and forbs (and at the highest site one pteridophyte), but again, we sampled a minimum of five species in each category. All samples were taken from the outer canopy.

Measurements were made on the young but fully expanded leaves attached to the cut branch.

Our analyses focus on four leaf traits: (i) leaf mass per unit area (\( M_a \, \text{g biomass m}^{-2} \)), (ii) the maximum capacity of carboxylation at 25 °C (\( V_{c,max25} \, \mu \text{mol C m}^{-2} \text{s}^{-1} \)), (iii) the ratio of leaf-internal to ambient CO\(_2\) partial pressure (\( \chi \), unitless) and (iv) leaf nitrogen content per unit area (\( N_{\text{area}} \, \text{g m}^{-2} \)). (see Table 1 for definitions of parameters and other abbreviations frequently used in the text.) The \( M_a \) was obtained from the measurements of leaf area and dry weight following standard protocols (Cornelissen et al. 2003). Leaf area was taken as the projected area of a leaf, or leaflet for compound leaves, using a LiDE 220 Scanner (Canon Inc., Huntington, NY, USA). Dry weight was obtained after air-drying for several days and then after oven-drying at 75 °C for 48 h to constant weight. The \( M_a \) value of one species at each site was the average of three measurements made on leaves from multiple individuals. Leaf nitrogen content was measured using an Isotope Ratio Mass Spectrometer (Thermo Fisher Scientific Inc., Carlsbad, CA, USA). The \( N_{\text{area}} \) was calculated from \( M_a \) and leaf nitrogen content. Leaf nitrogen content (for \( N_{\text{area}} \)) and stable carbon isotope (\( \delta^{13} \)C, for \( \chi \)) measurements were made on pooled samples of leaves from multiple individuals.

We used a portable infrared gas analyzer system (LI-6400; Li-Cor Inc., Lincoln, NE, USA) to make the leaf gas-exchange measurements. Sunlit terminal branches from the upper canopy were collected and re-cut under water immediately prior to measurement. Measurements were made in the field with relative humidity and chamber block temperature close to that of the
Table 1. Parameters and abbreviations frequently used in the text. The table provides information on the meaning and units.

| Parameters and abbreviations | Unit | Description |
|------------------------------|------|-------------|
| $\chi$                       | Pa Pa$^{-1}$ | The ratio of leaf-internal to ambient CO$_2$ partial pressures |
| $M_a$                        | g biomass m$^{-2}$ | Leaf mass per area |
| $N_{area}$                   | g m$^{-2}$ | Leaf nitrogen content per area |
| $V_{cmax}$                   | $\mu$molC m$^{-2}$ s$^{-1}$ | The maximum capacity of carboxylation |
| $V_{cmax25}$                 | $\mu$molC m$^{-2}$ s$^{-1}$ molC m$^{-2}$ day$^{-1}$ | The maximum capacity of carboxylation at standard 25 $^\circ$C |
| $T_g$                        | $^\circ$C | Mean temperature during the growing season (mean daily temperature above a baseline of 0 $^\circ$C) |
| $D_0$                        | kPa | Mean vapor pressure deficit during the growing season |
| $R_0$                        | $\mu$mol photon m$^{-2}$ s$^{-1}$ | Mean photosynthetically active radiation during the growing season |
| $R_{LAI}$                    | mol photon m$^{-2}$ day$^{-1}$ | Mean leaf-area-index-weighted photosynthetically active radiation during the growing season |
| $f$                          | day day$^{-1}$ | The ratio of gsl to the number of days in the year |
| MAP                          | mm | Mean annual precipitation |
| $\alpha_p$                  | mm mm$^{-1}$ | The ratio of annual actual evapotranspiration to annual potential evapotranspiration |
| $T_{dl}$                     | $^\circ$C | Mean daytime temperature of July |
| $\Gamma^+$                   | Pa | The photosynthetic compensation point |
| $c_a$                        | Pa | Ambient CO$_2$ partial pressure |
| $c_i$                        | Pa | Internal CO$_2$ partial pressure |
| $\beta$                     | unitless | The ratio at 25 $^\circ$C of the unit costs of maintaining carboxylation and transpiration capacities (estimated as 146) |
| $K$                          | Pa | The effective Michaelis–Menten coefficient of Rubisco |
| $K_C$                       | Pa | The Michaelis–Menten coefficients of Rubisco for carboxylation |
| $c$                          | unitless | A constant proportional to the unit carbon cost for the maintenance of electron transport capacity (0.41) |
| LL                           | day | Leaf longevity |
| $b$                          | day | The potential age when leaves can no longer photosynthesize and assimilate CO$_2$ |
| $k$                          | g biomass mol C$^{-1}$ | Scaling factor |
| $I_{obs}$                    | mol photon m$^{-2}$ day$^{-1}$ | The photosynthetically active radiation absorbed by leaves |
| CC                           | gC gC$^{-1}$ | A constant representing the construction carbon cost per unit leaf mass carbon |
| $A_a$                        | g biomass m$^{-2}$ day$^{-1}$ | Daily carbon assimilation rate per unit leaf area |
| $\varphi_0$                 | $\mu$mol C $\mu$mol$^{-1}$ photon | The intrinsic quantum efficiency of photosynthesis |
| $N_{rubisco}$                | g m$^{-2}$ | Nitrogen content in Rubisco enzymes |
| $N_{structure}$              | g m$^{-2}$ | Nitrogen content in leaf structure |

\[
\chi = \frac{\Delta + f' \Gamma^+}{c_a} - a_s - \frac{a_s}{b'} - a_s,
\]

where $a_s$, $b'$, and $f'$ are the fractionations associated with diffusion in air (4.4‰), Rubisco carboxylation (30‰) and photorespiration (16‰), respectively. $\Gamma^+$ is the photosynthetic compensation point and $c_a$ is the ambient CO$_2$ partial pressure.

**Climate data**

In situ climate data were only available for five (1785, 2782, 2993, 3251 and 3943 m) of the 18 sampled sites. We therefore estimated the climate at each site consistently by interpolation between a larger set of weather stations in the region (17 stations, see Figure S1 available as Supplementary.
data at *Tree Physiology* online) for the period from January 2017 to December 2019 (http://data.cma.cn/data/ccddetail/dataCode/SURF_CLI_CHN_MUL_MON.html) to create seasonal climatologies of monthly maximum and minimum temperatures, fraction of sunshine hours, water vapor pressure and precipitation. These 17 stations range in elevation from 422 to 3951 m. We then used the elevationally sensitive ANUSPLIN interpolation scheme (Hutchinson and Xu 2004) to provide estimates of these meteorological variables at each of the sites. The monthly estimates at each site were converted to daily values by linear interpolation in order to calculate the bioclimatic estimates of these meteorological variables at each of the sites. We also calculated the ratio of gsl to the number of days in the year (f) and the leaf-area-index-weighted R0 (R40) to represent the effect of light interception by different layers in the canopy (Dong et al. 2017). The average leaf area index during July and August (i.e., the months the trait data were collected) in 2018 and 2019 was derived from the MODIS leaf area index product (MCD15A3H: https://modis.gsfc.nasa.gov/). An annual moisture index (\( \alpha_p \), an estimate of the ratio of annual actual evapotranspiration to annual potential evapotranspiration) was calculated from the monthly temperature, precipitation and fraction of sunshine hours at each site using the simple processed algorithms for simulating habitats (SPASH) model (Davis et al. 2017). Given the large difference between daytime and night-time temperatures at high elevations, we also calculated the mean daytime temperature of July (\( T_{dJ} \)) by approximating the daily temperature cycle with a sine curve:

\[
T_{dJ} = T_{\text{max}} \left\{ \frac{1}{2} + \frac{(1 - x^2)^{1/2}}{2 \cos^{-1}(1-x)} \right\} + T_{\text{min}} \left\{ \frac{1}{2} - \frac{(1 - x^2)^{1/2}}{2 \cos^{-1}(1-x)} \right\},
\]  

(2)

where \( T_{\text{max}} \) is the mean daily maximum air temperature, \( T_{\text{min}} \) is the mean daily minimum air temperature and \( x = \tan \phi \tan \delta \), where \( \phi \) is site latitude and \( \delta \) is the average solar declination in July.

Comparison of the interpolated bioclimatic variables with the values calculated using in situ data at the five sites where such data are available (see Figure S2 available as Supplementary data at *Tree Physiology* online) suggests that the ANUSPLIN interpolation provides robust estimates of the patterns of variation in climate across sites although, except for July temperature, the absolute values differ.

**Trait data analysis**

Analyses of the trait data focused on the predominant deciduous component of each community. We used redundancy analysis (RDA: Legendre and Legendre 2012) to determine the main patterns of trait variation using species average values from each site, assess how much of this variation is explained by environmental factors and determine the correlations between traits and environment. The RDA was performed using the vegan package in R (Oksanen et al. 2017). In order to compare the trait variability within and across sites, we calculated the coefficient of variation (CV: Brown 1998), a standardized measure of the dispersion of a frequency distribution, for the data set as a whole and at each site, for each of the traits independently.

We used generalized additive models (GAMs) to analyze trait variability with \( \alpha_p \) and elevation. The GAMs (Hastie and Tibshirani 1990) allow flexible relationships between response and predictor variables to be fitted to the data, avoiding the need to assume the form of the function in advance. Convex hulls were used to exclude areas of the fitted surface that were not well constrained by observations. The GAMs were fitted using the mgcv package (Wood 2001), and \( \alpha \)-convex hull was produced using alphahull package in R (Rodriguez Casal and Pateiro López 2010).

**Trait prediction**

We used existing optimality based models of \( \chi \) and \( V_{cmax25} \) and new models of \( M_a \) and \( N_{area} \) to predict the distribution of traits with climate and elevation across the sites. We used gsl as a proxy for the LL of deciduous plants. Specific photosynthetic traits adjust to the environmental conditions over different timeframes (Xu and Baldocchi 2003, Jiang et al. 2020), so we tried two alternative measures of temperature (\( T_g \) and \( T_{dJ} \)) as predictors. The models for \( \chi \) and \( V_{cmax25} \) apply for both deciduous and evergreen species.

**The model for \( \chi \)** This model is based on the assumption of evolutionary optimality in the trade-off between the costs of transpiration and carbon gain. The least-cost hypothesis predicts that plants minimize the total costs of photosynthesis, i.e., the requirement to maintain capacities for both carboxylation and transpiration (Wright et al. 2003, Prentice et al. 2014). Using the standard photosynthesis model due to Farquhar et al. (1980), Wang et al. (2017b) showed that \( \chi \) could be predicted by:

\[
\chi = \frac{\Gamma^*}{c_a} + \frac{\xi (1 - \Gamma^*/c_a)}{\xi + \sqrt{\Delta_0}},
\]  

(3)

where

\[
\xi = \sqrt{\frac{b(K + \Gamma^*)}{1.6 \eta}},
\]  

(4)

and

\[
K = K_c \left( 1 + \frac{P_o}{K_o} \right).
\]  

(5)
Here $\Gamma^*$ is the photorespiratory compensation point, and $c_a$ is the ambient CO$_2$ partial pressure. The $\eta$ is the viscosity of water relative to its value at 25 °C. The $\beta$ is the ratio at 25 °C of the unit costs of maintaining carboxylation and transpiration capacities. Based on a global compilation of leaf $^{13}$C measurements, Wang et al. (2017b) estimated $\beta = 146$. The $K$ is the effective Michaelis–Menten coefficient of Rubisco. The $K_c$ and $K_o$ are the temperature-dependent Michaelis–Menten coefficients for carboxylation and oxygenation, with reference values at 25 °C of 39.97 Pa and 27.48 kPa, respectively (Bernacchi et al. 2001). The $P_o$ is the ambient partial pressure of O$_2$. The composite variable $\xi$ determines the sensitivity of $\chi$ to $D_0$. This dependence is influenced by temperature (via $\Gamma^*$, $K$ and $\eta$) and O$_2$ pressure (via $K$) according to Eqs (4) and (5).

### The model for $V_{cmax25}$

The coordination hypothesis states that plants coordinate light-limited and Rubisco-limited photosynthesis rates so as to be equal under average daytime conditions (Chen et al. 1993). This coordination ensures that the use of absorbed light is maximized without incurring additional maintenance costs for $V_{cmax}$. The $V_{cmax}$ acclimated to growth temperature can be predicted from the universal model of carbon uptake proposed by Wang et al. (2017b):

$$V_{cmax} = \varphi_0 R_o \left( \frac{c_i + K}{c_i + 2\Gamma^*} \right) \sqrt{1 - \left( \frac{c_v}{c_i} \right)^{0.5}},$$  \hspace{1cm} (6)

$$m = \left( \frac{\xi - \Gamma^*}{c_i + 2\Gamma^*} \right),$$  \hspace{1cm} (7)

where $\varphi_0$ is the intrinsic quantum efficiency of photosynthesis (0.085 $\mu$mol C $\mu$mol$^{-1}$ photon), and $c_i$ is the leaf-internal CO$_2$ partial pressure, which is the product of observed $\chi$ and $c_a$. The $c$ is a constant proportional to the unit carbon cost for the maintenance of electron transport capacity (a value of 0.41 was estimated from an independent global data set on photosynthetic capacities). The $m$ represents the effect of subsaturating CO$_2$ on the light-limited rate of photosynthesis.

The kinetic response of Rubisco to temperature allows $V_{cmax25}$ to be estimated from $V_{cmax}$ at growth temperature ($T_g$), by the following relationship:

$$V_{cmax} = V_{cmax25} f_v,$$  \hspace{1cm} (8)

$$f_v = e^{H_o (T_g - 298.15)/(298.15T_g R)} \times \left[ \frac{1 + e^{(298.15\Delta S - H_d)/(298.15R)}}{1 + e^{(T_g \Delta S - H_d)/(T_g R)}} \right],$$  \hspace{1cm} (9)

where $H_o$ is the activation energy (71,513 J mol$^{-1}$), $R$ is the universal gas constant (8.314 J mol$^{-1}$ K$^{-1}$), $H_d$ is the deactivation energy (200,000 J mol$^{-1}$) and $\Delta S$ is an entropy term (J mol$^{-1}$ K$^{-1}$) calculated using a linear relationship with $T_g$, with a slope of 1.07 J mol$^{-1}$ K$^{-2}$ and intercept of 668.39 J mol$^{-1}$ K$^{-1}$ (Kattge and Knorr 2007).

## A new model for $M_a$

The $M_a$ contributes to determining how much leaf area can be displayed for a given amount of carbon allocated to above-ground tissues (Cui et al. 2019). There is a universal trade-off between $M_a$ and LL across growth forms, plant functional types (PFTs) and biomes, known as the ‘leaf economics spectrum’ (Wright et al. 2004). The spectrum runs from a ‘fast’ to a ‘slow’ economic strategy. Plants adopting a fast economic strategy have rapid returns on investment (low $M_a$) and short longevity (low LL), while plants adopting the slow strategy have high $M_a$ and high LL.

Here we propose a novel model for $M_a$, which combines three optimality-based predictions. We start from the model proposed by Kikuzawa (1991). By assuming that the average net carbon gain by a leaf during its lifetime is maximized, this model provides an optimality-based prediction of the trade-off between $M_a$ and LL:

$$LL = \sqrt{\frac{2b \times CC \times M_a}{A_a}}.$$  \hspace{1cm} (10)

Here $b$ (day) is the potential age at which leaves can no longer photosynthesize, CC (gC g$^{-1}$) is the construction cost per unit mass of leaf carbon and $A_a$ (g biomass m$^{-2}$ day$^{-1}$) is the daily mass of leaf carbon and inversely proportional to LL across growth forms, plant functional types (PFTs) and biomes (Xu et al. 2017), and $k$ is a scaling factor (30 g biomass mol C$^{-1}$).

Third, the coordination hypothesis allows optimal values of $V_{cmax}$ to be predicted by equating the Rubisco-limited assimilation rate with the electron transport limited rate under typical daytime conditions that include temperature, vapor pressure deficit, ambient CO$_2$ and the photosynthetically active radiation absorbed by leaves ($I_{abs}$). The model has the mathematical form of a ‘light-use efficiency model’: that is, modeled total photosynthesis over any period is proportional to the total light absorbed during that period, which is consistent with classical studies on crop growth (Wang et al. 2017b). For this derivation, we made the simplifying assumption that the maximum rate of electron transport ($I_{max}$) is large enough that the square-root term in Eq. (6) can be neglected. We substituted Eqs (8) and (9) into (11) to predict $b$ from $M_a$ and $V_{cmax}$, which is then predictable from $\varphi_0$, $I_{abs}$, $c_i$, $\Gamma^*$ and $K$. In this way, we obtained
a theoretical prediction of $M_a$:

$$M_a = \varphi_0 \text{LL} \, k \, \sqrt{\frac{(c_1 - \Gamma^*) (c_1 + K)}{(2 \mu CC_f) (c_1 + 2 \Gamma^*)^2}}.$$ (12)

In addition to the implied proportionality of $M_a$ with both absorbed light and LL, Eq. (12) indicates the existence of a composite temperature effect due to the temperature dependencies of $\chi$, $\Gamma^*$, $K$ and $f_v$. In order to separate these dependencies, estimate the net effect of temperature more easily and account for the moisture effect detected in the China Plant Trait database (Wang et al. 2018), we obtained the partial derivative of $\ln(M_a)$ in Eq. (12) with respect to temperature ($T_g$) and evaluated the result under standard environmental conditions. This predicts a decline in $\ln(M_a)$, for a given LL and $I_{\text{abs}}$, of $\approx 3\%$ per degree increase in growth temperature ($T_g$). In addition, all the constants ($\varphi_0$, $u$, $k$, $CC$ and reference values of $f_v$, $K$, $c_1$ and $\Gamma^*$ at 25 $^\circ$C) are combined into a single parameter $C_1$ to reduce the complexity of the model. A linearized equation for predicted $M_a$ can then be derived as:

$$\ln(M_a) = \ln(I_{\text{abs}}) - 0.03 \, T_g + \ln(\text{LL}) + \ln(C_1),$$ (13)

where $C_1$ is a free parameter. For deciduous species, there is an additional constraint on LL by gsl in Eq. (13), thus we obtained the equation for deciduous species:

$$\ln(M_a) = \ln(I_{\text{abs}}) - 0.03 \, T_g + \ln(f) + \ln(C_2),$$ (14)

where $f$ is the ratio of gsl to the number of days in the year. Thus, information on the number of days in a year is considered in the free parameter ($\ln(C_2) = \ln(C_1) + \ln(365)$), resulting in changing of $C_1$ to $C_2$. The $C_1$ and $C_2$ are unknown a priori but could be estimated from observations.

Although not included in this theoretical derivation, a strong negative effect of increasing moisture availability on $M_a$ has been reported (Meng et al. 2015). We used the ratio of actual to potential evapotranspiration ($\alpha_p$) as an index of moisture availability in order to estimate this effect from the data. Thus, parameter $C_2$ is further replaced by $C_3$ to denote the parameter difference in Eq. (14) and Eq. (15) after the moisture effect is included.

We used an independent data set of $\ln(M_a)$ for 621 deciduous species from the China Plant Trait database (Wang et al. 2018) to estimate the parameter $C_3$. Using $R_{\text{AI}}$ to represent the averaged light absorbed by leaves, we regressed the observations of $\ln(M_a)$ against $\ln(R_{\text{AI}})$, $T_g$, $\ln(f)$ and $\ln(\alpha_p)$ and obtained an estimate of $\ln(C_3)$ of 1.70. The predictors in this analysis explained 53% of the variation in $M_a$, and the fitted slopes of $R_{\text{AI}}$, $T_g$ and $\ln(f)$ were quantitatively consistent with their theoretical values as given in Eq. (14). Thus, the final model for $M_a$ was:

$$\ln(M_a) = 1.22 \ln(R_{\text{AI}}) + 0.78 \ln(f) - 0.06 \, T_g - 0.60 \, \ln(\alpha_p) + 1.70.$$ (15)

A simple model for $N_{\text{area}}$ The $N_{\text{area}}$ represents the sum of nitrogen in both metabolic and structural components of a leaf. Dong et al. (2017) proposed a model to predict $N_{\text{area}}$ from $M_a$ and $V_{\text{cmax25}}$ by assuming (based on previously published analyses) that (i) $V_{\text{cmax25}}$ is proportional to nitrogen in Rubisco and (ii) non-photosynthetic nitrogen is almost proportional to $M_a$. The model of Dong et al. (2017) is as follows:

$$N_{\text{area}} = 9.5 N_{\text{rubisco}} + N_{\text{structure}},$$ (16)

$$N_{\text{structure}} = 10 - 2.67 M_a 0.99$$ (17)

and

$$N_{\text{rubisco}} = 0.003135 V_{\text{cmax25}}.$$ (18)

The coefficient of $N_{\text{rubisco}}$ in Eq. (16) reflects the allocation of total metabolic nitrogen to Rubisco, which however, varies among species. We used the observed $M_a$ and $V_{\text{cmax25}}$ in this study to estimate $N_{\text{structure}}$ and $N_{\text{rubisco}}$ in Eqs (17) and (18), then fitted a regression of metabolic nitrogen (estimated as the difference between $N_{\text{area}}$ and $N_{\text{structure}}$) against $N_{\text{rubisco}}$ to estimate this coefficient for the deciduous species from the Gongga sites. We obtained a value for the coefficient of $N_{\text{rubisco}}$ of 7.2, which is within the predicted range given in Dong et al. (2017).

However, there is considerable uncertainty in Eq. (18), which describes the maximal catalytic turnover rate of Rubisco at 25 $^\circ$C (von Caemmerer et al. 1994, Harrison et al. 2009) as well as in Eqs (16) and (17). To simplify the calculations and avoid these uncertainties, we adopted an alternative method to estimate $N_{\text{area}}$ directly by regression as a linear combination of all observed $M_a$ and $V_{\text{cmax25}}$ (without intercept) in this study, yielding a simpler model that applies to non-nitrogen-fixing plants:

$$N_{\text{area}} = 0.02 M_a + 0.003 V_{\text{cmax25}}.$$ (19)

We used this simple model to predict $N_{\text{area}}$ first from observed—and then from predicted—$V_{\text{cmax25}}$ and $M_a$. In this way, we could first test whether $N_{\text{area}}$ is indeed predictable from $V_{\text{cmax25}}$ and $M_a$ in our data set and then test whether $N_{\text{area}}$ is predictable from the climate data alone. In order to examine the impact of nitrogen fixation on this relationship, we also included ‘N-fixer’ as a factor in this linear model. Partial residuals from the regression model for $N_{\text{area}}$ were plotted using the visreg package (Breheny and Burchett 2017).
Estimating the contribution of individual predictor variables

The contribution of each predictor variable to trait variation was calculated in three steps. At Step 1, we created a baseline by averaging the values of each predictor variable across the 18 sites to create a data set for an ‘average’ site. We used this average site data to calculate baseline trait values. At Step 2, we changed one predictor variable at a time to the actual value at that site, keeping all the other variables constant at the average site value. We then calculated trait values using these new inputs. At Step 3, the contribution of each predictor variable was calculated as the difference between the traits simulated at Step 2 and the baseline value of the traits from Step 1. This procedure allowed us to separate out the individual influences of changes in air pressure with elevation, $T_d$ and $D_0$ on $\chi$, the influence of changes in air pressure with elevation, $T_d$ and $R_0$ on $V_{\text{cmax25}}$, as well as the impact of $\chi$ itself on $V_{\text{cmax25}}$. It also allowed us to separate the effects of $T_g$ and $R_{\text{LAI}}$ on $M_a$ and the effects of LL (indexed by gsl) and moisture (indexed by the ratio of annual actual evapotranspiration to annual potential evapotranspiration) on $M_a$.

Uncertainty of the model predictions

The uncertainty of trait prediction can come from two sources: parameter values and input data. To evaluate the parameter uncertainty, we calculated the uncertainty of each parameter separately and combined them using the standard error propagation formula:

$$u^2(y) = \sum_i \left( \frac{\partial m}{\partial n_i} \right)^2 u^2(n_i),$$

where $u(y)$ is the standard uncertainty of the trait, $\partial m/\partial n_i$ is the sensitivity to variable $n_i$ (obtained by differentiating the individual equations) and $u(n_i)$ is the standard uncertainty of $n_i$. The uncertainty of predicted $M_a$ and $N_{\text{area}}$ values arises from the uncertainties in the coefficients fitted by regression and additional observed $M_a$ and $V_{\text{cmax25}}$ for $N_{\text{area}}$. The uncertainty of $\chi$ and $V_{\text{cmax25}}$ arises from the values of the various ecophysiological quantities in the prediction equations and additional observed $\chi$ for $V_{\text{cmax25}}$, which show some degree of variation among species.

Model evaluation

We evaluated model performance by comparing the observed mean trait value at each site with predictions of each trait, using $r$ and root mean square error (RMSE) between the observed and predicted values across the sites. We compared the $R^2$ explained by the optimality models and statistical models. To test whether the optimality-based models can capture the climate variability, we also fitted multiple linear regressions of the site-mean trait values against the driving climate data which serve as a statistical benchmark. All statistics were performed in R 3.1.3.

Results

Trait variation related to climate

The four climate variables together accounted for 22.2% of the trait variation as shown in the RDA. The first axis explained 16.9% of the variability in the observations. On this axis, variability was negatively related with temperature and positively related with $R_0$ (Figure 2). The second axis reflected gradients in moisture ($\alpha_p$ and vapor pressure deficit). Variability in $\chi$ was shown to be controlled by moisture, although with a small influence from temperature. The $V_{\text{cmax25}}$ varied positively with radiation, and negatively with temperature and moisture, in the opposite direction from $\chi$. Temperature had a small positive influence on $M_a$ but moisture had a negative impact, reflecting the fact that leaves were thicker in hotter and drier environments. The $N_{\text{area}}$ was mainly controlled by radiation and moisture and covaried with $M_a$ and $V_{\text{cmax25}}$.

Observed and predicted trait variation with elevation

All observed traits showed non-linear relationships with elevation (Figure 3). Trait distributions in climate space also showed non-linear relationships. (Figure 4). These non-linear relationships arose because although temperature (as measured by either $T_g$ or $T_d$) decreased monotonically with elevation, the moisture-related variables in the Gongga Mountain region had non-linear relationships with elevation (Figure 1c): the lowest
and uppermost sites had lower mean annual precipitation (MAP) and $\alpha_p$ than the sites at intermediate elevations (see Table S1 available as Supplementary data at Tree Physiology online). The combination of these different trends in individual climate variables led to a complex pattern of trait variability. The $M_a$ and $N_{\text{area}}$ were large under dry conditions and high elevation. The $V_{\text{cmax}25}$ increased along elevation and moisture gradients. The $\chi$ was lower under dry conditions and low elevation. Nevertheless, $M_a$, $V_{\text{cmax}25}$ and $N_{\text{area}}$ tended to increase overall with elevation, while $\chi$ showed an overall decrease with elevation. There was no trend in the CV of any of the traits with elevation (see Figure S3 available as Supplementary data at Tree Physiology online). Within-site CV values were larger than across-site CV values across the sites with $r$ values of 0.68 and 0.67, respectively (see Figure S4 available as Supplementary data at Tree Physiology online). However, predicted $M_a$ using $T_g$, $\chi$ using $T_g$ and predicted $N_{\text{area}}$ using $N_{\text{structure}}$ and $N_{\text{rubisco}}$ were underestimated, and $V_{\text{cmax}25}$ using $T_g$ was overestimated (see Figure S5 available as Supplementary data at Tree Physiology online). Using $T_{\text{dj}}$ instead of $T_g$ improved the predictions of $V_{\text{cmax}25}$ and $\chi$ but degraded the prediction for $M_a$ (Figure 5, see Figure S6 available as Supplementary data at Tree Physiology online). The predicted $\chi$ values using $T_{\text{dj}}$ were better than those using $T_g$, and the best-fit model could predict the values across the sites with $r = 0.71$ and RMSE = 0.06 despite the bias, with median values of $\chi$ underpredicted at most sites (Figure 3). The uncertainties of predicted $V_{\text{cmax}25}$ and $N_{\text{area}}$ were much narrower than the observed ranges. All parameters in the $N_{\text{area}}$ models contributed almost equally to the uncertainty, while the parameter $c$ was the major source of uncertainty for $V_{\text{cmax}25}$. The large uncertainty of $M_a$ and $\chi$ mainly resulted from the intercept and the parameter $\beta$, respectively (see Figure S7 available as Supplementary data at Tree Physiology online).

The $N_{\text{area}}$ was shown to be strongly correlated with both $M_a$ and $V_{\text{cmax}25}$ ($P < 0.001$) (Figure 6, see Figure S8 available as Supplementary data at Tree Physiology online). However, there was a significant effect of including nitrogen fixation (‘N-fixer’) as a factor. At any given $M_a$ or $V_{\text{cmax}25}$, $N_{\text{area}}$ was slightly higher
Predictability of leaf traits with climate and elevation

Figure 4. The distribution of predicted trait values in a climate space defined by elevation and a moisture index ($\alpha_p$) using GAMs. The traits are (a) leaf mass per area ($M_a$), (b) the maximum capacity of carboxylation standardized to 25°C ($V_{\text{cmax25}}$), (c) the ratio of leaf-internal to ambient CO2 partial pressure ($\chi$) and (d) leaf nitrogen content per unit area ($N_{\text{area}}$). Trait values are indicated by the color scale.

Table 2. The comparison between $R^2$ of statistical models (multiple linear regressions of the site-mean trait values against the driving climate data) and optimality models. For $M_a$, $V_{\text{cmax25}}$ and $\chi$, '$T_g$' in parentheses represents the predicted traits using mean temperature during growing season, '$T_dJ$' represents the predicted traits using daytime temperature in July. For $N_{\text{area}}$, '$M_a + V_{\text{cmax25}}$' represents the predicted $N_{\text{area}}$ using observed $M_a$ and $V_{\text{cmax25}}$ in Eq. (19).

| Traits            | Statistical model | Optimality model |
|-------------------|-------------------|------------------|
| $M_a$ ($T_g$)     | 0.55              | 0.33             |
| $V_{\text{cmax25}}$ ($T_dJ$) | 0.45              | 0.60             |
| $\chi$ ($T_dJ$)  | 0.49              | 0.51             |
| $N_{\text{area}}$ ($M_a + V_{\text{cmax25}}$) | 0.65              | 0.84             |

in the nitrogen-fixing species. The prediction of $N_{\text{area}}$ directly from $M_a$ and $V_{\text{cmax25}}$ with our simple method (Eq. 19) was marginally closer to the data than the prediction from $M_a$ and $V_{\text{cmax25}}$ via $N_{\text{structure}}$ and $N_{\text{rubisco}}$ (see Figure S5 available as Supplementary data at Tree Physiology online). The predicted site-mean $N_{\text{area}}$ with our new method but from predicted $M_a$ and $V_{\text{cmax25}}$ was also not significantly different from the observed $N_{\text{area}}$ ($P = 0.08$). These ‘fully predicted’ $N_{\text{area}}$ values were within the range of observations at most sites but were underestimated at low elevation due to the underestimation of predicted $M_a$ (see Figure S5 available as Supplementary data at Tree Physiology online).

**Contribution of climate and elevation to trait variations**

Vapor pressure deficit and temperature were shown to be the most important factors influencing the variation in $\chi$ between sites at different elevations in the Gongga Mountain region, but with opposing effects. Elevation made little contribution to the variation of $\chi$. The $V_{\text{cmax25}}$ was influenced most by temperature and radiation, but elevation also had a small impact on $V_{\text{cmax25}}$. The effects of all the predictors were important for $M_a$ (Figure 7).

**Discussion**

Optimality models have shown skill in predicting the trait variations along the elevation gradient in the Gongga Mountain region, without site- or region-specific calibration of parameters. The $r$ of optimality models was generally higher than statistical models (Table 2). The $r$ of the optimal $M_a$ model was 0.73 when four predictions at low elevation were excluded. This finding suggests that the optimality models considering the underlying mechanisms are better than the statistical models and supports the general validity of these models. The new model for $M_a$—calibrated using an independent set of measurements—correctly predicted patterns in the community-mean $M_a$ of deciduous plants at the Gongga Mountain sites. When the coefficients of $I_{\text{abs}}, f, T_g$ and $\alpha_p$ were calibrated

**Contribution of climate and elevation to trait variations**

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Figure 5. Site-mean values of traits. The traits are (a) leaf mass per area ($M_a$), (b) the maximum capacity of carboxylation standardized to 25 $^\circ$C ($V_{cmax25}$), (c) the ratio of leaf-internal to ambient CO$_2$ partial pressure ($\chi$) and (d) leaf nitrogen content per unit area ($N_{area}$). Observations are site-mean values and predictions are the best versions of different driven data at each site: predicted $M_a$ using mean temperature during the growing season ($T_g$), predicted $V_{cmax25}$ and $\chi$ driven by daily temperature in July ($T_dJ$) and predicted $N_{area}$ using observed $M_a$ and $V_{cmax25}$. The solid line is the 1:1 line.

Figure 6. Partial residual plots showing leaf nitrogen content per unit area ($N_{area}$) as a function of leaf mass per area ($M_a$) and the maximum capacity of carboxylation standardized to 25 $^\circ$C ($V_{cmax25}$) with nitrogen-fixer as an interaction term. (a) The $N_{area}$ as a function of $M_a$ and (b) $N_{area}$ as a function of $V_{cmax25}$. Blue, nitrogen-fixing plants (N-fixer); red, non-nitrogen-fixing plants (non-N-fixer).
Figure 7. The modeled contributions of individual climate variables for each trait at each site. The traits are (a) the ratio of leaf-internal to ambient CO₂ partial pressures (χ), (b) the maximum capacity of carboxylation standardized to 25 °C (V_{max25}) and (c) leaf mass per area (M_a). The gray bars show the changes in predicted trait values compared with the reference level driven by site-mean environment. The green bars show the elevation effect on χ and V_{max25} due to the changes in air pressure. The red bars show the effects of average daytime temperature in July on χ and V_{max25} and the effect of growing season mean temperature on M_a, respectively. The blue bars show the effect of vapor pressure deficit (D_o) on χ and then the effect of χ on V_{max25}. The yellow bars show the effect of growing season mean radiation on V_{max25} and the effect of leaf-area-index-weighted growing season mean radiation on M_a, respectively. The effects of LL (indexed by gsl) and moisture (indexed by the ratio of annual actual evapotranspiration to annual potential evapotranspiration) on M_a are shown in purple and blue.

with the sampled data, the values obtained were 0.99, 0.52, −0.03 and −0.75, which were not significantly different from the values obtained using the China Plant Trait database but were closer to the values for I_{abs} and T_g deduced from the theory. We did not apply the new model to evergreen species because we had no information about their LL. Leaf longevity is strongly related to M_a (Kikuzawa 1991, Reich et al. 1997, Wright and Westoby 2002, Wright et al. 2004). According to the leaf economics spectrum, the LL and M_a of deciduous plants are smaller than those of evergreen plants (Wright et al. 2004). However, LL cannot be reliably estimated in the field without monitoring over a long period (Cornelissen et al. 2003). If such data were available, it would be possible to extend the M_a model to evergreen species.

We have developed a simplified approach to predict N_{area}. This approach produced results close to those obtained using the two-step approach put forward by Dong et al. (2017). The agreement between predictions using the two methods suggests that the hypothesis underpinning both, namely that N_{area} can be predicted as the sum of a photosynthetic component related to V_{max25} and a non-photosynthetic component proportional to M_a (Evans 1989, Lambers and Poorter 1992, Onoda et al. 2004), is reasonable. However, our simpler approach does not require explicit specification of the relative allocation to the metabolic and structural components and, by removing the intermediate steps, reduces the uncertainties and improves the predictions. We have shown that M_a and V_{max25} are predictable from climate and that fully predicted N_{area} values lie within the
range of observations at most sites (see Figure S5 available as Supplementary data at Tree Physiology online). This interpretation differs from some previous studies in which leaf nitrogen availability, implicitly assumed to be related to soil nitrogen availability, is used to predict $V_{cmax25}$ (Luo et al. 2004). There is evidence that soil nutrients, particularly phosphorus rather than nitrogen, can influence $N_{area}$ and $V_{cmax25}$ (He et al. 2014, Gvozdevaite et al. 2018). However, there is growing evidence (i) that LMA exerts a major control on $N_{area}$ (Dong et al. 2017) and (ii) that climate variables are the dominant drivers of $V_{cmax}$. Smith et al. (2019) found that climate variables accounted for about two-thirds of global variation in $V_{cmax}$; soil fertility indices accounted for about one-third. Liang et al. (2020), in a meta-analysis of soil nitrogen enhancement experiments, showed a 2–4 times greater effect on the leaf area and biomass (i.e., whole-plant carbon allocation responses) than on leaf-level $N_{area}$ and $V_{cmax}$. In other words, consistent with optimality theory, plants react to nutrient deficiency more by reducing leaf area, and increasing below-ground carbon investment, than by developing suboptimal leaves. Thus, a key implication of our results is that leaf nitrogen content can be predicted from climate alone. No global analysis of $N_{area}$ is yet available, but the consistency of results for Australia (Dong et al. 2017), Peru (Peng et al. 2020) and this study strongly supports the idea. Moreover, further work should focus on improving $M_a$ prediction since fully predicted $N_{area}$ is underestimated at low elevation due to the underestimation of $M_a$. We have also shown that $R_0$ is positively related to $N_{area}$—consistent with widespread observations that leaf nitrogen is higher at the top of the canopy (Hirose and Werger 1987, Chen et al. 1993) and the optimality hypothesis that nitrogen is unequally allocated within the canopy so as to maximize photosynthesis at each canopy level (Werger and Hirose 1991, Peltoniemi et al. 2012).

Our analyses provide insights into the timescales on which leaf trait acclimation and adaptation operate. Since optimality models implicitly consider acclimation and adaptation in physiological processes, the use of climate inputs at the appropriate timescale—which resulted in better predictions—might provide insight on the corresponding adaptation/acclimation timescale of a trait. We showed that $T_g$ was a better predictor than $T_{dl}$ for $M_a$, suggesting that $M_a$ adapts to the whole growing season environment. The adaptation of $M_a$ to long-term temperature is consistent with the fact that deciduous leaves are built at the beginning of the growing season with one-time carbon investment from the previous year and maximize carbon gain per day, and in turn, net carbon gain during the whole growing season (Kikuzawa 1991). However, although predictions of $V_{cmax25}$ have commonly been made using long-term temperature inputs such as $T_g$ (Wang et al. 2017a, Smith et al. 2019), our results show this can lead to a mis-estimation of $V_{cmax25}$. Using $T_{dl}$ (i.e., daytime during the month the plants were sampled) gives a better prediction, suggesting that $V_{cmax25}$ adapts to environmental conditions during the previous few weeks. Several studies have shown that photosynthetic traits can acclimate quickly to temperature changes (Smith and Dukes 2017, Smith et al. 2017) by regulating intrinsic biochemical characteristics, such as Rubisco content or catalytic turnover rate (Cavanagh and Kubien 2014). Our model data comparison also suggests that $\chi$ acclimates to $T_{dl}$ rather than $T_g$. The least-cost hypothesis underlying the model of $\chi$ considers the total cost of maintaining plant carboxylation and transpiration. Both metabolic processes function mainly in the daytime and can be adjusted rapidly. Therefore, the regulation of $\chi$ is also expected to acclimate to daytime temperature at a weekly to monthly scale, consistent with our finding that $\chi$ is better predicted using $T_{dl}$ than $T_g$. The $\chi$ is highly plastic compared with $M_a$ (Dong et al. 2017), and seasonal variations in $\chi$ for deciduous species have been observed in many studies (Chen and Chen 2007, Ma et al. 2010, McKown et al. 2013); however, the correlation of leaf phenology with seasonal changes in the growth environment of deciduous leaves indicates a need to disentangle their effects in the future. Given that different processes have different timescales for acclimation/adaptation, model inputs should be selected to reflect this.

We have focused on predicting community-mean trait values. Although between-site variation is larger than within-site variation for all traits, nevertheless, there is considerable variability at each site. This variability presumably reflects the within-canopy heterogeneity in bioclimate and in particular in radiation. There are large differences in the photosynthetic traits between sunlit and shaded leaves, and it has also been shown that sunflecks contribute greatly to the photosynthesis of shaded leaves. Our model for $M_a$ is sensitive to radiation inputs. By using $R_{LAI}$ to estimate the average light level absorbed by the leaves within the canopy to drive the $M_a$ model, we were able to obtain relatively good predictions of the community-mean values except at the lowest sites, which may be attributable to disturbance, since many people live at lower elevations in this region. This approach would be insufficient to model within-canopy variability. However, site-based radiation measurements could be used in order to test whether this optimality-based model could predict within-site variation, given appropriate inputs. The within-canopy heterogeneity of other bioclimatic factors may also be important in the choice of appropriate model inputs (Blonder et al. 2018) and for testing the applicability of optimality-based models to explain the within-site variability.

The comparison between the observed and simulated traits allows us to identify mechanisms that are missing from the current optimality framework. For example, our analysis emphasizes the importance of soil moisture constraints. The RDA showed that $V_{cmax25}$ was positively associated with soil moisture, indexed by $\alpha_p$. We found significant relationships between $\alpha_p$ and the residuals of predicted $\chi$ and $V_{cmax25}$. Some hydraulic
traits, including the ratio of leaf-to-sapwood area and specific sapwood hydraulic conductance, also showed significant correlations with photosynthetic traits (see Figure S9 available as Supplementary data at *Tree Physiology* online), suggesting coordination between photosynthesis and water transport. Many studies have shown a strong coordination between hydraulic and photosynthetic traits across species (Brodribb 2009, Scoffoni et al. 2016, Zhu et al. 2018), especially when the hydraulic structure plays a crucial role in limiting the photosynthesis process under water stress (Tyree and Sperry 1989). Lin et al. (2015) analyzed a large global data set and found a positive relationship between wood density and carbon cost per unit water use. We have detected a significant positive effect of wood density on \( V_{\text{cmax}25} \). Further empirical analysis on the coordination between photosynthetic and hydraulic traits over a larger environmental gradient is required. The coordination of photosynthesis and hydraulic traits has already been considered in models to predict stomatal response (Sperry et al. 2017) and vegetation response to drought (Eller et al. 2018), and has been shown to produce improved predictions under water-limited conditions. Our results underline the need to consider aspects of water limitation, in addition to the stomatal response to vapor pressure deficit, in order to predict key plant traits.

Empirical analyses have shown that LL is positively related to potential evapotranspiration and vapor pressure deficit (Wright et al. 2004). In our model, to predict \( M_a \), the effect of \( \alpha_p \) was based on an empirical analysis of an independent global trait data set because there is currently no theory to explain the impact of moisture on optimal LL. Using local data to calibrate the parameters for the theoretical model of \( M_a \) showed that the estimated effect of \( \alpha_p \) is stronger than that indicated by the China Plant Trait database. The RMSE of predictions using the two different sets of calibrated parameters showed larger differences in the lowest values, where the soil moisture constraint is more severe. Given that the effects of other climate variables on \( M_a \) are well captured by the model, it would be worthwhile to try to identify and incorporate the mechanism of moisture impact on optimal LL.

The large functional diversity within sites may result from species attributes, biotic factors or microenvironment (Violle et al. 2014, Pappas et al. 2016). The model uncertainty analysis may provide a new way to estimate the functional diversity. Uncertainty analysis showed that the parameters \( \beta \) and \( c \), representing unit costs for the maintenance of carboxylation, electron transport and transpiration, are the main contributors to uncertainty in \( \chi \) and \( V_{\text{cmax}25} \), respectively (see Figure S7 available as Supplementary data at *Tree Physiology* online). Empirical analysis has shown substantial interspecific variation in \( \beta \), but the current model of \( \chi \) uses a single value of \( \beta \) for all species (Wang et al. 2017b). Using a single value estimated from the published values of photosynthetic capacity (Kattge and Knorr 2007, Wang et al. 2017b) for the parameter \( c \) in the model of \( V_{\text{cmax}25} \), similarly, cannot fully represent its variation among species. Predictions using average values of \( \beta \) and \( c \) estimated from published data could cause mismatches with observed values, such as the predicted \( \chi \) being lower than median observed value at many sites (Figure 3). At the same time, parameter uncertainty due to species variation also represents functional diversity in the community, which could in principle be considered in ecosystem models by specifying a realistic range of values for each parameter. Meanwhile, modeling functional diversity still needs further work both in theory and application.

**Implications for terrestrial ecosystem models**

Optimality theory relies on the concept that natural selection requires plants to acclimate or adapt to prevailing environmental conditions. The development of optimality-based models therefore focuses on identifying the trade-offs between competing requirements. We have shown that optimality-based models for four key traits related to photosynthesis, \( M_a \), \( N_{\text{area}} \), \( V_{\text{cmax}} \) and \( \chi \), predict community-level variability with elevation and climate in the Gongga region, with no need for site- or regional-scale calibration. This finding adds to the growing number of studies showing that patterns of variation in these traits along climate gradients are predictable (Meng et al. 2015, Wang et al. 2017a).

Optimality-based models could be beneficially incorporated into vegetation- or land-surface models since they provide a natural way of accounting for trait variability within PFTs, or across vegetation types, as a function of environmental gradients. The prediction of continuous trait variation with environment would obviate the need to specify parameter values separately for different PFTs (Kucharik et al. 2000, Sitch et al. 2003, Kim et al. 2018) or to account for within-PFT variability probabilistically (see e.g., Kelley et al. 2014). Moving from PFT-based parameters to optimality-based formulations would have the desirable effect of reducing the number of parameters that have to be specified. Moreover, models should improve in realism if the parameter values are allowed to adjust to changing environmental conditions.

However, some issues need to be addressed before implementing optimality-based trait models into vegetation models. First, the timescales of acclimation and adaptation differ between traits. Thus, it is important to ensure that the variability of a given trait is predicted using the appropriate climate information, for example, daytime temperature over a week or month (rather than a climatological growing season average) in the case of \( V_{\text{cmax}25} \). Second, although soil moisture can limit photosynthesis, we lack theoretical understanding of the coordination between plant photosynthesis and hydraulics required to account for this constraint within the current optimality-based modeling framework. Third, the current framework does not account for within-site trait variability and thus does not account...
for functional diversity within communities. Nevertheless, our study suggests a promising way forward to improve both the robustness (with fewer parameters) and realism (considering the acclimation and adaptation of traits) of terrestrial ecosystem models through the prediction of continuous trait variation along environmental gradients.

**Supplementary data**

Supplementary data for this article are available at Tree Physiology Online.

**Data availability**

The trait and climate data are available from Zenodo (Xu et al. 2020). The codes to produce the predicted trait values along with a readme file can be found on GitHub (https://github.com/Huiying-Xu/PTG).

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**Conflict of interest**

None declared.

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**Authors’ contributions**

H.X. carried out the analyses and prepared the manuscript with contributions from all co-authors. H.W. and I.C.P. designed and carried out the fieldwork and contributed to the analyses. H.W. and I.C.P developed the model of $M_\text{s}$. G.W. and X.S. provided climate data for the five sites.

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