Global photosynthetic capacity is optimized to the environment

Abstract
Earth system models (ESMs) use photosynthetic capacity, indexed by the maximum Rubisco carboxylation rate (\(V_{\text{cmax}}\)), to simulate carbon assimilation and typically rely on empirical estimates, including an assumed dependence on leaf nitrogen determined from soil fertility. In contrast, new theory, based on biochemical coordination and co-optimization of carboxylation and water costs for photosynthesis, suggests that optimal \(V_{\text{cmax}}\) can be predicted from climate alone, irrespective of soil fertility. Here, we develop this theory and find it captures 64% of observed variability in a global, field-measured \(V_{\text{cmax}}\) dataset for C\(_3\) plants. Soil fertility indices explained substantially less variation (32%). These results indicate that environmentally regulated biophysical constraints and light availability are the first-order drivers of global photosynthetic capacity. Through acclimation and adaptation, plants efficiently utilize resources at the leaf level, thus maximizing potential resource use for growth and reproduction. Our theory offers a robust strategy for dynamically predicting photosynthetic capacity in ESMs.

Keywords
Carbon cycle, Carboxylation, coordination, ecophysiology, electron transport, \(J_{\text{max}}\), light availability, nitrogen availability, temperature, \(V_{\text{cmax}}\).

Ecosystem and Earth system models are highly sensitive to the representation of photosynthetic processes (Rogers et al. 2017a). In the majority of these models, C\(_3\) photosynthesis is simulated using well-established biochemical theory (Farquhar et al. 1980). The applicability of the theory relies on knowledge of photosynthetic capacity, which varies both among species and over time and space, in response to environmental conditions (Ali et al. 2015; Smith & Dukes 2018).

Photosynthetic capacity is also known to correlate with leaf nitrogen (N) across plant types as a result of the N used to build photosynthetic machinery (Walker et al. 2014). Many global models use these empirical relationships to predict the maximum rate of Rubisco carboxylation (\(V_{\text{cmax}}\); \(\mu\text{mol m}^{-2} \text{s}^{-1}\)), a primary determinant of photosynthetic capacity (Rogers 2014). This approach inherently assumes that variation in \(V_{\text{cmax}}\) is driven by variation in N allocated to leaves, which is itself prescribed or calculated from N availability in soils. This leads to a positive relationship between \(V_{\text{cmax}}\) and...
soil N availability. This approach was shown to perform well in a comparison of several model formulations (Walker et al. 2017). However, there are several important limitations to the N-supply approach for predicting $V_{\text{cmax}}$. First, observed relationships between field-measured $V_{\text{cmax}}$ and leaf N per leaf area ($N_a$) are often only weak (e.g. $r^2 = 0.3$; Niinemets et al. 2009). Second, an increase in $V_{\text{cmax}}$ per leaf $N_a$ at lower soil N availability (Ainsworth & Rogers 2007; Kattge et al. 2009; Maire et al. 2012) suggests that high $V_{\text{cmax}}$ can be achieved under low soil N. Third, the N-supply approach is necessarily empirical, yet it is only with mechanistic models that we stand to reliably predict responses to future, novel conditions.

Photosynthetic coordination theory provides an approach to predict dynamic responses of photosynthetic capacity to environmental constraints. Originally proposed by Von Caemmerer & Farquhar (1981) and further developed by Chen et al. (1993), Maire et al. (2012) and Wang et al. (2017c), it states that photosynthesis tends to be equally limited by electron transport and carboxylation under average environmental conditions. Notably, while this implicitly assumes dynamic nutrient partitioning within leaves, it does not assume any nutrient availability constraint on carboxylation rates, electron transport rates or the partitioning of nitrogen between the two. While this response may be possible under any given amount of N availability, here, we present a ‘strong’ form of the coordination theory, which assumes that plants are able to acquire the N necessary to build leaves that can photosynthesize at the fastest possible rate given light availability and biophysical constraints, for example, through increased belowground allocation (Drake et al. 2011; Terrer et al. 2016). This is quite different, in formulation and consequences, from other interpretations that focus on the partitioning of a fixed amount of N to $V_{\text{cmax}}$ versus $J_{\text{max}}$ (e.g. Ali et al. 2016).

In this study, we tested a theoretical framework for predicting $V_{\text{cmax}}$ from first principles at the global scale. Building on work from Dong et al. (2017), Wang et al. (2017b) and Togashi et al. (2018b), our approach works by combining photosynthetic coordination theory with ‘least-cost’ theory for understanding investments in carboxylation and water transport capacities for photosynthesis (Wright et al. 2003; Prentice et al. 2014). The least-cost hypothesis posits that these investments are co-optimized in relation to environmental properties such that a given photosynthetic rate is achieved at the lowest total cost (i.e. respiration). From this principle, one can predict the optimal CO2 drawdown during photosynthesis (i.e. intercellular to atmospheric CO2 or $C_i;C_a$) as a function of site temperature, vapour pressure deficit and atmospheric pressure (Prentice et al. 2014; Wang et al. 2017c). By drawing together the least-cost and coordination theory, an important step forward is possible: as outlined in the Methods, $V_{\text{cmax}}$ can in theory be predicted as a function of light availability ($J$), temperature ($T$), vapour pressure deficit ($D$) and atmospheric pressure (as indexed by elevation, $z$).

Here, we test this proposition, using a dataset of 3672 values of $V_{\text{cmax}}$ from 201 sites from across the globe. First, we tested our quantitative predictions for individual effects of $J$, $T$, $D$ and $z$ on $V_{\text{cmax}}$ and compared model-predicted $V_{\text{cmax}}$ to observed $V_{\text{cmax}}$ values. Second, we examined the sensitivity of our $V_{\text{cmax}}$ predictions to $I$, $T$, $D$ and $z$ as well as leaf traits not included in the model, namely leaf nitrogen per leaf area ($N_a$) and leaf mass per area ($LMA$). Finally, we used six soil indices to explore the relative influence of soil N and water supply and environmental constraints on $V_{\text{cmax}}$. Using these data, we indirectly tested the proposition that leaf N concentrations more strongly reflect ‘demand’ for N (the need to support a given $V_{\text{cmax}}$ itself optimized to climate) rather than ‘supply’ of N (from the soil).

**MATERIALS AND METHODS**

**Observational $V_{\text{cmax}}$ dataset**

An observational dataset of $V_{\text{cmax}}$ values was built by combining independent data reported to be from top canopy, natural vegetation from Baha et al. (2017), Carswell et al. (2000), De Kauwe et al. (2016), Dominguez et al. (2010, 2015), Ellsworth & Crous (2016), Keenan & Niinemets (2016), Maire et al. (2015), Meir et al. (2002), Niinemets et al. (2015), Rogers et al. (2017b), Serbin et al. (2015), Smith & Dukes (2017a), Tarvainen et al. (2013), Togashi et al. (2018a,b), the TRY plant trait database (Kattge et al. 2011), Wang et al. (2017a) and Wohlfahrt et al. (1999) (Figure S1 and S2). $V_{\text{cmax}}$ values in the dataset were derived from either net photosynthesis ($A_{\text{net}}$) to intercellular CO2 ($C_i$; 56% of the total dataset) curves or from point measurements of $A_{\text{net}}$ and $C_i$ using the one-point method (44%; method presented in De Kauwe et al. (2016); see discussion of the limitations of this method in the Supplementary Information). The dataset includes latitude, longitude and leaf temperature at the time of measurement for each point and, for a subset of the data, leaf nitrogen content per unit leaf area ($N_a$; gN m$^{-2}$; 57% of the dataset) and leaf mass per unit leaf area ($LMA$; g m$^{-2}$; 60% of the dataset). Latitude and longitude were used to extract effective growing season mean temperature ($T_g$; °C), atmospheric vapour pressure deficit ($D_g$; Pa) and incoming photosynthetically active radiation ($I_g$; µmol m$^{-2}$ s$^{-1}$) for each site from monthly, 1901–2015, 0.5° resolution data provided by the Climatic Research Unit (CRU TS3.24.01) (Harris et al. 2014). Growing season was operationally defined as months with mean temperatures greater than 0 °C. The elevation ($z$; m) at each site at 0.5° resolution was obtained from the WFDEI meteorological forcing dataset (Weedon et al. 2014). The ratio of actual evapotranspiration to equilibrium evapotranspiration (Priestley-Taylor coefficient, $\zeta$), which represents the plant-available surface moisture, was calculated at each 0.5° resolution site using the SPLASH model run at a monthly timescale (Davis et al. 2017). Soil cation exchange capacity (CEC; cmolc kg$^{-1}$), soil pH, soil C:N ratio, soil silt content (%) and soil clay content (%) at 0–40 cm depth were extracted from 1 km global data provided by ISRIC SoilGrids database (www.soilgrids.org). These soil data were available for 97% of the total dataset.

**Theoretical model of $V_{\text{cmax}}$**

The theoretical model of $V_{\text{cmax}}$ was developed from the theory presented by Wang et al. (2017c) and Dong et al. (2017) by
combining the coordination theory of photosynthesis (Maire et al. 2012) with the least-cost hypothesis (Wright et al. 2003; Prentice et al. 2014). The combination of the two theories is done by calculating an optimal intercellular CO₂ concentration under average environmental conditions (Cᵢ), which is then used to calculate optimal V'_{cmax} under the same conditions (V'_{cmax}). These calculations were made using light, temperature, vapour pressure deficit, elevation and atmospheric CO₂ as inputs. We first present the formulations for calculating the Cᵢ values used in the optimal V'_{cmax} prediction following Prentice et al. (2014). We then describe how we use coordination theory to predict optimal V'_{cmax} (equation 20 below).

Optimal Cᵢ calculation

The optimal intercellular CO₂ concentration under average environmental conditions (Cᵢ; Pa) was calculated using a theoretical derivation of the optimal ratio (γ) of Cᵢ to atmospheric CO₂ partial pressure (Cᵢ/Pa), based on least-cost theory from Prentice et al. (2014):

$$\gamma = \frac{\Gamma^*}{C_a} + \left(1 - \frac{\Gamma^*}{C_a}\right) \frac{\xi}{\sqrt{D_g}}$$

where

$$\xi = \sqrt{\beta \frac{K + \Gamma^*}{1.6\eta}}$$

(2)

where ξ defines the sensitivity of γ to D_g and is related to the carbon cost of water (Medlyn et al. 2011; Prentice et al. 2014), Γ* (Pa) is the CO₂ compensation point in the absence of mitochondrial respiration, and K (Pa) is as follows:

$$K = K_c \left(1 + \frac{O_i}{K_o}\right)$$

(3)

where K_c (Pa) and K_o (Pa) are Michaelis–Menten coefficients of Rubisco activity for CO₂ and O₂, respectively, and O_i (Pa) is the intercellular O₂ concentration. A consideration of O₂ concentrations is included to account for declines in carboxylation that occur as a result of Rubisco oxygenation. Values of K and Γ* are temperature dependent and were calculated using the equations and parameters of Bernacchi et al. (2001) using T_g. The term β (unitless) in equation 2 is the ratio (b/a) of dimensionless cost factors describing the carbon cost of maintaining photosynthetic proteins to support assimilation at a given rate under normal daytime conditions (b) and the carbon cost of maintaining a transpiration stream to support assimilation at the same rate (a) (Prentice et al. 2014). We used a constant β, estimated as 146, calculated under standard conditions (T_g = 25 °C, D_g = 1 kPa, z = 0) from γ values derived from leaf stable carbon isotope data (Cornwell 2017) and equations 1 and 2, as in Wang et al. (2017c). η* is the viscosity of water relative to its value at 25 °C, calculated using temperature and elevation as in Huber et al. (2009). In cases where C_a was unknown, we used the year of measurement to estimate C_a from global estimates used by the NASA GISS model, which utilizes a combination of measurements and modelling techniques to estimate a global average C_a (https://data.giss.nasa.gov/modelforce/ghg-gases/Fig 1A.ext.txt).

Figure 1 Sensitivity of the theoretical model to environmental drivers. Sensitivity of the theoretical maximum rate of Rubisco carboxylation (V'_{cmax}; black, solid lines) and ratio of intercellular to atmospheric CO₂ concentration (γ; grey dotted lines, panels f, g and h) to the main environmental parameters within the model: growing season mean for irradiances (Iₙ, panels a and e), air temperature (Tₙ, panels b and f) and vapour pressure deficit (Dₙ, panels c and g), as well as elevation (z, panels d and h). In panels a, b, c and d, V'_{cmax} values were mean centred to aid in comparison across environmental parameters. In panels e, f, g and h, values were mean centred and scaled (divided by the standard deviation) to aid comparison of V'_{cmax} and γ sensitivities. Sensitivity analyses were done while keeping all other environmental variables at standard levels: Iₙ = 800 μmol m⁻² s⁻¹, Tₙ = 25 °C, Dₙ = 1 kPa, z = 0 km. Note: γ is insensitive to Iₙ, and as such, no dashed grey line was plotted.

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Optimal $V_{\text{cmax}}$ calculation

We calculated the optimal maximum rate of Rubisco carboxylation under average environmental conditions ($V'_{\text{cmax}}$) by assuming that, optimally, plants will coordinate the allocation of resources to photosynthesis such that under typical environmental conditions:

$$A_c = A_j$$

where $A_c$ (µmol m$^{-2}$ s$^{-1}$) is the photosynthetic rate limited by the maximum rate of Rubisco carboxylation ($V_{\text{cmax}}$; µmol m$^{-2}$ s$^{-1}$):

$$A_c = V_{\text{cmax}} m_c$$

where

$$m_c = \frac{C_j - \Gamma}{C_j + K}$$

where $C_j$(Pa), $\Gamma$ (Pa) and $K$ (Pa) are calculated as in the previous section.

$A_j$ (µmol m$^{-2}$ s$^{-1}$) is the photosynthetic rate limited by the electron transport rate for the regeneration of ribulose-1,5-bisphosphate ($\text{RuBP}$; $J$; µmol m$^{-2}$ s$^{-1}$):

$$A_j = \left(\frac{J}{4}\right)^m$$

where

$$m = \frac{C_j - \Gamma}{C_j + 2\Gamma}$$

$J$ is a saturating function of irradiance, converging on $J_{\text{max}}$ (µmol m$^{-2}$ s$^{-1}$) at high levels:

$$\theta J^2 - (\varphi J + J_{\text{max}}) J + \varphi J J_{\text{max}} = 0$$

where $I$ is the incident photosynthetically active photon flux density (µmol m$^{-2}$ s$^{-1}$), $\theta$ (unitless) is the curvature of the light response curve, and $\varphi$ is the realized quantum yield of photosynthetic electron transport (mol mol$^{-1}$) (Farquhar & Wong 1984). We adopted a value of $\varphi$ of 0.257 mol mol$^{-1}$, which yielded a slope between the measured and predicted $V_{\text{cmax}}$ values near 1. This $\varphi$ value is within the range of values observed by independent, leaf-level studies (0.26 in soya bean (June 2005), 0.23 in soya bean (Harley et al. 1985), 0.28 in Eucalyptus pauciflora (Kirschbaum & Farquhar 1987), and 0.26 in a seven-species analysis (Ehleringer & Björkman 1977)). The curvature term, $\theta$, is related to the distribution of light intensity relative to the distribution of photosynthetic capacity, assumed to be 0.85, consistent with observations (June 2005). Eqn 9 can be substituted into eqn 7 to yield

$$A_j = \left(\frac{m}{4}\right)^{\varphi J + J_{\text{max}}} \pm \sqrt{(\varphi J + J_{\text{max}})^2 - 4\varphi J J_{\text{max}}}}{2\theta}$$

where

$$c = \frac{\partial A_j}{\partial J_{\text{max}}}$$

c is then given by

$$c = \left(\frac{m}{4}\right) \frac{\partial}{\partial J_{\text{max}}} \left(\varphi J + J_{\text{max}} - \sqrt{(\varphi J + J_{\text{max}})^2 - 4\varphi J J_{\text{max}}}}{2\theta}\right)$$

which simplifies to

$$c = \frac{m}{8\theta} \left(1 - \frac{\partial}{\partial J_{\text{max}}} \left(\varphi J + J_{\text{max}} - 2\varphi J \right) \sqrt{(\varphi J + J_{\text{max}})^2 - 4\varphi J J_{\text{max}}} \right)$$

which can be solved as

$$c = \frac{m}{8\theta} \left(1 - \frac{\varphi J + J_{\text{max}} - 2\varphi J}{\sqrt{(\varphi J + J_{\text{max}})^2 - 4\varphi J J_{\text{max}}} \right)$$

Equation 14 can be rearranged to:

$$J_{\text{max}} = \varphi J _{\text{at}}$$

where

$$\sigma = -(1 - 2\theta) + \sqrt{(1 - \theta) \left(\frac{\varphi}{m} \left(1 - \frac{\varphi}{m} \right) - 4\theta\right)}$$

For the calculation of $\sigma$, $c$ was assumed to be non-varying and derived as 0.053 under standard conditions (see Supplementary Information). We then inserted the solution for $J_{\text{max}}$ into eqn 10 and solved for $A_j$:

$$A_j = \frac{\varphi J_{\text{at}} \sigma^*}{\theta}$$

where

$$\sigma^* = 1 + \sigma - \sqrt{(1 + \sigma)^2 - 4\sigma^2}$$

Finally, eqns 5 and 17 were used to replace $A_c$ and $A_j$ in equation 4 and solve for an intermediate rate of $V_{\text{cmax}}$, which we term $V_{\text{cmax}*}$:

$$V_{\text{cmax}*} = \varphi I \left(\frac{m}{m_c}\right) \sigma^*$$

Equation 19 incorporates the temperature response of $m$ and $m_c$. However, $V_{\text{cmax}}$ itself (i.e. the saturation point of the Michaelis–Menten curve) is also sensitive to temperature. As such, we used a formulation from Kattge & Knorr (2007) to incorporate this temperature response, which yielded $V_{\text{cmax}*}$ predicted or predicted $V_{\text{cmax}}$ acclimated to varying environmental conditions:

$$V'_{\text{cmax}*} = \left(V_{\text{cmax}*} e^{\frac{\Delta S}{R T_g}}\right) + \frac{1 + e^{\frac{\Delta S}{R T_g}}}{1 + e^{\frac{\Delta S}{R T_g}}}$$

where $H_d$ is the deactivation energy (200 000 J mol$^{-1}$), $H_a$ is the activation energy (71,513 J mol$^{-1}$), $R$ is the universal gas constant (8,314 J mol$^{-1}$ K$^{-1}$), $\Delta S$ is an entropy term (J mol$^{-1}$ K$^{-1}$), $T_g$ is the growing season temperature in K.

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and \( T_o \) is the optimum temperature in K, assumed to be the temperature at which \( V_c^{\text{max}} \) is operating. \( T_o \) was estimated based on its relationship to growth temperature (Kattge & Knorr 2007):

\[
T_o = 177.884 + 0.44T_g
\]  

(21)

\( \Delta S \) was calculated based on a linear relationship with \( T_g \) from Kattge & Knorr (2007), with a slope of \(-1.07 \text{ J mol}^{-1} \text{ K}^{-1}\) and intercept of \(668.39 \text{ J mol}^{-1} \text{ K}^{-1}\) (Kattge & Knorr 2007).

In addition to \( C' \), the resulting theoretical prediction of optimal \( V_c^{\text{max}} \) (Eq. 20) requires only two free parameters: \( \theta \) (unitless), the curvature of the light response curve, and \( \phi \), the quantum yield of photosynthetic electron transport (mol mol\(^{-1}\)).

### Model-data comparison

To perform the model-data comparison, we standardized each observed \( V_c^{\text{max}} \) value (\( V_{c^{\text{max(\text{mean})}}} \)) to its \( T_g \) (i.e. \( V_{c^{\text{max(\text{obs})}}} \)) using temperature response formulations from Kattge & Knorr (2007):

\[
V_{c^{\text{max(\text{obs})}}} = V_{c^{\text{max(\text{mean})}}}e^{\frac{-h_a(T_a-T_{\text{mean}})}{RT_{\text{mean}}}} - \frac{1 + e^{\frac{r_{\text{max}}(\Delta S-H_a)}{RT_{\text{mean}}}}}{1 + e^{\frac{g(T_g-Y)-H_a}{RT_{\text{mean}}}}}
\]  

(22)

where \( T_{\text{mean}} \) is the leaf temperature at which the measurement was taken (K), \( V_{c^{\text{max(\text{mean})}}} \) is the measured \( V_c^{\text{max}} \) and \( \Delta S \) was calculated as in eqn. 20 from \( T_g \) following Kattge & Knorr (2007). Next, we used the theoretical model described above to predict \( V_c^{\text{max}} \) values at the \( T_g \) for each observation (i.e. \( V_{c^{\text{max(\text{obs})}}} \)). We then aggregated the predicted and observed \( V_c^{\text{max}} \) values by latitude and longitude at a resolution of 0.5 °C to match the climatological data. Finally, we used Model II Reduced Major Axis slope-fitting (R package ‘lmmodel2’ (Legendre 2014)) to compare predicted and observed rates of \( V_c^{\text{max}} \) at each site. To examine the ability of our model to simulate the ratio of \( J_c^{\text{max}} \) to \( V_c^{\text{max}} \) (\( J_c^{\text{max}}/V_c^{\text{max}} \)), we ran a similar comparison of predicted and observed \( J_c^{\text{max}}/V_c^{\text{max}} \) at each of the 90 sites where \( J_c^{\text{max(\text{obs})}}/V_c^{\text{max(\text{obs})}} \) data were available. Note, that due to the similarity between Eqs. 20 and 22 necessarily applied to predicted and observed data for comparison, we explored the potential for a spurious correlation between modelled and observed data due to a common element (Chayes 1971) (Supplementary Information). Additionally, because some \( V_c^{\text{max}} \) values in the observational dataset were derived using the one-point method (method presented in De Kauwe et al. 2016), we ran a similar model-data comparison as above using only data derived using \( A_{\text{net-}C} \) curves (Supplementary Information).

Following direct comparison, we calculated the model bias (\( B \)) in \( V_{c^{\text{max}}} \) predictions at each site as

\[
B = \frac{V_{c^{\text{max(\text{pred})}}}}{V_{c^{\text{max(\text{obs})}}}} - 100
\]  

(23)

We then explored \( B \) as a function of the primary environmental drivers in the model, \( T_g, T_s, D_g \) and \( z \), as well as secondary environmental variables soil cation exchange capacity, soil pH, soil C:N ratio, soil silt content, soil clay content, a soil water content index (\( z \)), leaf mass per area (\( LMA \)) and leaf nitrogen content (\( N_o \)) using multiple linear regression. A single regression model was first fit using the four primary drivers. Following this, a second model was fit that included the four primary drivers and each of the six soil variables, which were available for 193 of 201 sites (97%).

Two additional models were fit that included all primary drivers and one of \( LMA \) or \( N_o \), which were available for 112 (56%) and 98 (49%) of 201 sites, respectively. All analyses were performed in R version 3.5.0.

As a further examination of the influence of soil variables on \( V_{c^{\text{max}}(\text{obs})} \), we fit three separate models using the 193 sites for which soil data were available. The first model, similar to above, only included \( V_{c^{\text{max(\text{pred})}}} \). The second model only included the six soil variables: soil cation exchange capacity, soil pH, soil C:N ratio, soil silt content, soil clay content and \( z \). The third model included both \( V_{c^{\text{max(\text{pred})}}} \) and all six soil variables. The three models were compared using Akaike information criteria (AIC). We also performed a similar comparison using leaf \( N_o \) values for the 98 sites that had \( N_o \) data. For comparisons of models with and without soil variables, each model was fit using only the 193 sites where soil data were available. Similarly, for comparisons of models with and without \( N_o \), each model was fit using only the 98 sites where \( N_o \) data were available. This ensured that model comparisons were done using identical datasets. For all models, we visually examined residual plots following model fitting to ensure that necessary assumptions for model comparisons were met (Zuur et al. 2009). We also calculated the variance inflation factor (VIF) for each model predictor to assess the degree of collinearity. In all cases, VIF values were less than 5 and, in the case of all discussed significant predictors (i.e. \( P < 0.05 \)), values were less than 3, indicating that collinearity did not have a large impact on our interpretations (Zuur et al. 2009).

### Comparison to CANTRIP database

To examine the potential influence of canopy position on our model-data comparison, we examined a subset of the \( V_{c^{\text{max(\text{obs})}}} \) values in the dataset (CANTRIP) (Keenan & Niinemets 2016) that were standardized to top of the canopy light values (\( Q_{\text{int}} = 40 \text{ mol m}^{-2} \text{ d}^{-1} \)). These values were determined using individual canopy scaling relationships, which were applied to 109 individual plant canopies (Niinemets et al. 2015). Separate model-data comparisons, as described above, were performed for the full dataset without the CANTRIP data and with only the CANTRIP data. We used Student’s \( t \)-test to examine whether the difference between modelled and observed data differed between the non-CANTRIP and the CANTRIP data. Both the CANTRIP and non-CANTRIP datasets were normally distributed and had similar standard deviations.

### RESULTS

#### Predicted response of optimal \( V_{c^{\text{max}}} \) to environmental drivers

In response to increased light availability, our model predicted a positive, linear response of optimal \( V_{c^{\text{max}}} \) (i.e. \( V_{c^{\text{max}}} \)). This effect was driven by increases in electron transport under increased light, which led to a necessary increase in \( V_{c^{\text{max}}} \) for carboxylation rate-limited photosynthesis to match electron
transport rate-limited photosynthesis. Similarly, our model predicted a nonlinear increase in $V'_{cmax}$ with temperature (Fig. 1). This was the result of an increase in electron transport with temperature as well as an increased affinity of Rubisco for O$_2$, which also caused an increase in $\chi$. As a consequence, the predicted ratio of $J'_{max}$ to $V'_{cmax}$ decreased with increasing temperatures (Figure S5). The model predicted slight increases in $V'_{cmax}$ with increased vapour pressure deficit and elevation due to reduced stomatal conductance (Fig. 1).

**Model-data comparison**

When compared to the global database, our theoretical model captured 64% of the total variation in $V'_{cmax[obs]}$ values (Fig. 2). After tuning the model to have a slope near 1, the intercept of the relationship between observed and predicted values had a 95% confidence interval (CI) that bracketed 0 (mean = $-2.01$, 95% CI: $-5.49$, 1.12). The model performed similarly well using only data derived from $A_{net}$-C$_i$ curves ($r^2 = 0.68$; Supplementary Information). Our theoretical model was also able to capture 61% of the variation in $J'_{max[obs]}$/$V'_{cmax[obs]}$ at the 90 sites that contained $J'_{max[obs]}$ data (Figure S3). The slope and intercept of the relationship between observed and predicted $J'_{max[obs]}$/$V'_{cmax[obs]}$ values had 95% confidence intervals (CI) that bracketed 0 (slope $= 0.94$, 95% CI: 0.79, 1.12; intercept $= -0.44$, 95% CI: $-0.99$, 0.02). In both cases, there was a slight overprediction of values on average across sites (Fig. 2 and Figure S3).

**Model biases – environmental drivers**

Our theoretical model showed a positive bias with growing season mean irradiance (Fig. 3 and Table S1; $F_{1,196} = 11.54$, $P < 0.01$). This was driven by an overprediction in wet, tropical regions (Fig. 2), potentially due to an overestimation of incoming light in dense tropical forests. To explore whether this was due to an overestimation of light availability, we compared the accuracy of our theory using high-light $V'_{cmax[obs]}$ estimates from the CANTRIP database (Keenan & Niinemets 2016), which are not influenced by canopy shading. The model tended to underpredict the CANTIRIP $V'_{cmax[obs]}$ rates to a greater degree than non-CANTIRIP rates (Figure S4; $t_{76.2} = -2.912$, $P < 0.01$). This result suggests that some data in the observational dataset may have been collected from leaves growing under non-maximum light conditions.

The warmest and driest environments in our dataset ($D_g > 1.5$ kPa) showed the greatest underestimation of $V'_{cmax[obs]}$, leading to a slight negative bias overall (Fig. 3; $F_{1,196} = 7.66$, $P < 0.01$). The model also tended to overpredict $V'_{cmax[obs]}$ at elevations above c. 1500 m (Fig. 3), which led to a significant positive bias in our model with elevation ($F_{1,196} = 11.62$, $P < 0.01$). There was no systematic bias in our model related to $T_g$ (Fig. 3; $F_{1,196} = 2.19$, $P = 0.14$).

**Model biases – leaf traits**

When evaluated across variation in $N_a$ our theory showed a negative bias, indicating an overestimation of $V'_{cmax[obs]}$ among low $N_a$ sites and underestimation at high $N_a$ sites (Fig. 4 and Table S2; $F_{1,92} = 29.67$, $P < 0.01$). To explore the relative impact of $N_a$ versus climate and environmental variables driving the optimality model, we fit three linear regression models predicting $V'_{cmax[obs]}$: one with $V'_{cmax[pred]}$, a second with $N_a$, and a third with $V'_{cmax[pred]}$ and $N_a$ each using the same subset of the dataset where $N_a$ was reported ($n = 98$ sites). The fit of the model that included both $V'_{cmax[pred]}$ and $N_a$ (AIC = 724.5, $r^2 = 0.67$) was slightly better than the model that included just $V'_{cmax[pred]}$ (AIC = 741.7, $r^2 = 0.60$) and substantially better than the model that included $N_a$ (AIC = 828.4, $r^2 = 0.03$), suggesting that, while $N_a$ did add significant predictive value, environmental constraints and light availability (indexed by $V'_{cmax[pred]}$) are the dominant drivers of photosynthetic capacity. Our theory showed no bias in response to LMA (Fig. 4 and Table S3; $F_{1,106} = 0.09$, $P = 0.76$).
Model biases – soil characteristics

For the 193 sites with soil data, we used a linear model to explore the relative influence of soil nutrient and water supply on bias in our theory. Of six indices of soil nutrient and water availability (soil cation exchange capacity (CEC), soil C:N ratio, soil pH, soil silt content, soil clay content and α), only soil pH had a significant influence (Fig. 5 and Table S4; pH: $F_{1,182} = 10.14$, $P < 0.01$; all others: $P > 0.05$). The negative relationship between model bias and pH indicated that our theoretical model tended to overpredict $V'_{\text{cmax,obs}}$ as soil acidity increased. To assess the relative influence of climate and
soil on $V'_{\text{cmax(obs)}}$, we quantified the influence of the soil metrics on model predictive ability by comparing three models for predicting $V'_{\text{cmax(obs)}}$: one based on site climate and elevation (indexed by $V'_{\text{cmax(pred)}}$), a second model with the six metrics of soil nutrient and water availability only, and a third model based on both climate and soils. The fit of the model that included both $V'_{\text{cmax(pred)}}$ and soil variables (AIC = 1529.3; $r^2 = 0.68$) was slightly better than the model that only included $V'_{\text{cmax(pred)}}$ (AIC = 1536.4; $r^2 = 0.64$) and substantially better than the model that only included the soil variables (AIC = 1669.1; $r^2 = 0.32$). These results suggest that soil variables (pH in particular) add statistically significant greater ability to predict $V'_{\text{cmax(obs)}}$ over biophysical constraints and light availability alone, but that the dominant drivers of $V'_{\text{cmax(obs)}}$ are captured by our theory.

**DISCUSSION**

The broad fidelity of our theory to observations suggests that, across large spatial and phylogenetic scales, realized $V'_{\text{cmax}}$ is principally determined by the optimization of photosynthetic processes in response to environmental conditions. Predicted carboxylation capacity is largest in tropical and subtropical regions of the world (Fig. 6), where temperatures and incoming solar radiation are highest. This effect not only follows from the observations presented here (Fig. 2), but also results from temperature (e.g. Smith & Dukes 2017b) and light (e.g. Meir et al. 2007) gradient studies. These results suggest that future, warmer conditions may favour increased photosynthetic potential, although this may be balanced by decreases in $V'_{\text{cmax}}$ as a result of elevated CO₂ (Ainsworth & Rogers 2007).

![Figure 5](image-url)
Nonetheless, there were some significant biases in our model predictions that warrant further discussion. The linear model results indicated a positive bias with light availability, suggesting that the observational data were less sensitive to light availability than predicted by the theory. It is possible that this was driven by individual variation in the realized quantum yield of photosynthetic electron transport (φ), which is the product of the intrinsic quantum efficiency and leaf absorptance of incoming radiation. Previous studies have suggested that intrinsic quantum efficiency and leaf absorptance are not driven by light availability (Evans & Poorter 2001) and, for intrinsic quantum efficiency, that observed variability may be due to measurement technique rather than meaningful biological variation (Skillman 2008). This suggests that the bias in the light response may be due to variability in leaf position and angle, which influence the actual light reaching the leaf surface. Our comparison to the CANTRIP dataset (Keenan & Niinemets 2016) indeed suggests that measured leaves likely were not receiving full sunlight, which would have contributed to the model overestimation that we observed. The combined impact of light availability, leaf position and canopy architecture is a major research need for scaling from leaf to whole-plant responses at large scales.

Unlike with light availability, there was no bias in our model related to temperature, indicating that the temperature response predicted tends to follow similar responses seen in the global dataset. Notably, the response is also similar to that seen in meta-analytical (Kattge & Knorr 2007) and controlled-environment (Scafaro et al. 2017; Smith & Dukes 2017b) studies. Nonetheless, temperature was an important determinant of optimal \( V'_{\text{cmax}} \) rates (Figure 1). Our theory suggests that as temperature increases, higher \( V'_{\text{cmax}} \) is necessary to support increased electron transport up to their optima. This effect is amplified by a greater stimulation of \( K_c \) compared to the CO₂ compensation point, \( I^* \), with temperature (Bernacchi et al. 2001). This phenomenon is also observable as a reduction in the optimal ratio of \( J'_{\text{max}} \) to \( V'_{\text{cmax}} \) at higher temperatures (Figure S5), an effect consistent with previous studies (e.g. Medlyn et al. 2002; Kattge & Knorr 2007; Crous et al. 2013; Smith & Dukes 2017b).

It is worth noting that our theory predicts \( V'_{\text{cmax}} \) rates at the average growing season temperature (i.e. \( V'_{\text{cmax}} \)), rather than at a standardized temperature. Indeed, \( V'_{\text{cmax}} \) at a standardized temperature is likely to be better correlated to \( N_a \) than \( V'_{\text{cmax}} \) is to \( N_a \) because \( V'_{\text{cmax}} \) at a standardized temperature is a proxy for Rubisco content rather than a realized rate. This possibly explains the relatively weaker trend seen here compared to other studies (e.g. Kattge et al. 2009; Walker et al. 2014). Nonetheless, our strategy allows for a prediction of \( V'_{\text{cmax}} \) that is as good or better than a recent approach for estimating \( V'_{\text{cmax}} \) at a standardized temperature from dynamic allocation of leaf N (Ali et al. 2016). Predicting \( V'_{\text{cmax}} \) under typical growth conditions is likely more useful for vegetation modelling because it allows for predictions of \( V'_{\text{cmax}} \) at temperatures near to the temperatures regularly experienced by plants in a given environment, rather than at a common temperature (e.g. 25 °C), which may be atypical for that environment. Thus, \( V'_{\text{cmax}} \) would vary temporally owing to comparatively modest diurnal or day-to-day temperature variation rather than across large temperature gradients, which will minimize potential predictive errors due to the choice of temperature response functions used to scale \( V'_{\text{cmax}} \).

Our approach could be extended to examine the influence of temporal variation in environmental conditions on optimal \( V'_{\text{cmax}} \) predictions. Due to the scale of our analyses and a lack of consistent, high-resolution environmental data, we used monthly mean data (Harris et al. 2014) to create our predictions. While our predictions were able to pick up large spatial trends, the ability of our model to simulate temporal variation is untested here. Better temporal data, coupled with a firmer understanding of the timescale of photosynthetic acclimation, should lead to better temporal predictions.

Our model showed a bias with soil pH, a proxy for soil fertility and leaf \( N_a \). The soil pH effect may be due to the

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**Figure 6** Globally predicted optimal rates of \( V'_{\text{cmax}} \). Global ‘present-day’ optimal rates of maximum Rubisco carboxylation (\( V'_{\text{cmax}} \)) computed using mean growing season irradiance, air temperature, vapour pressure deficit and elevation. Values were calculated at 0.5° resolution using effective growing season mean temperature (\( T_g^* \), °C), atmospheric vapour pressure deficit (\( D_p \), Pa) and incoming photosynthetically active radiation (\( I_g \), \( \mu \text{mol m}^{-2} \text{s}^{-1} \)) for each location from monthly data provided by the Climatic Research Unit (CRU TS3.24.01) (Harris et al. 2014). Growing season was defined as months having temperatures greater than 0 °C. Elevation (\( z \), m) at each location was obtained from the WFDEI meteorological forcing dataset (Weedon et al. 2014). Atmospheric CO₂ was assumed to be 400 \( \mu \text{mol mol}^{-1} \) at \( z = 0 \) m and converted to Pa for each location based on \( z \).
negative effect of soil acidity on nutrient availability, which has been linked to lower rates of photosynthesis (Maire et al. 2015). However, because soil acidity tends to correlate with rainfall (Slessarev et al. 2016), the overprediction may partly be the result of an overestimation of light availability in wet, tropical regions, as mentioned above. The leaf $N_c$ effect indicated that the model underestimated $V'_{c_{\max}}$ in high $N_c$ leaves. This is not surprising, as a substantial amount of leaf $N_c$ is used for Rubisco (Evans 1989). However, neither soil pH nor leaf $N_c$, although significant, provided substantial additional explanatory power over climate. By contrast, a substantial portion of $V'_{c_{\max}}$ is explained by climate alone.

One possible downside to our approach to predicting $V'_{c_{\max}}$ is that our theory, as presented here, does not explicitly include an index of soil moisture and only implements moisture influences through vapour pressure deficit impacts on $C_t$. While it is still uncertain how soil moisture influences $V'_{c_{\max}}$ (Smith et al. 2014), models that include soil water stress impacts on $V_{c_{\max}}$ tend to match observations better than those that do not (Keenan et al. 2010). Nonetheless, our model did not show any bias in relation to an index of soil water availability, $\alpha$. The least-cost theory, as originally presented (Wright et al. 2003), does implicitly assume soil moisture costs to photosynthesis and future work devoted to including these costs explicitly into the quantitative theory could improve model predictions. Optimality based plant hydraulic transport models (e.g. Sperry et al. 2017) could be used for this purpose.

Our findings are consistent with the hypothesis that photosynthetic demand drives leaf nitrogen content, rather than the other way around. This was previously suggested by Evans (1989), after which photosynthetic theory has been used to successfully predict leaf nitrogen concentrations (Dong et al. 2017). However, most current carbon cycle models utilize leaf N content to predict $V_{c_{\max}}$ even those that do not include an interactive N cycle (Smith & Dukes 2013). Our data suggest that leaf N concentration is likely the consequence of demand for $V_{c_{\max}}$. Even so, our theory presents an avenue for reliably predicting $V'_{c_{\max}}$ at global scales without needing to predict $N_c$, which would reduce model uncertainty.

While we found that collinearity of our data likely had no effect on the results presented here (see VIF analysis in Methods), some degree of collinearity in climate and environmental variables is unavoidable when using natural gradient data. A potential next step in testing our theory is to tailor controlled-environment studies to assess the individual response of each operational explanatory power over climate. By contrast, a substantial portion of $V_{c_{\max}}$ is explained by climate alone.

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In conclusion, we have developed and tested a theory for predicting environment-dependent optimal rates of $V'_{c_{\max}}$ against an observational dataset. The agreement between data and theory suggests that plants, through acclimation, adaptation or some combination of the two, are assimilating carbon in an efficient manner by preferentially allocating resources to rate-limiting processes. This allows for greater resources to be used for non-photosynthetic processes, such as growth, storage and reproduction, which are important in competitive environments.

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AUTHOR CONTRIBUTIONS

NGS, TFK, ICP and HW designed the study and developed the theoretical model. IJW and UN provided input during early stages of the study. NGS performed the analyses. NGS, HW, IJW, ÜN, KYC, TFD, RG, FYI, JK, ELK, VM, AR, SPS, LT, HFT, PAT, MW, LKW and SXZ provided data for the analyses. All authors contributed to the writing of the manuscript.

DATA AVAILABILITY

Model code can be found at https://github.com/SmithEco physLab/optimal_vcmax_R (https://doi.org/10.5281/zenodo. 1482044). No new data were used in the analyses. Investigators should refer to the citations provided in the Methods section for data access. Please contact Nick Smith (nick.smith@ttu.edu) with any issues.

REFERENCES


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**SUPPORTING INFORMATION**

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