1 Remote sensing of young leaf photosynthetic capacity in tropical and subtropical

2	evergreen broadleaved forests
3	Xueqin Yang <sup>1, 2, 6,#</sup> , Qingling Sun <sup>2,#</sup> , Liusheng Han <sup>3</sup> , Jie Tian <sup>2</sup> , Wenping Yuan <sup>4</sup> , Liyang Liu <sup>5</sup> , Wei Zheng <sup>2</sup> , Mei
4	Wang <sup>2</sup> , Yunpeng Wang <sup>1,6</sup> , Xiuzhi Chen <sup>2,*</sup>
5	<sup>1</sup> Guangzhou Institute of Geochemistry, Chinese Academy of Sciences, Guangzhou 510640, China
6	<sup>2</sup> Guangdong Province Data Center of Terrestrial and Marine Ecosystems Carbon Cycle, School of Atmospheric
7	Sciences, Sun Yat-sen University, Zhuhai 519082, China;
8	<sup>3</sup> School of Civil Engineering and Geomatics, Shandong University of Technology, Zibo 255000, China;
9	<sup>4</sup> College of Urban and Environmental Sciences, School of Urban Planning and Design, Peking University, Beijing
10	100871, China;
11	<sup>5</sup> Laboratoire des Sciences du Climat et de l'Environnement, IPSL, CEA-CNRS-UVSQ, Université Paris-Saclay,
12	91191 Gif sur Yvette, France;
13	<sup>6</sup> College of Earth and Planetary Sciences, University of Chinese Academy of Sciences, Beijing 101408, China;
14	<sup>#</sup> These authors contributed equally;
15	<sup>*</sup> Correspondance: Xiuzhi Chen ( <u>chenxzh73@mail.sysu.edu.cn</u> )
16	
1/	Abstract
18	Determining the large-scale Rubisco carboxylation maximum rate ( $V_{c,max25}$ ) in relation to leaf age is
19	essential for evaluating the photosynthetic capacity of canopy leaves in global forests. Young leaves ( $\leq 180$
20	days), which exhibit higher $V_{c,max25}$ compared to old leaves (>180 days), are key to controlling the seasonality
21	of leaf photosynthetic capacity in tropical and subtropical evergreen broadleaved forests (TEFs).
22	Nevertheless, quantifying the leaf photosynthetic capacity of different age across TEF's remains challenging,
25	nevel methodology that loverages neighborhood nivels analysis with poplinger least squares entimization to
24	derive the $V_{\rm exc}$ of the young leaves at 0.25° spatial resolution. This approach utilizes satellite based solar
25	induced chlorophyll fluorescence (SIE) products spanning from 2001 to 2018, which were reconstructed
20	using both the TROPOMI (Tronospheric Monitoring Instrument) SIF and MODIS reflectance data (RTSIF)
28	Validations against <i>in situ</i> observations demonstrate that the newly developed $V_{c,max^25}$ products accurately
29	capture the seasonality of voung leaves in South America and subtropical Asia, with correlation coefficients
30	of 0.84, 0.66, and 0.95, respectively. The $V_{c max^{25}}$ of the young leaves simulated from the RTSIF-derived gross
31	primary production (GPP) is effectively correlated (R>0.51) with that dissolved from the global Orbiting
32	Carbon Observatory-2 (OCO-2)-based SIF (GOSIF) GPP. Furthermore, the gridded $V_{c,max25}$ dataset for young
33	leaves successfully detects the green-up regions during the dry seasons in the tropics. Overall, this study
34	presents the first satellite-based $V_{c,max25}$ dataset specifically targeting photosynthetically efficient young
35	leaves, providing valuable insights for modeling large-scale photosynthetic dynamics and carbon cycle in
36	TEFs. Herein, we provide the $V_{c,max25}$ time series derived from RTSIF GPP as the primary dataset,
37	supplemented by GOSIF-derived and FLUXCOM products. These $V_{c,max25}$ products are available at
38	https://doi.org/10.5281/zenodo.14807414 (Yang et al., 2025).
39	

40 Keywords: maximum rate of carboxylation ( $V_{c,max25}$ ), leaf age, photosynthesis, tropical and subtropical forest.

# **1. Introduction**

43 The maximum carboxylation rate  $(V_{c,max})$  is a critical leaf trait that strongly influences the seasonal

- 44 variations in canopy photosynthesis across tropical and subtropical evergreen broadleaved forests (TEFs;
- 45 Chen et al., 2022a; Wu et al., 2018). This relationship stems from the high correlation between the  $V_{c,max}$  and
- 46 nitrogen-related plant functional traits (Lu et al., 2020; Dechant et al., 2020) including leaf nitrogen and
- 47 chlorophyll content (Lu et al., 2020). However, leaf nitrogen content varies substantially at a large scale due
- 48 to the influence of multiple biotic and abiotic factors (Quebbeman and Ramirez, 2016), such as leaf lifespan
- 49 (Onoda et al., 2017), leaf temperature (Verheijen et al., 2013), light intensity (Hikosaka, 2014) and species
- 50 (Evans, 1989). Leaf nitrogen content inversion from remote sensing data at a large scale remains challenging
- 51 (Knyazikhin et al., 2013), hindering the accurate mapping of  $V_{c,max}$  at regional to global scales.
- The  $V_{c,max}$  at 25°C (hereafter denoted as  $V_{c,max25}$ ) serves as a benchmark in most ecosystem models for simulating various  $V_{c,max}$  values at different temperatures. For instance, the Farquhar-von Caemmerer-Berry (FvCB) leaf photosynthetic model, widely adopted for simulating plant photosynthesis across ecosystems (Farquhar et al., 1980; Sun et al., 2015), relies on  $V_{c,max25}$  as a key parameter in determining leaf photosynthetic capacity. However,  $V_{c,max25}$  varies considerably among tree species, with even 2-3-fold differences observed within the same species (Orndahl et al., 2022). Research on this issue remains limited and inconclusive, largely due to the complex interplay of seasonal constraints such as water availability and
- light, which affect leaf flushing and shedding processes across different climatic zones (Brando et al., 2010;
  Yang et al., 2021). Recent advancements have led to the development of two independent satellite remote
- sensing approaches for estimating of  $V_{c,max25}$  at a global scale. The first approach to deriving  $V_{c,max25}$  is via leaf chlorophyll content (LCC) (Luo et al., 2019; Lu et al., 2020), as chlorophyll harvests light and provides
- energy for reactions in the Calvin-Benson-Bassham (CBB) cycle of photosynthesis (Luo et al., 2019).
   Moreover, chlorophyll harvests light energy and powers reactions in the CBB cycle (Luo et al., 2019), V<sub>c.max25</sub>
- exhibits strong coordination with LCC as plants optimize their photosynthetic nitrogen resources (Croft et
- al., 2020; Xu et al., 2022a; Xu et al., 2022b). This LCC-based method enables reliable  $V_{c,max25}$  estimation across various spatiotemporal scales. The second approach estimates  $V_{c,max25}$  using solar-induced chlorophyll
- 68 fluorescence (SIF) (Mohammed et al., 2019), which serves as a robust proxy for global gross primary
- 69 production (GPP) mapping (Mohammed et al., 2019; Frankenberg et al., 2011). Both LCC- and SIF-derived 70  $V_{c,max25}$  products present distinct advantages and limitations. Notably, multispectral satellite data can retrieve
- LCC at significantly higher spatial and temporal resolutions than SIF measurements (Chen et al., 2022a). Nevertheless, LCC retrieval from remote sensing data is susceptible to uncertainty in the vegetation structural parameters employed in the derivation (Luo et al., 2019). Converting LCC to  $V_{c,max^{25}}$  relies on empirical
- parameters employed in the derivation (Luo et al., 2019). Converting LCC to  $V_{c,max25}$  relies on empirical relationships for different plant functional types (PFTs), introducing substantial uncertainties (Chou et al.,
- 75 2020; Croft et al., 2017; Houborg et al., 2013; Houborg et al., 2015). In comparison, while SIF directly 76 correlates with photosynthetic rates, most satellite-based SIF products suffer from relatively coarse spatial 77 and temporal resolutions (Liu et al., 2024; Chen et al., 2022a). A recent study has demonstrated that 78 TROPOMI SIF data, characterized by high spatial and temporal resolution, exhibit a linear relationship with
- 78 TROPOMI SIF data, characterized by high spatial and temporal resolution, exhibit a linear relationship with 79 GPP and contain robust signals for  $V_{c,max25}$  (Chen et al., 2022a). Consequently, TROPOMI SIF has been 80 extensively employed for modeling photosynthesis across various ecosystems (Yang et al., 2023).

TEFs account for 40-50% of the carbon sinks in global forest ecosystems, playing a vital role in the global carbon cycle (Yang et al., 2023; Lu et al., 2021). Despite TEFs maintaining a perennial canopy cover, TEFs exhibit pronounced seasonal variability in photosynthetic activity (Wu et al., 2016). This seasonality is primarily attributed to shifts in canopy leaf age structure (Chen et al., 2021; Chen et al., 2022a), which are predominantly driven by climatic seasonality (Li et al., 2021b; Yang et al., 2021). Recent studies have revealed that young leaves ( $\leq 180$  days) generally exhibit higher  $V_{c,max25}$  than old ones (>180 days), thereby dominating the seasonal dynamics of leaf photosynthetic capacity in TEFs (Locke and Ort, 2014; Wu et al.,

- 88 2016). Consequently, accurately mapping seasonality of  $V_{c,max25}$  seasonality in young leaves is essential for
- 89 modeling tropical and subtropical photosynthesis at continental scales. However, current satellite-based
- 90 approaches face challenges in distinguishing  $V_{c,max25}$  across leaf age cohorts, primarily due to the complex
- 91 interactions between climate drivers and leaf phenology (Jensen et al., 2015; Song et al., 2020). These 92 limitations hinder the seasonal characterization of  $V_{c,max25}$  of young leaves. Additionally, Earth system models
- 92 limitations hinder the seasonal characterization of  $V_{c,max25}$  of young leaves. Additionally, Earth system models 93 (ESMs) often struggle to capture the seasonal variations in  $V_{c,max25}$  across different leaf age categories (Atkir
- 93 (ESMs) often struggle to capture the seasonal variations in  $V_{c,max25}$  across different leaf age categories (Atkin 94 et al., 2014; Ali et al., 2016). A key unresolved challenge remains the insufficient understanding of how
- seasonal changes in water and light availability regulate leaf emergence and shedding patterns.
- 96 To address the aforementioned gaps in mapping  $V_{c,max25}$  of young leaves, we categorized the canopy foliage of TEFs into two distinct leaf age groups: young (≤180 days) and old (>180 days) leaves. We then 97 98 proposed a novel neighbor-based approach to estimate the maximum carboxylation rate ( $V_{c,max25}$ ) for young 99 leaves cohort by assuming a constant for the older cohort (Yang et al., 2023). This assumption is supported by previous research indicating that  $V_{c,max25}$  in old leaves exhibits minimal variation over time (Chen et 100 al.,2019; Albert et al., 2018). This study aims to achieve three key objectives: (1) to develop a global gridded 101 102 dataset capturing seasonal variability of  $V_{c,max25}$  in young leaves across TEFs from 2001 to 2018; (2) to validate the dataset against ground-based measurements and dissolved V<sub>c,max25</sub> data from GOSIF-derived GPP 103 datasets; (3) to analyze the spatiotemporal patterns of  $V_{c,max25}$  in young leaves across TEFs. The resulting 104 105  $V_{c,max25}$  dataset enhances our understanding of tropical and subtropical phenology by quantifying the 106 photosynthetic seasonality of young leaves. Furthermore, it provides valuable insights for refining tropical 107 phenological models within ESMs.
- 108

# 109 2. Materials and methods

#### 110 2.1 Study area

111 The studied TEFs were identified by selecting pixels marked as EBF (Evergreen Broadleaf Forest; Sulla-112 Menashe et al., 2018) on MODIS MCD12C1 land cover maps at 0.05° spatial resolution (see **Fig. 1**). TEFs 113 in South America are the largest tropical rainforests in the world and mainly located at 18°N ~22°S and 114 40~90°W, followed by TEFs in tropical Africa (12°N~12°S, 2.5~37.5°E). TEFs in tropical Asia are mainly 115 located in the Malay Archipelago, Asian Peninsula and northern Australia (30°N~14°S, 85~155°E).

116

# 117 **2.2** Data sources for mapping the $V_{c,max25}$ of young leaves

The continental scale GPP (referred to as RTSIF-derived GPP) at a resolution of 0.125° and spanning 118 from 2001 to 2018 was derived from TROPOMI (Tropospheric Monitoring Instrument) SIF data, according 119 120 to the relationships between the SIF and GPP delineated by Chen et al. (2021), which used a constant value 121 of 15.343 to transform the SIF to the GPP (see Sect. 2.4.1). Monthly meteorological data, including the air temperature (T<sub>mean</sub>) from the ERA5-Land dataset (Zhao et al., 2020), vapor pressure deficit (VPD) from 122 123 ERA5-Land (Yuan et al., 2019), and downward shortwave solar radiation (SW) provided by the Breathing Earth System Simulator (BESS; Ryu et al., 2018), were used to calculate the Michaelis-Menton constant for 124 125 carboxylase ( $K_C$ ), the Michaelis–Menton constant for oxygenase ( $K_0$ ), the CO<sub>2</sub> compensation point ( $\Gamma^*$ ), dark respiration  $(R_d)$ , and thus to calculate the  $A_n$  parameter according to the equations in Table S1 (see the 126 127 Supplement). All datasets were collected and harmonized to a spatial resolution of 0.125°. Further details 128 regarding the satellite and input data are provided in Table 1.

129

# 130 **2.3 Data for validating the** $V_{c,max25}$ **of young leaves**

131 The  $V_{c,max25}$  of the young leaves and canopy-averaged leaves from *in situ* observations were collected to 132 validate the  $V_{c,max25}$  seasonality simulated from RTSIF-derived GPP by the proposed model (**Table S2**). Field

measurements of monthly  $V_{c,max25}$  for young leaves and canopy-averaged leaves were conducted at the 133 Santarem Primary Forest Ecosystem Research Station (BR-Sa1) during August and December 2012 (Albert 134 et al., 2018). Annual V<sub>c,max25</sub> observations for canopy-averaged leaves were acquired over a 12-months period 135 from 2004 to 2016 at the Guyaflux Forest Ecosystem Research Station (GF-Guy) (Wang et al., 2022), from 136 2003 2009 137 to at the Dinghushan Forest Ecosystem Research Station (CN-Din) 138 (https://fluxnet.org/data/fluxnet2015-dataset/), and in November 2012 at the Mbam–Djerem National Park 3 (MDJ-03) (Ferreira Domingues et al., 2015). The V<sub>c,max25</sub> of young leaves and canopy-averaged leaves for 139 the BR-Sa1 site were directly obtained from the literature, whereas for the remaining three sites, only existing 140 141 literature was available, which reported only the  $V_{c,max25}$  of canopy-averaged leaves. To evaluate the simulated  $V_{c,max25}$  of young leaves, the dissolved method (see Sect. 2.5.1) was used to derived the true values of  $V_{c,max25}$ 142 for young leaves, based on a monthly leaf-age-dependent leaf area index (Lad-LAI) product (Yang et al., 143 144 2023). Furthermore, gross primary production retrieved from OCO-2 Solar induced chlorophyll fluorescence (referred to as GOSIF-derived GPP) data at a spatial resolution of 0.05° for the period 2001-2018, and gross 145 primary production from eddy covariance flux tower measurements (referred to as FLUXCOM GPP) data at 146 147 a spatial resolution of  $0.5^{\circ}$  for the period 2001-2013, were used to evaluate the uncertainty of the proposed model in simulating monthly gridded  $V_{c,max25}$  of young leaves (Table 1). 148

149

## 150 **2.4 Methods for simulating the** $V_{c,max25}$ of young leaves

151 Fig. 2 shows the practical procedures applied to produce the seasonal dynamic product of the  $V_{c,max25}$  of young leaves. The 'leaf demographic-identical (LDO)' hypothesis proposes that the leaf cohorts can be 152 153 classified into three categories on the basis of their growth, development and lifespan: young leaf (less than 154 60 days), mature leaf (between 60 days and 180 days), and old leaf (greater than 180 days) (Wu et al., 2017b). To ensure comparability between the observations and simulations and simplify the calculations, we 155 categorized the leaf area index (LAI) and the corresponding net CO<sub>2</sub> assimilation rate  $(A_n)$  into two groups 156 based on leaf age: those with a leaf age greater than 180 days were considered 'old', and those with a leaf 157 age less than 180 days were considered 'young' (Chen et al., 2020). Since the total GPP of the leaf cohort 158 159 remained constant and the leaf cohorts were composed of leaves of different ages, we calculated the total 160 GPP as a sum of the GPP of each leaf age cohort. The total GPP was simulated using the FvCB photochemical model by combining the LAI groups (young leaf  $LAI_Y$  vs. old leaf  $LAI_O$ ; Equation 1) and the corresponding 161 162 net assimilation rates of CO<sub>2</sub> (young  $A_{n, sat Y}$  vs. old leaf  $A_{n, sat O}$ ; Equation 1) (Farquhar et al., 1980).

163  $GPP_{total} =$ 

 $GPP_{total} = LAI_Y \times A_{n,sat_Y} + LAI_O \times A_{n,sat_O}$ (1)

where  $LAI_Y$  represents the LAI of young leaves ( $\leq 180$  days) and LAI<sub>O</sub> represents the LAI of old leaves ( $\geq 180$  days).  $A_{n, sat\_Y}$  and  $A_{n, sat\_O}$  represent the net CO<sub>2</sub> assimilation rates of young and old leaves, respectively. The sum of  $LAI_Y$  and  $LAI_O$  was set as the total canopy LAI. GPP<sub>total</sub> refers to the total gross primary production of the canopy.

The gridded GPP data over the whole TEFs were derived from SIF (denoted as RTSIF-derived GPP) 168 using a linear SIF-GPP regression model (see Sect. 2.4.1), which was established based on in situ GPP from 169 76 eddy covariance (EC) sites (Chen et al., 2022b). The majority of the TEFs retain leaves year-round, and 170 171 their total LAI shows marginally small spatial and seasonal changes (Wu et al., 2016; Fig. S1). Therefore, 172 previous modeling studies have assumed a constant value for the total LAI in TEFs (Cramer et al., 2001; Arora and Boer, 2005; De Weirdt et al., 2012). Based on this, we collected observed seasonal LAI dynamics 173 174 in TEFs from previously published literature, which showed a constant value of LAI at around 6.0 (Fig. S1; 175 Table S3). Consequently, we streamlined the data to assume that the seasonal LAI was broadly equivalent to 6.0 in TEFs. This assumption was also found to be reasonable in the region of the TEFs by Yang et al. (2023). 176 The  $V_{c,max25}$  values for old leaves were set to 20  $\mu$ mol m<sup>-2</sup> s<sup>-1</sup> according to previous ground-based 177

- observations (Chen et al., 2020; Zhou et al., 2015) in our method. The  $A_{n, sat_O}$  can be calculated according to the FvCB biochemical model (Farquhar et al., 1980; Bernacchi et al., 2003; see Sect. 2.4.2).  $A_{n, sat_Y}$  can be expressed as the function of  $V_{c,max25}$  for young leaves (see Sect. 2.4.2). Consequently, only  $LAI_Y$  and  $V_{c,max25}$ of young leaves remains as the final parameters to be solved in Equation 1.
- 182 The complexity of model is evident due to the two parameters that needed to be solved. To overcome 183 the challenge of the calculation, we assumed that the four adjacent pixel points had homogeneous plant functional types (PFTs) and had consistent leaf age cohorts. The LAI and  $V_{c,max25}$  of young leaves were 184 estimated using nonlinear least squares and constraints on the basis of the GPP values with the four 185 186 neighboring pixels according to Equation 1. The optimal  $V_{c,max25}$  was determined by minimizing the residual 187 while satisfying the positivity constraint (i.e.,  $V_{c,max25}>0$ ). The input gridded dataset consisted of the GPP obtained from the RTSIF and climatic data such as Tmean, VPD and SW. The spatial resolution of these data 188 was homogeneously resampled to 0.125°, resulting in a spatial resolution of 0.25° for the map of the output 189  $V_{c,max25}$  of young leaves. To further validate the robustness and reliability of our neighborhood pixel method, 190 191 we conducted sensitivity analysis by systematically varying the number of neighborhood pixels, ultimately 192 generating the  $V_{c,max25}$  product with 0.5° spatial resolution. In the optimization process, an mean  $V_{c,max25}$  value 193 was determined by assuming that the leaf cohort was completely young. A reasonable adjustment for the  $V_{c,max25}$  of young leaf value was then determined based on previously published literature (Chen et al., 2021; 194 195 Yang et al., 2023) and the initial value. Importantly, the difference between the finely optimized  $V_{c,max25}$  of 196 young leaves and the initial value could often be significant and outside the margin of error. Therefore, an 197 appropriate adjustment for the  $V_{c,max25}$  of young leaves value needs to be carefully determined (He et al., 2019). All analyses were performed using MATLAB (R2 version). 198
- 199

#### 200 2.4.1 Calculating the GPP (RTSIF-derived GPP) from TROPOMI SIF

201 SIF is a widely used proxy for canopy photosynthesis (Yang et al., 2015; Dechant et al., 2020). Here, 202 we used a long-term reconstructed TROPOMI SIF dataset (RTSIF; Chen et al., 2022b) to estimate GPP seasonality. Previous analyses showed that RTSIF was strongly linearly correlated to eddy covariance (EC) 203 GPP and used 15.343 as a transformation coefficient to convert RTSIF to GPP (Chen et al., 2022b). We 204 collected seasonal GPP data observed at four EC sites from the FLUXNET2015 tier 1 dataset (Table S4; 205 Pastorello et al., 2020) and validated the Chen et al. (2022) simple SIF-GPP relationship (Fig. S2 in the 206 Supplement). Results confirmed the robustness of the Chen et al. (2022b) simple SIF-GPP relationship for 207 208 estimating the GPP seasonality in TEFs (R>0.49). Despite the potential overestimation (Fig. S2a) or 209 underestimation (Fig. S2d) of the magnitudes, the RTSIF-derived GPP mostly captured the seasonality of the 210 EC GPP at all four sites ( $d_{phase} < 0.29$ ).

211

218

# 212 **2.4.2** Calculating the net CO<sub>2</sub> assimilation rate

The net CO<sub>2</sub> assimilation rate is a significant parameter characterizing the photosynthetic rate. According to Farquhar's (1980) biochemical model (FvCB), the net CO<sub>2</sub> assimilation rate ( $A_n$ ) depends on the most limiting conditions for photosynthesis (RuBisCO saturation  $A_c$ , RuBP saturation  $A_j$ , or TPU saturation  $A_p$ ) and the intensity of dark respiration (R<sub>d</sub>, Bernacchi et al., 2013). The net CO<sub>2</sub> assimilation rate (either  $A_{n, sat_v}$  or  $A_{n, sat_v}$ ) can be expressed by the following equation:

#### 219 (1) Calculation of $A_c$

$$A_n = \min(A_c, A_j, A_p) - R_d \tag{2}$$

220 When the CO<sub>2</sub> pressure is low ( $C_i < 300 \ \mu mol \ mol^{-1}$ ), the net photosynthesis rate is mainly constrained 221 by the activity and quantity of the carboxylase RuBisCO. The Rubisco-limited photosynthetic rate  $A_c$  can be 222 calculated using the following equation under a limited carboxylation rate:

223 
$$A_c = V_{cmax} \times \frac{C_i - \Gamma^*}{C_i + K_c \times (1 + \frac{O}{K_c})}$$
(3)

where  $\Gamma^*$  represents the CO<sub>2</sub> compensation point and  $C_i$  is the intercellular CO<sub>2</sub> pressure.  $K_c$ ,  $K_0$ , O, and  $\Gamma^*$  are estimated based on the leaf temperature using **Equation 4** to calculate their values at the given temperature, which is used to convert from their values at 25°.

$$P = P_{25} \times e^{\frac{(T_k - 298.15) \times \Delta H_p}{r \times T_k \times 298.15}}$$
(4)

where *P* is the parameter at each temperature that varies with temperature, including the Michaelis constant for O<sub>2</sub> ( $K_0$ ), the Michaelis constant for CO<sub>2</sub> ( $K_0$ ), the intercellular concentration (*O*) and the CO<sub>2</sub> compensation point ( $\Gamma^*$ ).  $P_{25}$  denotes the constant temperature dependence parameter at 25°C (Bernacchi et al., 2001); specifically,  $K_c$ ,  $K_0$ ,  $\Gamma^*$  and *O* at 25°C are equal to 404.9 µmol mol<sup>-1</sup>, 278.4 mmol mol<sup>-1</sup>, 42.75 µmol mol<sup>-1</sup>, and 210 mmol mol<sup>-1</sup>, respectively.  $\Delta H_p$  is the activation energy and varies with the temperature and parameters. *r* is the standard gas constant (8.314 J mol<sup>-1</sup> K<sup>-1</sup>).  $T_k$  is the leaf temperature (unit: K).

Using the stomatal conductance model, the internal  $CO_2$  concentration ( $C_i$ , Equation 5) was estimated to depend on the atmospheric  $CO_2$  concentration instead of the ambient relative moisture (Xu et al. 2017; Lin et al., 2015; Medlyn et al., 2011).

237 
$$C_i = 380 \times (1 - \frac{1}{1.6 \times (1 + \frac{3.77}{\sqrt{VPD}})})$$

238 where  $C_i$  represents the internal CO<sub>2</sub> concentration.

239 (2) Calculation of  $A_j$ 

240 When the concentration of  $CO_2$  is high, leaf photosynthesis is constrained by RuBP regeneration. The 241 photosynthetic rate ( $A_j$ ) is then limited by electron transport and calculated using the following equation:

242

227

$$A_j = J \times \frac{C_i \cdot \Gamma^*}{4 \times (C_i + 2 \times \Gamma^*)}$$
(6)

(5)

where *J* is the electron transport rate for leaf photosynthesis. It is a quadratic function of the full electron transfer rate ( $J_e$ ) and maximum electron transfer rate ( $J_{max}$ ) (Luo et al., 2001; Bernacchi et al., 2013). The maximum electron transport rate ( $J_{max}$ ), the maximum carboxylation rate ( $V_{c,max25}$ ), and the CO<sub>2</sub> compensation point in the absence of mitochondrial respiration (r\*) were used to determine the Michaels– Menten constants for oxygenation and carboxylase. For the detailed calculation process, refer to **Equations 7-9**.

249 
$$J = \frac{J_e + J_{\max} - \sqrt{(J_e + J_{\max})^2 - 4 \times J_e \times J_{\max} \times \theta}}{2 \times \theta}$$
(7)

263

$$J_e = PAR_{total} \times \partial \times \varphi \times \kappa$$

$$J_{max} = J_{max,25} \times e^{\left(\left(\frac{25-T_{opt}}{\Phi}\right)^2 - \left(\frac{T_k - 273.15 - T_{opt}}{\Phi}\right)^2\right)}$$
(8)
(9)

where  $J_{\text{max}}$  denotes the maximum electron transfer rate at a given temperature and varies with temperature.  $J_{\text{max},25}$  is the maximum electron transfer rate at 25°C, is usually assigned  $1.67 \times V_{c,max25}$  in TEFs.  $T_{opt}$  is the optimum temperature for electron transfer.  $J_e$  is a function of canopy photosynthetically active radiation (*PAR<sub>total</sub>*) and can be calculated by inputting SW and LAI; for details, refer to Weiss and Norman (1985) and Ryu et al. (2018).  $\theta$ ,  $\partial$ ,  $\varphi$ , and  $\Phi$  are constants and equal to 0.7, 0.85, 0.5, and 0.85, respectively (Xu et al., 2017; Yang et al., 2023).  $\kappa$  is a function of the optimal temperature, which represents the maximum quantum efficiency of PSII photochemistry.

#### $259 \qquad (3) Calculation of A_p$

The rate of photosynthesis is limited by the export of triose phosphate.  $A_p$  represents the photosynthetic capacity to export or utilize the photosynthetic products for the different LAI cohorts, as determined by multiple field observations.

- $A_p = c \times V_{c,max25} \tag{10}$
- 264 The ratio of the interior foliar CO<sub>2</sub> concentration to the environmental CO<sub>2</sub> concentration was fixed at

265 0.5 for C<sub>3</sub> species and 0.7 for C<sub>4</sub> species based on previous investigations (Fabre et al., 2019; Mcclain and
266 Sharkey, 2019; Yang et al., 2016).

- 267
- 268

#### 2.5 Methods for evaluating the simulated *V<sub>c,max25</sub>* of young leaves

This study assessed the proposed algorithms in three ways: (1) monthly in situ  $V_{c,max25}$  observations 269 obtained from the literature; (2) annually dissolved  $V_{c,max25}$  from the GOSIF-derived GPP; and (3) a monthly 270 Lad-LAI product covering the entire TEF region, derived from the RTSIF product by Yang et al. (2023). 271 However, in situ  $V_{c,max25}$  observations of young leaves remain scarce, with only one site (BR-Sa1; see Sect. 272 273 **2.3**) providing monthly  $V_{c,max25}$  data. To compensate for the lack of ground-based validation, seasonal  $V_{c,max25}$ of canopy-averaged leaves was collected from three additional sites (GF-Guy, MDJ-03 and CN-Din), and the 274 275  $V_{c,max25}$  of young leaves at these sites was estimated using the dissolved method (see Sect. 2.5.1) based on 276 the Lad-LAI product (Yang et al., 2023). To evaluate the efficiency and reliability of the newly proposed methodology, we compared the gridded  $V_{c,max25}$  of young leaves simulated from RTSIF-derived GPP using 277 the proposed method with that estimated from GOSIF-derived GPP and the Lad-LAI product using dissolved 278 279 method. To investigate the reliability of the neighborhood-based subdivision technique, we conducted a comparative analysis for the  $V_{c.max25}$  of young leaves derived from RTSIF-derived GPP using 2×2 (0.25° 280 resolution) and  $4\times 4$  (0.5° resolution) neighboring pixels. To assess the uncertainties stemming from the 281 282 estimation of gross primary production, we incorporated two additional GPP products, GOSIF-derived and 283 FLUXCOM GPP (Jung et al., 2019; Yang et al., 2023), along with the original RTSIF-derived GPP, resulting in three distinct versions of the  $V_{c,max25}$  of young leaves products. 284

285

#### 286 **2.5.1** Dissolved method for evaluating the $V_{c,max25}$ of young leaves

The total GPP can be expressed as the sum of the GPP of the young and old cohorts. The GPP of each 287 288 leaf age cohort is a function of the corresponding LAI cohort and net CO<sub>2</sub> assimilation rate. In accordance 289 with related studies, the  $V_{c,max25}$  of old leaves is presumed to be a constant value (Chen et al., 2020). When 290 the LAI of different leaf ages is known, only the  $V_{c,max25}$  of the young leaves remains unknown in Equation 291 1. The value of the  $V_{c,max25}$  of the young leaves can be determined by solving the aforementioned Equation 292 1. This method involves dividing GPP into young and old cohort according to leaf age, with the  $V_{c,max25}$  of young leaves being directly solved by using the Lad-LAI product, hence the term 'dissolved method'. At 293 present, there is a lack of available data regarding the ground  $V_{c,max25}$  of different leaf ages. The dissolved 294 295 method is employed to validate the reasonableness of the proposed algorithm.

296

#### 297 2.5.2 K-means method for classification

We analyzed the spatial patterns of  $V_{c,max25}$  across TEFs using the K-means clustering analysis. K-means 298 algorithm is an iterative algorithm that tries to partition the dataset into K predefined distinct non-overlapping 299 300 subgroups (clusters) where each data point belongs to only one group (Ikotun et al., 2023). It tries to make 301 the intra-cluster data points as similar as possible while also keeping the clusters as different as possible. It assigns data points to a cluster such that the sum of the squared distance between the data points and the 302 cluster's centroid (arithmetic mean of all the data points that belong to that cluster) is at the minimum. Intra-303 304 cluster homogeneity increases as variation decreases, indicating greater similarity among constituent data points. The way k-means algorithm works is as follows: 305

306 (1) Specify number of clusters K.

307 (2) Initialize centroids by first shuffling the dataset and then randomly selecting K data points for the 308 centroids without replacement.

- 309 (3) Iterate until convergence (i.e., cluster assignments remain unchanged between iterations).
- 310 (4) Compute the sum of the squared distance between data points and all centroids.
- 311 (5) Assign each data point to the closest cluster (centroid).
- 312 (6) Compute the centroids for the clusters by taking the average of all points that belong to each cluster.
- 313

#### 314 2.5.3 Random forests regression

315 Random Forests (RF) is a widely used ensemble learning method that constructs multiple decision trees 316 through bootstrapped sampling of the training data and aggregates their predictions to enhance model robustness in regression tasks (Yang et al., 2022). This method is particularly effective in capturing non-317 linear relationships and interactions among predictor variables, making it well-suited for complex ecological 318 319 datasets. In this study, we employed RF regression to identify the dominant climatic drivers of tropical forest 320 dynamics across the entire tropical region as well as within three major tropical forest regions. The model was trained using climate variables as predictors and  $V_{c,max25}$  of young leaves as dependent variables. We 321 322 utilized the feature importance scores derived from RF to rank the influence of three climatic variables on 323 forest dynamics across different regions, providing insights into the spatial heterogeneity of climate-forest 324 interactions.

325

329

#### 326 2.5.4 Precision evaluation index

Both the root mean square error (RMSE, Equation 11) and Pearson's correlation coefficient (R,
 Equation 12) were employed to evaluate the model capabilities.

$$RMSE = \sqrt{\frac{\sum_{i=1}^{N} (V_i - U_i)^2}{N}}$$
(11)

330 
$$R = \frac{\sum_{i=1}^{N} (V_i - \bar{V}) (U_i - \bar{U})}{\sqrt{\sum_{i=1}^{N} (V_i - \bar{V})^2} \sqrt{\sum_{i=1}^{N} (U_i - \bar{U})^2}}$$
(12)

331 
$$SD_{S} = \sqrt{\frac{1}{N} \sum_{i=1}^{n} (V_{i} - \bar{V})^{2}}$$
(13)

332 
$$SD_m = \sqrt{\frac{1}{N} \sum_{i=1}^n (U_i - \overline{U})^2}$$
 (14)

 $LCS = 2SD_SSD_m(1-R)$ (15)

where *N* is the total point extracted from the  $V_{c,max25}$  products simulated from RTSIF-derived GPP;  $V_i$  and *U<sub>i</sub>* represent the monthly simulated and observed  $V_{c,max25}$  in situ measurements, respectively; and  $\overline{V}$  and *U<sub>i</sub>* are the mean values of the simulated and observed  $V_{c,max25}$  in situ measurements. Moreover, the continental *V<sub>c,max25</sub>* simulated from the proposed model was compared against that the dissolved from GOSIF-derived GPP and Lad-LAI in TEFs. *SD<sub>S</sub>*, *SD<sub>m</sub>*, and LCS represent the standard deviation of the simulation, standard deviation of the measurement, and the lack of correlation weighted by the standard deviations (phase-related difference; d<sub>phase</sub>), respectively.

341

#### 342 **2.6** Quality control (QC) for young leaves *V<sub>c,max25</sub>* product

We provided information on data quality control (QC) along with the  $V_{c,max25}$  of young leaves product (Fig. S3). In the QC system (Table S5), data quality was divided into four levels: Level 1 represents the highest quality, Level 2 and Level 3 represent good and acceptable quality, respectively, and Level 4 should be used with caution. This QC product was generated based on Pearson's correlation coefficients (R) and the root mean square error (RMSE), which were obtained by comparing the seasonal  $V_{c,max25}$  estimated from RTSIF- and GOSIF-derived GPP.

#### 350 **3. Results**

351 **3.1** Validation of the gridded  $V_{c,max25}$  seasonality of young leaves using *in situ* observations

The seasonality of simulated mean  $V_{c,max25}$  for both all canopy leaves and young leaves was evaluated 352 with in situ measurements at 4 sites: CN-Din site in southern China (23.17°N, 112.54°E), MDJ-03 site in 353 Congo (5.98°S, 12.87°E), BR-Sa1 (2.86°S, 54.96°W) and GF-Guy (5.28°N, 52.93°W) sites in southern 354 355 America. Overall, the estimated mean V<sub>c,max25</sub> of canopy-averaged leaves (black line) ranged from 20 µmol  $m^{-2} s^{-1}$  to 60 µmol  $m^{-2} s^{-1}$ , and their seasonal fluctuations agreed well with the *in situ* mean  $V_{c,max25}$  (red dots) 356 (Fig. 3). In contrast,  $V_{c,max25}$  of the young leaves (green line) exhibited higher values compared with those of 357 canopy-averaged leaves, ranging from between 40 and 80 µmol m<sup>-2</sup> s<sup>-1</sup>. This finding is consistent with 358 previous studies that young leaves were more photosynthetically effective than old leaves (Urban et al., 2008; 359 360 Albert et al., 2018; Menezes et al., 2022). Specifically, our simulations can capture well the seasonal patterns of  $V_{c,max25}$  across different sites. At the BR-Sa1 site, the estimates were correlated well with the observed 361 mean V<sub>c,max25</sub> for all (R=0.85) and young leaves (R=0.84), which both increased during the dry season 362 (approximately between June and December) (Fig. 3a, 3b). At the GF-Guy site, in situ mean V<sub>c.max25</sub> of all 363 364 canopy leaves showed considerable seasonality, while the  $V_{c,max25}$  of young leaves remain more stable (Fig. **3c**). Our estimations also performed well in simulating the  $V_{c,max25}$  of all canopy leaves (R=0.95) and that of 365 366 young leaves (R=0.66) (Fig. 3d). In contrast, at the Din site in subtropical Asia, both  $V_{c,max25}$  for canopy-367 averaged leaves and young leaves increased during the wet-season period, with the highest precipitation occurring in June or July (Fig. 3e). This is similar in the MDJ-03 site, where both  $V_{c,max25}$  for all canopy 368 369 leaves and young leaves also increased during the wet-season period but with larger seasonal variations. Our model showed the best simulations of  $V_{c,max25}$  of young leaves at Din site (canopy-averaged leaves: R=0.84; 370 young leaves: R=0.95). Nevertheless, many more long-term in situ measurements are needed to determine 371 372 the reliability of these time series fluctuations.

Then, we analyzed the spatial patterns of  $V_{c,max25}$  across TEFs using the K-means clustering analysis. Results showed that  $V_{c,max25}$  for young leaves cohorts had evident seasonal dynamics, bringing influences on canopy photosynthesis. **Fig. S4** shows the timeseries fluctuations in  $V_{c,max25}$  for the young leaves in ten individual regions, as clustered using K-means analysis. Results show the amplitude of  $V_{c,max25}$  for young leaves is smaller in regions closer to the equator and larger in regions farther away from the equator.

378

# 379 **3.2** Validation of the $V_{c,max25}$ of young leaves simulated from RTSIF-derived GPP against 380 that dissolved from GOSIF-derived GPP

The  $V_{c,max25}$  of young leaves simulated from RTSIF-derived GPP demonstrated significant correlations 381 (R ranges from 0.51 to 0.87) with those dissolved from GOSIF-derived GPP (Fig. 4). Across the Amazon, 382 more than 69.78% of pixels have a high EBF fraction (>90%). The spatial clustering pattern aligns with the 383 onset of the dry season (cf. Tang and Dubayah, 2017), suggesting that the clustering analysis effectively 384 differentiates climate regions within the Amazon. The relatively homogeneous environmental conditions 385 386 across these sub-regions create similar plant growth environments, leading to a more constrained range of  $V_{c,max25}$  values and pronounced clustering effects in sub-regions A1–A5. Notably, sub-region A3, located in 387 the northwestern Amazon near coastal and mountainous areas, forms two distinct clustering zones. Statistical 388 analysis revealed strong seasonal correlations between the  $V_{c,max25}$  of young leaf simulated from RTSIF-389 derived and GOSIF-derived GPP, with R>0.80 occupy 91.68% (Fig. 5a-c) and RMSE<11.59 occupy 91.68% 390 391 (Fig. 5d-f) of the TEFs. The K-means spatial clustering analysis revealed strong agreement between the 392  $V_{c,max25}$  of young leaves simulated by RTSIF-derived and GOSIF-derived GPP in the low-latitudes (Amazon R1: R=0.90; Amazon R2: R=0.94; Amazon R4: R=0.87; Amazon R5: R=0.77; Congo R6: R=0.91; Congo 393

R7: R=0.97; Asia R8: R=0.86; Asia R9: R=0.84; Fig. S5) compared to higher-latitude areas (Amazon R3:
R=0.60; Amazon R10: R=0.50; Fig. S5). This latitudinal gradient was similarly reflected in RMSE values,
with lower errors in equatorial regions (Amazon R1: RMSE=1.78; Amazon R2: RMSE=2.17; Amazon R4:
RMSE=4.67; Congo R6: RMSE=3. 26; Congo R7: RMSE=4.73; Asia R8: RMSE=3.38; Asia R9:
RMSE=5.86; Fig. S6) versus higher-latitude zones (Amazon R5: RMSE=14.85; Amazon R10: RMSE=6.92;
Fig. S6).

400

#### 401 **3.3** Comparison of the seasonal $V_{c,max25}$ of young leaves with the leaf age product

402 While field measurements have identified distinct seasonal patterns in the  $V_{c,max25}$  of young leaves across 403 TEFs, the sparse distribution of observation sites hinders comprehensive assessment of these variations. To 404 address this issue, we conducted K-means clustering analysis on simulated  $V_{c,max25}$  maps to evaluate their 405 spatial coherence relative to the leaf age product developed by Yang et al. (2023).

406 The spatial distribution of clustered  $V_{c,max25}$  of young leaves, derived from satellite vegetation signals 407 (Fig. 6a-c), closely aligned with climate-based classifications from Chen et al. (2021) (Fig. 6d-f). These 408 patterns showed strong correspondence with the Lad-LAI clusters based on endogenous climate variables 409 reported by Yang et al. (2023) (Fig. 6g-i). Collectively, these results demonstrate similar spatial clustering patterns. In the middle (R2) and northern (R3) Amazon (Fig. 7a), the seasonal variation in the  $V_{c,max25}$  of 410 young leaves (Fig. 8b, c) was consistent with that of the BR-Sa1 and GF-Guy sites, where young leaves 411 412 increase during the dry seasons. Moreover, the seasonality of the  $V_{c,max25}$  of young leaves in subtropical Asia 413 (Fig. 8f) mirrored patterns observed at the CN-Din site, where young leaves conversely increase during the 414 wet seasons. The  $V_{c.max25}$  of young leaves peaked in July in sub-region R10, which was located between sub-415 regions R8 and R9, where the  $V_{c,max25}$  of young leaf exhibited a bidirectional phenology (Fig. 8j). The four equatorial sub-regions (R1, R2, R7, and R8) exhibited distinct phenological patterns compared to areas near 416 417 Tropic of Capricorn and Cancer. These equatorial regions demonstrated significantly dampened seasonal 418 variation in the  $V_{c,max25}$  of young leaves, with characteristic bidirectional peaks occurring in March and August (Fig. 8a,d,e,g). This bimodal pattern contrasts sharply with the unimodal seasonality observed at 419 420 regions far from equator, consistent with previous findings by Li et al. (2021a). 421

# 422 **3.4** Partial correlations between the seasonal $V_{c,max25}$ of young leaves and individual 423 climatic factors

To assess the climatic controls on  $V_{c,max25}$  of young leaves, we performed spatial partial correlation 424 analyses on climate drivers such as vapor pressure deficit (VPD), air temperature (T<sub>mean</sub>), and downward 425 shortwave solar radiation (SW) (Fig. 9), previously established as critical determinants of leaf phenology in 426 427 TEFs (Yang et al., 2023; Yang et al., 2021; Li et al., 2021a). The V<sub>c,max25</sub> of young leaves exhibited a strong 428 correlation with the three climate drivers (Fig. 9). We then analyzed the relative importance of three climate 429 drivers in influencing  $V_{c,max25}$  using the machine-learning model of the Random Forests (RF) method (Fig. 430 10, section 2.5.3). Shortwave radiation exhibited particularly notable positive correlations (R>0.34) with 431  $V_{c,max25}$  across almost all regions with the exception of Amazon sub-region R4 (R=0.17) (Fig. S8), and the 432 shortwave radiation was also the most contributing factor (Fig. 10a). This underscoring the dominant role of 433 radiation in regulating canopy photosynthesis in TEFs. Although seasonal temperature fluctuations were modest (Fig. S7), likely due to minor temperature gradients,  $T_{mean}$  still exhibited a positive correlation with 434  $V_{c,max25}$  of young leaves. However, at the global scale, T<sub>mean</sub> had the least influence compared to VPD and 435 SW (Fig. 10a). Notably, in the Asia region,  $T_{mean}$  emerged as the primary driver of  $V_{c,max25}$  variability and 436 437 showed a strong positive correlation in the Asia sub-region R10 (R = 0.88, Fig. S8). Notably, VPD and  $T_{mean}$ 

- in sub-region Congo R7 (R=-0.70) and  $T_{mean}$  in sub-region Congo R6 (R=-0.64) (**Fig. S8**). These two factors primarily governed the spatial variability of  $V_{c,max25}$  across the Congo (**Fig. 10 c**). This variability primarily stems from the canopy turnover patterns, where leaf aging during rainy seasons reverses during dry periods (Li et al., 2021a; Yang et al., 2023; Yang et al., 2021). As a result, the seasonality of leaf photosynthetic capacity tended to show an inverse trend to the seasonality of the leaf age, as expected Chen et al. (2020).
- 444

# 445 **3.5** Evaluating potential uncertainties in the *V<sub>c,max25</sub>* of young leaves

The seasonal variations in the  $V_{c,max25}$  of young leaves using 4×4 neighboring pixels were closely aligned with those observed in the 0.25° products utilizing a grid of 2×2 pixels (**Fig. S9**). Results showed a highly linear correlation between the simulated 0.25° resolution and 0.5° resolution consistent (R>0.99), with the root mean square error (RMSE) being maintained below 0.66 (**Fig. 11**). This evidence demonstrating that the neighbor-based decomposition approach reliably generates the consistent  $V_{c,max25}$  products across different spatial scales.

452 Three distinct versions for the gridded  $V_{c,max25}$  of young leaves products from RTSIF- and GOSIF-453 derived GPP and FLUXCOM GPP at various spatial resolutions (Figs. S10-S12) were produced in this study. 454 While minor differences existed among these products, they showed strong spatial consistency (Fig. 12) and high similarity in geographic distribution patterns (R: 0.87~0.96, P<0.001; Fig. 13). All three GPP-derived 455 456  $V_{c,max25}$  products exhibited consistent seasonal patterns across the ten sub-regions (Fig. 12). Validation against 457 in situ measurements demonstrated that the RTSIF-derived product achieved optimal performance, showing both the highest correlation (R=0.85) and minimal deviation (RMSE=13.69) with ground observations (Fig. 458 459 13). These results collectively indicate that the  $V_{c,max25}$  of young leaves products reliably capture 460 photosynthetic seasonality across the ten sub-regions.

# 461462 **4. Discussion**

463 Tropical forests, marked by no obvious seasonal shifts in greenness but distant variations in leaf age cohorts (Wu et al., 2016; Chen et al., 2020; Chavana-Bryant et al., 2017), show distinct leaf phenology 464 compared with temperate and boreal forests. The young leaves are the main leaf cohort to influence 465 466 photosynthesis (Oliveira et al., 2023; Sharma et al., 2017; Menezes et al., 2022), as photosynthetic capacity declines with leaf aging (Menezes et al., 2022; Wang et al., 2020). Understanding the leaf age-photosynthesis 467 468 relationships is therefore critical for assessing plant growth, ecosystem productivity, and carbon cycling in 469 evergreen forests (Albert et al., 2018). The leaf maximum carboxylation capacity ( $V_{c,max25}$ ) serves as a key parameter for modeling photosynthetic CO<sub>2</sub> absorption in Earth System Models (ESMs). However, most 470 471 ESMs typically employ static or annual mean  $V_{c,max25}$  values for each plant functional type (Stocker et al., 472 2020; Atkin et al., 2015). This empirical practice causes uncertainties in tropical forest biomes, which are 473 characterized by their extensive plant functional diversity (Echeverría-Londoño et al., 2018; Spicer et al., 2022) and variable photosynthetic capacity (Piao et al., 2019; Wu et al., 2017a). Furthermore, the  $V_{c,max25}$ 474 475 varies substantially within species due to leaf age, ambient growth temperatures, and the availability of water 476 and nutrients (Stefanski et al., 2020; Lu et al., 2022; Crous et al., 2022). Photosynthesis seasonality in tropical 477 evergreen forests is thus impacted by the replacement of old leaves with younger and more photosynthetically active foliage (Wu et al., 2016; Chen et al., 2020; Chavana-Bryant et al., 2017). These findings underscore 478 the importance of accurately quantifying the leaf age and integrating the leaf age information into  $V_{c,max25}$ 479 estimation to enhance simulation of the leaf CO2 assimilation in tropical forests. Currently, there is no 480 481 comprehensive continental-scale data are available on the leaf age-dependent  $V_{c,max25}$  variations throughout tropical evergreen forests. This data gap remains because the insufficient field observations for adequate 482 mapping (Hakala et al., 2019) and limitations in Earth System Models (ESMs) that rely on uncertain climatic 483

parameters (Brunner et al., 2021). These challenges hinder the application of remote sensing and land surface
 models (LSMs) to accurately model the seasonality of large-scale photosynthesis (Krause et al., 2022).

486 This study presents a novel continental-scale monthly gridded  $V_{c,max25}$  of young leaves. The newly developed dataset was validated at four sites (CN-Din in southern China, MDJ-03 in Congo, BR-Sa1, and 487 GF-Guy in South America) using the field measurements of the  $V_{c,max25}$ . We assessed the reliability of the 488 gridded V<sub>c,max25</sub> of young leaf across the entire TEFs through pixel-by-pixel validation against GPP-derived 489 estimates using the dissolved method and leaf age data from Yang et al. (2023). The results reveal substantial 490 age-dependent variation in  $V_{c.max25}$  (40-90 µmol m<sup>-2</sup> s<sup>-1</sup>), consistent with the ranges reported for tropical and 491 subtropical regions in current Earth System Models (Rogers, 2014). These findings highlight the necessity 492 493 of incorporating leaf-age information in future ESM designs. Moreover, the  $V_{c,max25}$  estimates successfully 494 captured the dry-season canopy greening patterns in the north of the equator, demonstrating prominent 495 advances in our ability to promptly monitor the photosynthetic capacity in tropical forests. Both direct and indirect evaluations confirm the robustness of these new photosynthetic products. In equatorial regions with 496 high annual rainfall and minimal dry seasons, canopy phenology exhibits subtler variations compared to 497 498 those forests near Tropic of Capricorn and Cancer (Yang et al., 2021). The new photosynthetic product 499 successfully captures the characteristic bimodal patterns of  $V_{c,max25}$  with limited seasonal amplitude in these areas. To converts the SIF data into GPP, a constant coefficient was used, and  $V_{c,max25}$  was assumed to be 500 501 uniformly distributed across all tropical evergreen forests, potentially introducing further uncertainties. This 502 assumption was reflected in the MSD assessment, where the bias component was predominant, especially 503 near the equator. Nevertheless, the impact of this on the seasonality of photosynthesis was minima; because 504 the phase-dependent component of the RMSE remained relatively insignificant.

The "leaf demographic-identical (LDO)" hypothesis categorizes the leaf cohorts into three distinct age 505 classes: new leaves (from 1 to 60 days), mature leaves (from 60 to 180 days), and old leaves (larger than 180 506 507 days), with corresponding mean  $V_{c,max25}$  values as reported by Wu et al. (2016). To enhance comparability 508 between observations and models, we further grouped leaves into two age classes. Leaf ages greater than 6 509 months are classified as a distinct old leaf class, while leaf ages less than 6 months are combined into a single 510 young leaf class. Menezes et al. (2022) reported that mature leaves (60-180 days) exhibited the highest 511 average  $V_{c,max25}$ , whereas older leaves (234–612 days) showed lower values (30.4 ± 1.2). The young leaves displayed a 23% higher  $V_{c,max25}$  than old leaves, with minimal variation in the latter. Notably, the link between 512 the older leaves and  $V_{c,max25}$  remains poorly understood in TEFs due to limited field data (Chen et al., 2020). 513 514 To address these simulation challenges, we treated  $V_{c,max25}$  of old leaves as a static value; potentially 515 introducing errors in photosynthetic performance predictions. This simplification may also affect the accuracy of  $V_{c,max25}$  and GPP seasonality in ESMs (De Weirdt et al., 2012). Moreover, additional uncertainties 516 517 stem from assumptions that neglect the spatial and temporal variations driven by the plant functional type diversity, which shifts with seasonal climate anomalies and high heterogeneity in diverse forest ecosystems. 518 519 These generalizations could also introduce inaccuracies in simulating seasonal variations in  $V_{c,max25}$ . Reflecting the inherent variability in photosynthetic behavior across leaf ages, the data revealed two distinct 520 responses: (1) certain species, such as P. tomentosa and P. caimito, exhibited marked reductions in  $V_{c,max25}$ 521 with age, whereas (2) others, such as M. angularis and V. parviflora, maintained consistent  $V_{c,max25}$  values 522 523 after reaching their peak. Menezes et al. (2022) identified a modest but significant correlation between the  $V_{c,max25}$  and leaf age due to these divergent patterns. Variations in the photosynthetic capacity at the ecosystem 524 level could be influenced by species composition and the distribution of plant functional groups within forests. 525 526 Furthermore, the seasonal fluctuations in  $V_{c,max25}$  of young leaves are closely associated with both plant

527 growth strategies and environmental factors. Higher  $V_{c,max25}$  values in young leaves during the early growing

season may reflect an adaptive strategy to quickly establish photosynthetic capacity, especially beneficial in competitive environments like tropical and subtropical forests. These seasonal variations directly impact a plant's carbon uptake capacity, potentially leading to increased carbon sequestration within plant biomass and influencing atmospheric  $CO_2$  concentrations, which could create feedback loops within the climate system.

533 In summary, we present a novel approach to develop a gridded dataset that incorporates leaf-age sensitivity into the photosynthesis parameters for TEFs at a continental scale. While some uncertainties 534 persist, we provide a monthly gridded  $V_{c,max25}$  of young leaves dataset. This innovation facilitates the 535 comprehensive phenological modeling in ESMs, which typically operate at coarser resolutions. These 536 improvements substantially enhance our ability to monitor and mechanistically interpret the spatiotemporal 537 variations in the  $V_{c,max25}$  of young leaves, providing essential data for the parameterization and assessment in 538 ESMs. Furthermore, as remote sensing technologies advance, we anticipate that the enhanced temporal and 539 spatial resolution of RTSIF-derived GPP will facilitate more accurate mapping of the photosynthesis products 540 541 in future studies.

542

549

# 543 **5. Data availability**

The 0.25 degree time-series  $V_{c,max25}$  data from 2001-2018 is presented in this paper as the main dataset. We also provided another two versions of  $V_{c,max25}$  generated from GOSIF-derived GPP and FLUXCOM GPP, respectively. The three datasets are available at https://doi.org/10.5281/zenodo.14807414 (Yang et al., 2025). These datasets are compressed in a GeoTiff format, with a spatial reference of WGS84. Each file in these datasets is named as follows: "V<sub>cmax25</sub> {GPP source}derived\_{YYYYMM}.tif".

#### 550 6. Conclusions

551 This study develop a novel monthly gridded dataset of  $V_{c,max25}$  in combination with ontogeny-dependent leaf age changes. The new  $V_{c,max25}$  of young leaves performs reasonably well in validating against three 552 independent datasets: including (1) in situ observations of the monthly  $V_{c,max25}$  records; (2) the  $V_{c,max25}$  product 553 dissolved from the GOSIF-derived GPP; (3) the leaf-age-dependent leaf area index product. Our results 554 555 demonstrate that the seasonal dynamics in V<sub>c,max25</sub> of young leaves are governed by distinct climatephenology regimes across tropical and subtropical evergreen broadleaved forests. Specifically, in the central 556 557 and southern Amazon, the V<sub>c,max25</sub> of young leaves decreased during dry season onset (approximately 558 February) but increased during wet season onset (approximately June). Conversely, the  $V_{c,max25}$  of young 559 leaves in subtropical Asia exhibited peak during the wet season (June or July), coinciding with maximum 560 rainfall. Near the equator, the  $V_{c.max25}$  of young leaves showed a bimodal seasonality with very slight 561 variations. The  $V_{c,max25}$  of young leaves products offer valuable insights into the adaptations of tropical and 562 subtropical forest to the ongoing climate change, while also serving to improve phenology parameterization in land surface models (LSMs). 563

564

565 **Supplement.** The supplement related to this article will be available online at once accepted.

566

567 Author contributions. XC designed the research and wrote the paper. XY and QS wrote the draft,
568 debugged algorithms and processed data. LH debugged algorithms and reviewed the paper. All the authors
569 edited and revised the paper.

- 570
- 571 **Competing interests.** The authors declare no competing interests.
- 572

573 Acknowledgement. We would like to thank the editor and reviewers for their valuable time in reviewing574 the manuscript.

575

## 576 **References**

- Albert, L. P., Wu, J., Prohaska, N., de Camargo, P. B., Huxman, T. E., Tribuzy, E. S., Ivanov, V. Y., Oliveira,
  R. S., Garcia, S., Smith, M. N., Oliveira Junior, R. C., Restrepo-Coupe, N., da Silva, R., Stark, S. C.,
  Martins, G. A., Penha, D. V., and Saleska, S. R.: Age-dependent leaf physiology and consequences for
  crown-scale carbon uptake during the dry season in an Amazon evergreen forest, New Phytol., 219, 870884, https://doi.org/10.1111/nph.15056, 2018.
- Ali, A. A., Xu, C., Rogers, A., Fisher, R. A., Wullschleger, S. D., Massoud, E., Vrugt, J. A., Muss, J. D.,
  McDowell, N. G., and Fisher, J. B.: A global scale mechanistic model of photosynthetic capacity (LUNA
  V1. 0), Geosci. Model Dev., 9, 587-606, https://doi.org/10.5194/gmd-9-587-2016, 2016.
- Arora, V. K. and Boer, G. J.: Fire as an interactive component of dynamic vegetation models, J. Geophys.
  Res. Biogeosci., 110, G02008, https://doi.org/10.1029/2005jg000042, 2005.
- Atkin, O. K., Meir, P., and Turnbull, M. H.: Improving representation of leaf respiration in large-scale
  predictive climate-vegetation models, New Phytol., 202, 743-748, https://doi.org/10.1111/nph.12686,
  2014.
- Atkin, O. K., Bloomfield, K. J., Reich, P. B., Tjoelker, M. G., Asner, G. P., Bonal, D., Boenisch, G., Bradford,
  M. G., Cernusak, L. A., Cosio, E. G., Creek, D., Crous, K. Y., Domingues, T. F., Dukes, J. S., Egerton,
  J. J. G., Evans, J. R., Farquhar, G. D., Fyllas, N. M., Gauthier, P. P. G., Gloor, E., Gimeno, T. E., Griffin,
- 593 K. L., Guerrieri, R., Heskel, M. A., Huntingford, C., Ishida, F. Y., Kattge, J., Lambers, H., Liddell, M.
- J., Lloyd, J., Lusk, C. H., Martin, R. E., Maksimov, A. P., Maximov, T. C., Malhi, Y., Medlyn, B. E.,
- 595 Meir, P., Mercado, L. M., Mirotchnick, N., Ng, D., Niinemets, U., O'Sullivan, O. S., Phillips, O. L.,
- 596Poorter, L., Poot, P., Prentice, I. C., Salinas, N., Rowland, L. M., Ryan, M. G., Sitch, S., Slot, M., Smith,
- N. G., Turnbull, M. H., VanderWel, M. C., Valladares, F., Veneklaas, E. J., Weerasinghe, L. K., Wirth,
  C., Wright, I. J., Wythers, K. R., Xiang, J., Xiang, S., and Zaragoza-Castells, J.: Global variability in
  leaf respiration in relation to climate, plant functional types and leaf traits, New Phytol., 206, 614-636,
  https://doi.org/10.1111/nph.13253, 2015.
- Bernacchi, C. J., Pimentel, C., and Long, S. P.: *In vivo* temperature response functions of parameters required
  to model RuBP-limited photosynthesis, Plant Cell and Environment, 26, 1419-1430,
  https://doi.org/10.1046/j.0016-8025.2003.01050.x, 2003.
- Bernacchi, C. J., Singsaas, E. L., Pimentel, C., Portis, A. R., and Long, S. P.: Improved temperature response
  functions for models of Rubisco-limited photosynthesis, Plant Cell and Environment, 24, 253-259,
  https://doi.org/10.1046/j.1365-3040.2001.00668.x, 2001.
- Bernacchi, C. J., Bagley, J. E., Serbin, S. P., Ruiz-Vera, U. M., Rosenthal, D. M., and Vanloocke, A.:
  Modelling C<sub>3</sub> photosynthesis from the chloroplast to the ecosystem, Plant Cell and Environment, 36,
  1641-1657, https://doi.org/10.1111/pce.12118, 2013.
- Brando, P. M., Goetz, S. J., Baccini, A., Nepstad, D. C., Beck, P. S. A., and Christman, M. C.: Seasonal and 610 611 interannual variability of climate and vegetation indices across the Amazon, Proceedings of the National States 612 Academy of Sciences of the United of America. 107. 14685-14690. https://doi.org/10.1073/pnas.0908741107, 2010. 613
- Brunner, M. I., Slater, L., Tallaksen, L. M., and Clark, M.: Challenges in modeling and predicting floods and
  droughts: A review, Wiley Interdiscip. Rev.: Water, 8, e1520, https://doi.org/10.1002/wat2.1520, 2021.
- 616 Chavana-Bryant, C., Malhi, Y., Wu, J., Asner, G. P., Anastasiou, A., Enquist, B. J., Cosio Caravasi, E. G.,
- Doughty, C. E., Saleska, S. R., Martin, R. E., and Gerard, F. F.: Leaf aging of Amazonian canopy trees

- as revealed by spectral and physiochemical measurements, New Phytol., 214, 1049-1063,
  https://doi.org/10.1111/nph.13853, 2017.
- Chen, J. M., Ju, W., Ciais, P., Viovy, N., Liu, R., Liu, Y., and Lu, X.: Vegetation structural change since 1981
  significantly enhanced the terrestrial carbon sink, Nat. Commun., 10, https://doi.org/10.1038/s41467019-12257-8, 2019.
- Chen, J. M., Wang, R., Liu, Y. H., He, L. M., Croft, H., Luo, X. Z., Wang, H., Smith, N. G., Keenan, T. F.,
  Prentice, I. C., Zhang, Y. G., Ju, W. M., and Dong, N.: Global datasets of leaf photosynthetic capacity
  for ecological and earth system research, Earth Syst. Sci. Data, 14, 4077-4093,
  https://doi.org/10.5194/essd-14-4077-2022, 2022a.
- 627 Chen, X., Huang, Y., Nie, C., Zhang, S., Wang, G., Chen, S., and Chen, Z.: A long-term reconstructed
   628 TROPOMI solar-induced fluorescence dataset using machine learning algorithms, Sci. Data, 9, 427,
   629 https://doi.org/10.1038/s41597-022-01520-1, 2022b.
- Chen, X., Maignan, F., Viovy, N., Bastos, A., Goll, D., Wu, J., Liu, L. Y., Yue, C., Peng, S. S., Yuan, W. P.,
  da Conceicao, A. C., O'Sullivan, M., and Ciais, P.: Novel Representation of Leaf Phenology Improves
  Simulation of Amazonian Evergreen Forest Photosynthesis in a Land Surface Model, J. Adv. Model.
  Earth Syst., 12, e2018MS001565, https://doi.org/10.1029/2018ms001565, 2020.
- Chen, X., Ciais, P., Maignan, F., Zhang, Y., Bastos, A., Liu, L. Y., Bacour, C., Fan, L., Gentine, P., Goll, D.,
  Green, J., Kim, H., Li, L., Liu, Y., Peng, S. S., Tang, H., Viovy, N., Wigneron, J. P., Wu, J., Yuan, W. P.,
  and Zhang, H. C.: Vapor Pressure Deficit and Sunlight Explain Seasonality of Leaf Phenology and
  Photosynthesis Across Amazonian Evergreen Broadleaved Forest, Global Biogeochem. Cycles, 35,
  e2018MS001565, https://doi.org/10.1029/2020gb006893, 2021.
- Chou, S., Chen, B., Chen, J., Wang, M., Wang, S., Croft, H., and Shi, Q.: Estimation of leaf photosynthetic
  capacity from the photochemical reflectance index and leaf pigments, Ecol. Indic., 110, 105867,
  https://doi.org/10.1016/j.ecolind.2019.105867, 2020.
- Cramer, W., Bondeau, A., Woodward, F. I., Prentice, I. C., Betts, R. A., Brovkin, V., Cox, P. M., Fisher, V.,
  Foley, J. A., Friend, A. D., Kucharik, C., Lomas, M. R., Ramankutty, N., Sitch, S., Smith, B., White, A.,
  and Young-Molling, C.: Global response of terrestrial ecosystem structure and function to CO<sub>2</sub> and
  climate change:: results from six dynamic global vegetation models, Global Change Biol., 7, 357-373,
  https://doi.org/10.1046/j.1365-2486.2001.00383.x, 2001.
- 647 Croft, H., Chen, J. M., Luo, X., Bartlett, P., Chen, B., and Staebler, R. M.: Leaf chlorophyll content as a proxy
  648 for leaf photosynthetic capacity, Global Change Biol., 23, 3513-3524, https://doi.org/10.1111/gcb.13599,
  649 2017.
- Croft, H., Chen, J., Wang, R., Mo, G., Luo, S., Luo, X., He, L., Gonsamo, A., Arabian, J., and Zhang, Y.: The
  global distribution of leaf chlorophyll content, Remote Sens. Environ., 236, 111479,
  https://doi.org/10.1016/j.rse.2019.111479, 2020.
- Crous, K. Y., Uddling, J., and De Kauwe, M. G.: Temperature responses of photosynthesis and respiration in 653 from boreal to tropical latitudes, New Phytol., 654 evergreen trees 234, 353-374, 655 https://doi.org/10.1111/nph.17951, 2022.
- De Weirdt, M., Verbeeck, H., Maignan, F., Peylin, P., Poulter, B., Bonal, D., Ciais, P., and Steppe, K.: Seasonal
  leaf dynamics for tropical evergreen forests in a process-based global ecosystem model, Geosci. Model
  Dev., 5, 1091-1108, https://doi.org/10.5194/gmd-5-1091-2012, 2012.
- Dechant, B., Ryu, Y., Badgley, G., Zeng, Y., Berry, J. A., Zhang, Y., Goulas, Y., Li, Z., Zhang, Q., Kang, M.,
  Li, J., and Moya, I.: Canopy structure explains the relationship between photosynthesis and sun-induced
  chlorophyll fluorescence in crops, Remote Sens. Environ., 241, 111733,

- 662 https://doi.org/10.1016/j.rse.2020.111733, 2020.
- Echeverría-Londoño, S., Enquist, B. J., Neves, D. M., Violle, C., Boyle, B., Kraft, N. J., Maitner, B. S.,
  McGill, B., Peet, R. K., and Sandel, B.: Plant functional diversity and the biogeography of biomes in
  North and South America, Front. Ecol. Evol., 6, 219, https://doi.org/10.3389/fevo.2018.00219, 2018.
- Evans, J. R.: Photosynthesis and Nitrogen Relationships in Leaves of C<sub>3</sub> Plants, Oecologia, 78, 9-19,
   https://doi.org/10.1007/BF00377192, 1989.
- Fabre, D., Yin, X., Dingkuhn, M., Clément-Vidal, A., Roques, S., Rouan, L., Soutiras, A., and Luquet, D.: Is
  triose phosphate utilization involved in the feedback inhibition of photosynthesis in rice under
  conditions of sink limitation?, J. Exp. Bot., 70, 5773-5785, https://doi.org/10.1093/jxb/erz318, 2019.
- Farquhar, G. D., Caemmerer, S. V., and Berry, J. A.: A biochemical-model of photosynthetic CO<sub>2</sub> assimilation
  in leaves of C<sub>3</sub> specials, Planta, 149, 78-90, https://doi.org/10.1007/bf00386231, 1980.
- Ferreira Domingues, T., Ishida, F. Y., Feldpausch, T. R., Grace, J., Meir, P., Saiz, G., Sene, O., Schrodt, F.,
  Sonké, B., and Taedoumg, H.: Biome-specific effects of nitrogen and phosphorus on the photosynthetic
  characteristics of trees at a forest-savanna boundary in Cameroon, Oecologia, 178, 659-672,
  https://doi.org/10.1007/s00442-015-3250-5, 2015.
- Frankenberg, C., Fisher, J. B., Worden, J., Badgley, G., Saatchi, S. S., Lee, J. E., Toon, G. C., Butz, A., Jung, 677 678 M., and Kuze, A.: New global observations of the terrestrial carbon cycle from GOSAT: Patterns of fluorescence with 679 plant gross primary productivity, Geophys. Res. Lett., 38, 680 https://doi.org/10.1029/2011GL048738, 2011.
- Hakala, K., Addor, N., Teutschbein, C., Vis, M., Dakhlaoui, H., and Seibert, J.: Hydrological modeling of
  climate change impacts, Encyclopedia of water: Science, technology, and society, 1-20,
  https://doi.org/10.1002/9781119300762.wsts0062, 2019.
- He, L., Chen, J. M., Liu, J., Zheng, T., Wang, R., Joiner, J., Chou, S., Chen, B., Liu, Y., and Liu, R.: Diverse
  photosynthetic capacity of global ecosystems mapped by satellite chlorophyll fluorescence
  measurements, Remote Sens. Environ., 232, 111344, https://doi.org/10.1016/j.rse.2019.111344, 2019.
- Hikosaka, K.: Optimal nitrogen distribution within a leaf canopy under direct and diffuse light, Plant, Cell
  Environ, 37, 2077-2085, https://doi.org/10.1111/pce.12291, 2014.
- Houborg, R., Cescatti, A., Migliavacca, M., and Kustas, W.: Satellite retrievals of leaf chlorophyll and
  photosynthetic capacity for improved modeling of GPP, Agric. For. Meteorol., 177, 10-23,
  https://doi.org/10.1016/j.agrformet.2013.04.006, 2013.
- Houborg, R., McCabe, M. F., Cescatti, A., and Gitelson, A. A.: Leaf chlorophyll constraint on model
  simulated gross primary productivity in agricultural systems, Int. J. Appl. Earth Obs. Geoinf., 43, 160176, https://doi.org/10.1016/j.jag.2015.03.016, 2015.
- Ikotun, A. M., Ezugwu, A. E., Abualigah, L., Abuhaija, B., and Heming, J.: K-means clustering algorithms:
  A comprehensive review, variants analysis, and advances in the era of big data, Information Sciences,
  622, 178-210, https://doi.org/https://doi.org/10.1016/j.ins.2022.11.139, 2023.
- Jensen, A. M., Warren, J. M., Hanson, P. J., Childs, J., and Wullschleger, S. D.: Needle age and season
  influence photosynthetic temperature response and total annual carbon uptake in mature Picea mariana
  trees, Ann. Bot., 116, 821-832, https://doi.org/10.1093/aob/mcv115, 2015.
- Jung, M., Koirala, S., Weber, U., Ichii, K., Gans, F., Camps-Valls, G., Papale, D., Schwalm, C., Tramontana,
  G., and Reichstein, M.: The FLUXCOM ensemble of global land-atmosphere energy fluxes, Sci. Data,
  6, https://doi.org/10.1038/s41597-019-0076-8, 2019.
- Knyazikhin, Y., Schull, M. A., Stenberg, P., Mõttus, M., Rautiainen, M., Yang, Y., Marshak, A., Latorre
   Carmona, P., Kaufmann, R. K., and Lewis, P.: Hyperspectral remote sensing of foliar nitrogen content,

- 706 Proc. Natl Acad. Sci. USA, 110, E185-E192, https://doi.org/10.1073/pnas.1210196109, 2013.
- Krause, A., Papastefanou, P., Gregor, K., Layritz, L. S., Zang, C. S., Buras, A., Li, X., Xiao, J., and Rammig,
  A.: Quantifying the impacts of land cover change on gross primary productivity globally, Sci. Rep., 12,
  18398, https://doi.org/10.1038/s41598-022-23120-0, 2022.
- Li, Q., Chen, X., Yuan, W., Lu, H., Shen, R., Wu, S., Gong, F., Dai, Y., Liu, L., Sun, Q., Zhang, C., and Su,
  Y.: Remote Sensing of Seasonal Climatic Constraints on Leaf Phenology Across Pantropical Evergreen
  Forest Biome, Earths Future, 9, e2021EF002160, https://doi.org/10.1029/2021ef002160, 2021a.
- Li, X., Du, H., Zhou, G., Mao, F., Zhang, M., Han, N., Fan, W., Liu, H., Huang, Z., and He, S.: Phenology
  estimation of subtropical bamboo forests based on assimilated MODIS LAI time series data, ISPRS J.
  Photogramm. Remote Sens., 173, 262-277, https://doi.org/10.1016/j.isprsjprs.2021.01.018, 2021b.
- Lin, Y.-S., Medlyn, B. E., Duursma, R. A., Prentice, I. C., Wang, H., Baig, S., Eamus, D., de Dios, Victor R.,
  Mitchell, P., Ellsworth, D. S., de Beeck, M. O., Wallin, G., Uddling, J., Tarvainen, L., Linderson, M.-L.,
  Cernusak, L. A., Nippert, J. B., Ocheltree, T. W., Tissue, D. T., Martin-StPaul, N. K., Rogers, A., Warren,
  J. M., De Angelis, P., Hikosaka, K., Han, Q., Onoda, Y., Gimeno, T. E., Barton, C. V. M., Bennie, J.,
  Bonal, D., Bosc, A., Löw, M., Macinins-Ng, C., Rey, A., Rowland, L., Setterfield, S. A., Tausz-Posch,
  S., Zaragoza-Castells, J., Broadmeadow, M. S. J., Drake, J. E., Freeman, M., Ghannoum, O., Hutley,
- Lindsay B., Kelly, J. W., Kikuzawa, K., Kolari, P., Koyama, K., Limousin, J.-M., Meir, P., Lola da Costa,
- A. C., Mikkelsen, T. N., Salinas, N., Sun, W., and Wingate, L.: Optimal stomatal behaviour around the
   world, Nat. Clim. Change, 5, 459-464, https://doi.org/10.1038/nclimate2550, 2015.
- Liu, Y., Chen, J. M., Xu, M., Wang, R., Fan, W., Li, W., Kammer, L., Prentice, C., Keenan, T. F., and Smith,
   N. G.: Improved global estimation of seasonal variations in C3 photosynthetic capacity based on eco evolutionary optimality hypotheses and remote sensing, Remote Sens. Environ., 313, 114338,
   https://doi.org/10.1016/j.rse.2024.114338, 2024.
- Locke, A. M. and Ort, D. R.: Leaf hydraulic conductance declines in coordination with photosynthesis,
  transpiration and leaf water status as soybean leaves age regardless of soil moisture, J. Exp. Bot., 65,
  6617-6627, https://doi.org/10.1093/jxb/eru380, 2014.
- Lu, X., Croft, H., Chen, J. M., Luo, Y., and Ju, W.: Estimating photosynthetic capacity from optimized
  Rubisco-chlorophyll relationships among vegetation types and under global change, Environ. Res. Lett.,
  17, 014028, https://doi.org/10.1088/1748-9326/ac444d, 2022.
- Lu, X., Ju, W., Li, J., Croft, H., Chen, J. M., Luo, Y., Yu, H., and Hu, H.: Maximum Carboxylation Rate
  Estimation With Chlorophyll Content as a Proxy of Rubisco Content, J. Geophys. Res. Biogeosci., 125,
  e2020JG005748, https://doi.org/10.1029/2020jg005748, 2020.
- Lu, X., Vitousek, P. M., Mao, Q., Gilliam, F. S., Luo, Y., Turner, B. L., Zhou, G., and Mo, J.: Nitrogen deposition accelerates soil carbon sequestration in tropical forests, Proc. Natl Acad. Sci. USA, 118, e2020790118, https://doi.org/10.1073/pnas.2020790118, 2021.
- Luo, X., Croft, H., Chen, J. M., He, L., and Keenan, T. F.: Improved estimates of global terrestrial
  photosynthesis using information on leaf chlorophyll content, Global Change Biol., 25, 2499-2514,
  https://doi.org/10.1111/gcb.14624, 2019.
- Luo, Y., Medlyn, B., Hui, D., Ellsworth, D., Reynolds, J., and Katul, G.: Gross primary productivity in duke
  forest: modeling synthesis of CO2 experiment and eddy–flux data, Ecol. Appl., 11, 239-252,
  https://doi.org/10.2307/3061070, 2001.
- McClain, A. M. and Sharkey, T. D.: Triose phosphate utilization and beyond: from photosynthesis to end
   product synthesis, J. Exp. Bot., 70, 1755-1766, https://doi.org/10.1093/jxb/erz058, 2019.
- 749 Medlyn, B. E., Duursma, R. A., Eamus, D., Ellsworth, D. S., Prentice, I. C., Barton, C. V. M., Crous, K. Y.,

- de Angelis, P., Freeman, M., and Wingate, L.: Reconciling the optimal and empirical approaches to
  modelling stomatal conductance, Global Change Biol., 17, 2134-2144, https://doi.org/10.1111/j.13652486.2010.02375.x, 2011.
- Menezes, J., Garcia, S., Grandis, A., Nascimento, H., Domingues, T. F., Guedes, A. V., Aleixo, I., Camargo,
  P., Campos, J., Damasceno, A., Dias-Silva, R., Fleischer, K., Kruijt, B., Cordeiro, A. L., Martins, N. P.,
- 755 Meir, P., Norby, R. J., Pereira, I., Portela, B., Rammig, A., Ribeiro, A. G., Lapola, D. M., and Quesada,
- 756 C. A.: Changes in leaf functional traits with leaf age: when do leaves decrease their photosynthetic
- capacity in Amazonian trees?, Tree Physiol., 42, 922-938, https://doi.org/10.1093/treephys/tpab042,
  2022.
- Mohammed, G. H., Colombo, R., Middleton, E. M., Rascher, U., Van Der Tol, C., Nedbal, L., Goulas, Y.,
  Pérez-Priego, O., Damm, A., and Meroni, M.: Remote sensing of solar-induced chlorophyll fluorescence
  (SIF) in vegetation: 50 years of progress, Remote Sens. Environ., 231, 111177,
  https://doi.org/10.1016/j.rse.2019.04.030, 2019.
- Oliveira, F. M., Knoechelmann, C. M., Wirth, R., Tabarelli, M., and Leal, I. R.: Leaf-cutting ant nests support
   less dense and impoverished seed assemblages in a human-modified Caatinga dry forest, Biotropica, 55,
   444-453, https://doi.org/10.1111/btp.13198, 2023.
- Onoda, Y., Wright, I. J., Evans, J. R., Hikosaka, K., Kitajima, K., Niinemets, Ü., Poorter, H., Tosens, T., and
  Westoby, M.: Physiological and structural tradeoffs underlying the leaf economics spectrum, New
  Phytol., 214, 1447-1463, https://doi.org/10.1111/nph.14496, 2017.
- Orndahl, K. M., Ehlers, L. P., Herriges, J. D., Pernick, R. E., Hebblewhite, M., and Goetz, S. J.: Mapping
  tundra ecosystem plant functional type cover, height, and aboveground biomass in Alaska and northwest
  Canada using unmanned aerial vehicles, Arct. Sci., 8, 1165-1180, https://doi.org/10.1088/17489326/ac9d50, 2022.
- 773 Pastorello, G., Trotta, C., Canfora, E., Chu, H., Christianson, D., Cheah, Y.-W., Poindexter, C., Chen, J., Elbashandy, A., Humphrey, M., Isaac, P., Polidori, D., Ribeca, A., van Ingen, C., Zhang, L., Amiro, B., 774 775 Ammann, C., Arain, M. A., Ardo, J., Arkebauer, T., Arndt, S. K., Arriga, N., Aubinet, M., Aurela, M., 776 Baldocchi, D., Barr, A., Beamesderfer, E., Marchesini, L. B., Bergeron, O., Beringer, J., Bernhofer, C., Berveiller, D., Billesbach, D., Black, T. A., Blanken, P. D., Bohrer, G., Boike, J., Bolstad, P. V., Bonal, 777 D., Bonnefond, J.-M., Bowling, D. R., Bracho, R., Brodeur, J., Bruemmer, C., Buchmann, N., Burban, 778 779 B., Burns, S. P., Buysse, P., Cale, P., Cavagna, M., Cellier, P., Chen, S., Chini, I., Christensen, T. R., 780 Cleverly, J., Collalti, A., Consalvo, C., Cook, B. D., Cook, D., Coursolle, C., Cremonese, E., Curtis, P. 781 S., D'Andrea, E., da Rocha, H., Dai, X., Davis, K. J., De Cinti, B., de Grandcourt, A., De Ligne, A., De Oliveira, R. C., Delpierre, N., Desai, A. R., Di Bella, C. M., di Tommasi, P., Dolman, H., Domingo, F., 782 Dong, G., Dore, S., Duce, P., Dufrene, E., Dunn, A., Dusek, J., Eamus, D., Eichelmann, U., ElKhidir, H. 783 A. M., Eugster, W., Ewenz, C. M., Ewers, B., Famulari, D., Fares, S., Feigenwinter, I., Feitz, A., Fensholt, 784 785 R., Filippa, G., Fischer, M., Frank, J., Galvagno, M., Gharun, M., Gianelle, D., Gielen, B., Gioli, B., Gitelson, A., Goded, I., Goeckede, M., Goldstein, A. H., Gough, C. M., Goulden, M. L., Graf, A., Griebel, 786 A., Gruening, C., Gruenwald, T., Hammerle, A., Han, S., Han, X., Hansen, B. U., Hanson, C., Hatakka, 787 J., He, Y., Hehn, M., Heinesch, B., Hinko-Najera, N., Hoertnagl, L., Hutley, L., Ibrom, A., Ikawa, H., 788 Jackowicz-Korczynski, M., Janous, D., Jans, W., Jassal, R., Jiang, S., Kato, T., Khomik, M., Klatt, J., 789 790 Knohl, A., Knox, S., Kobayashi, H., Koerber, G., Kolle, O., Kosugi, Y., Kotani, A., Kowalski, A., Kruijt, B., Kurbatova, J., Kutsch, W. L., Kwon, H., Launiainen, S., Laurila, T., Law, B., Leuning, R., Li, Y., 791 792 Liddell, M., Limousin, J.-M., Lion, M., Liska, A. J., Lohila, A., Lopez-Ballesteros, A., Lopez-Blanco, 793 E., Loubet, B., Loustau, D., Lucas-Moffat, A., Lucers, J., Ma, S., Macfarlane, C., Magliulo, V., Maier,

R., Mammarella, I., Manca, G., Marcolla, B., Margolis, H. A., Marras, S., Massman, W., Mastepanov, 794 M., Matamala, R., Matthes, J. H., Mazzenga, F., McCaughey, H., McHugh, I., McMillan, A. M. S., 795 796 Merbold, L., Meyer, W., Meyers, T., Miller, S. D., Minerbi, S., Moderow, U., Monson, R. K., 797 Montagnani, L., Moore, C. E., Moors, E., Moreaux, V., Moureaux, C., Munger, J. W., Nakai, T., Neirynck, J., Nesic, Z., Nicolini, G., Noormets, A., Northwood, M., Nosetto, M., Nouvellon, Y., Novick, 798 K., Oechel, W., Olesen, J. E., Ourcival, J.-M., Papuga, S. A., Parmentier, F.-J., Paul-Limoges, E., 799 800 Pavelka, M., Peichl, M., Pendall, E., Phillips, R. P., Pilegaard, K., Pirk, N., Posse, G., Powell, T., Prasse, 801 H., Prober, S. M., Rambal, S., Rannik, U., Raz-Yaseef, N., Reed, D., de Dios, V. R., Restrepo-Coupe, 802 N., Reverter, B. R., Roland, M., Sabbatini, S., Sachs, T., Saleska, S. R., Sanchez-Canete, E. P., Sanchez-Mejia, Z. M., Schmid, H. P., Schmidt, M., Schneider, K., Schrader, F., Schroder, I., Scott, R. L., Sedlak, 803 P., Serrano-Ortiz, P., Shao, C., Shi, P., Shironya, I., Siebicke, L., Sigut, L., Silberstein, R., Sirca, C., 804 Spano, D., Steinbrecher, R., Stevens, R. M., Sturtevant, C., Suyker, A., Tagesson, T., Takanashi, S., Tang, 805 Y., Tapper, N., Thom, J., Tiedemann, F., Tomassucci, M., Tuovinen, J.-P., Urbanski, S., Valentini, R., 806 807 van der Molen, M., van Gorsel, E., van Huissteden, K., Varlagin, A., Verfaillie, J., Vesala, T., Vincke, C., Vitale, D., Vygodskaya, N., Walker, J. P., Walter-Shea, E., Wang, H., Weber, R., Westermann, S., 808 Wille, C., Wofsy, S., Wohlfahrt, G., Wolf, S., Woodgate, W., Li, Y., Zampedri, R., Zhang, J., Zhou, G., 809 Zona, D., Agarwal, D., Biraud, S., Torn, M., and Papale, D.: The FLUXNET2015 dataset and the 810 811 **ONEFlux** processing pipeline for eddy covariance data, Sci. Data, 7, 225, 812 https://doi.org/10.1038/s41597-020-0534-3, 2020.

- Piao, S., Liu, Q., Chen, A., Janssens, I. A., Fu, Y., Dai, J., Liu, L., Lian, X., Shen, M., and Zhu, X.: Plant
  phenology and global climate change: Current progresses and challenges, Global Change Biol., 25,
  1922-1940, https://doi.org/10.1111/gcb.14619, 2019.
- Quebbeman, J. and Ramirez, J.: Optimal allocation of leaf-level nitrogen: Implications for covariation of
   V<sub>cmax</sub> and J<sub>max</sub> and photosynthetic downregulation, J. Geophys. Res. Biogeosci., 121, 2464-2475,
   https://doi.org/10.1002/2016JG003473, 2016.
- Rogers, A.: The use and misuse of V<sub>c,max</sub> in Earth System Models, Photosynth. Res., 119, 15-29, https://doi.org/10.1007/s11120-013-9818-1, 2014.
- Ryu, Y., Jiang, C., Kobayashi, H., and Detto, M.: MODIS-derived global land products of shortwave radiation
  and diffuse and total photosynthetically active radiation at 5 km resolution from 2000, Remote Sens.
  Environ., 204, 812-825, https://doi.org/10.1016/j.rse.2017.09.021, 2018.
- Sharma, U., Kataria, V., and Shekhawat, N.: In vitro propagation, ex vitro rooting and leaf micromorphology
  of Bauhinia racemosa Lam.: a leguminous tree with medicinal values, Physiol. Mol. Biol. Plants, 23,
  969-977, https://doi.org/10.1007/s12298-017-0459-2, 2017.
- Song, G., Wang, Q., and Jin, J.: Leaf photosynthetic capacity of sunlit and shaded mature leaves in a
  deciduous forest, Forests, 11, 318, https://doi.org/10.3390/f11030318, 2020.
- Spicer, M. E., Radhamoni, H. V. N., Duguid, M. C., Queenborough, S. A., and Comita, L. S.: Herbaceous
  plant diversity in forest ecosystems: patterns, mechanisms, and threats, Plant Ecolog., 223, 117-129,
  https://doi.org/10.1007/s11258-021-01202-9, 2022.
- Stefanski, A., Bermudez, R., Sendall, K. M., Montgomery, R. A., and Reich, P. B.: Surprising lack of
  sensitivity of biochemical limitation of photosynthesis of nine tree species to open-air experimental
  warming and reduced rainfall in a southern boreal forest, Global Change Biol., 26, 746-759,
  https://doi.org/10.1111/gcb.14805, 2020.
- 836 Stocker, B. D., Wang, H., Smith, N. G., Harrison, S. P., Keenan, T. F., Sandoval, D., Davis, T., and Prentice,
- 837 I. C.: P-model v1. 0: An optimality-based light use efficiency model for simulating ecosystem gross

- primary production, Geosci. Model Dev., 13, 1545-1581, https://doi.org/10.5194/gmd-13-1545-2020,
  2020.
- Sulla-Menashe, D., Woodcock, C. E., and Friedl, M. A.: Canadian boreal forest greening and browning trends:
  an analysis of biogeographic patterns and the relative roles of disturbance versus climate drivers,
  Environ. Res. Lett., 13, 014007, https://doi.org/10.1088/1748-9326/aa9b88, 2018.
- Sun, J., Sun, J., and Feng, Z.: Modelling photosynthesis in flag leaves of winter wheat (Triticum aestivum)
  considering the variation in photosynthesis parameters during development, Funct. Plant Biol., 42,
  1036-1044, https://doi.org/10.1071/FP15140, 2015.
- Tang, H., and Dubayah, R.: Light-driven growth in Amazon evergreen forests explained by seasonal
  variations of vertical canopy structure. Proc. Natl Acad. Sci. USA, 114(10): 2640-2644. 2017.
- Urban, O., Šprtová, M., Košvancová, M., Tomášková, I., Lichtenthaler, H. K., and Marek, M. V.: Comparison 848 of photosynthetic induction and transient limitations during the induction phase in young and mature 849 leaves from three poplar clones, Tree Physiol., 28, 1189-1197, 850 851 https://doi.org/10.1093/treephys/28.8.1189, 2008.
- Verheijen, L., Brovkin, V., Aerts, R., Boenisch, G., Cornelissen, J., Kattge, J., Reich, P. B., Wright, I. J., and
  Van Bodegom, P.: Impacts of trait variation through observed trait–climate relationships on performance
  of an Earth system model: a conceptual analysis, Biogeosciences, 10, 5497-5515,
  https://doi.org/10.5194/bg-10-5497-2013, 2013.
- Wang, S., Li, Y., Ju, W., Chen, B., Chen, J., Croft, H., Mickler, R. A., and Yang, F.: Estimation of Leaf
  Photosynthetic Capacity From Leaf Chlorophyll Content and Leaf Age in a Subtropical Evergreen
  Coniferous Plantation, J. Geophys. Res. Biogeosci., 125, e2019JG005020,
  https://doi.org/10.1029/2019jg005020, 2020.
- Wang, X., Chen, J. M., Ju, W., and Zhang, Y.: Seasonal variations in leaf maximum photosynthetic capacity
  and its dependence on climate factors across global FLUXNET sites, J. Geophys. Res. Biogeosci., 127,
  e2021JG006709, https://doi.org/10.1029/2021JG006709, 2022.
- Weiss, A. and Norman, J. M.: Partitioning solar radiation into direct and diffuse, visible and near-infrared
   components, Agric. For. Meteorol., 34, 205-213, https://doi.org/10.1016/0168-1923(85)90020-6, 1985.
- Wu, J., Serbin, S. P., Xu, X., Albert, L. P., Chen, M., Meng, R., Saleska, S. R., and Rogers, A.: The phenology
  of leaf quality and its within-canopy variation is essential for accurate modeling of photosynthesis in
  tropical evergreen forests, Global Change Biol., 23, 4814-4827, https://doi.org/10.1111/gcb.13725,
  2017a.
- Wu, J., Guan, K., Hayek, M., Restrepo-Coupe, N., Wiedemann, K. T., Xu, X., Wehr, R., Christoffersen, B.
  O., Miao, G., da Silva, R., de Araujo, A. C., Oliviera, R. C., Camargo, P. B., Monson, R. K., Huete, A.
  R., and Saleska, S. R.: Partitioning controls on Amazon forest photosynthesis between environmental
  and biotic factors at hourly to interannual timescales, Global Change Biol., 23, 1240-1257,
  https://doi.org/10.1111/gcb.13509, 2017b.
- Wu, J., Kobayashi, H., Stark, S. C., Meng, R., Guan, K., Tran, N. N., Gao, S., Yang, W., Restrepo-Coupe, N.,
  Miura, T., Oliviera, R. C., Rogers, A., Dye, D. G., Nelson, B. W., Serbin, S. P., Huete, A. R., and Saleska,
  S. R.: Biological processes dominate seasonality of remotely sensed canopy greenness in an Amazon
  evergreen forest, New Phytol., 217, 1507-1520, https://doi.org/10.1111/nph.14939, 2018.
- Wu, J., Albert, L. P., Lopes, A. P., Restrepo-Coupe, N., Hayek, M., Wiedemann, K. T., Guan, K., Stark, S. C.,
  Christoffersen, B., Prohaska, N., Tavares, J. V., Marostica, S., Kobayashi, H., Ferreira, M. L., Campos,
  K. S., da Silva, R., Brando, P. M., Dye, D. G., Huxman, T. E., Huete, A. R., Nelson, B. W., and Saleska,
- 881 S. R.: Leaf development and demography explain photosynthetic seasonality in Amazon evergreen

- forests, Science, 351, 972-976, https://doi.org/10.1126/science.aad5068, 2016.
- Xu, M., Liu, R., Chen, J. M., Liu, Y., Wolanin, A., Croft, H., He, L., Shang, R., Ju, W., and Zhang, Y.: A 21year time series of global leaf chlorophyll content maps from MODIS imagery, IEEE Trans. Geosci.
  Remote Sens., 60, 1-13, https://doi.org/10.1109/TGRS.2022.3204185, 2022a.
- Xu, M., Liu, R., Chen, J. M., Shang, R., Liu, Y., Qi, L., Croft, H., Ju, W., Zhang, Y., and He, Y.: Retrieving
  global leaf chlorophyll content from MERIS data using a neural network method, ISPRS J. Photogramm.
  Remote Sens., 192, 66-82, https://doi.org/10.1016/j.isprsjprs.2022.08.003, 2022b.
- Xu, X., Medvigy, D., Joseph Wright, S., Kitajima, K., Wu, J., Albert, L. P., Martins, G. A., Saleska, S. R.,
  and Pacala, S. W.: Variations of leaf longevity in tropical moist forests predicted by a trait-driven carbon
  optimality model, Ecol. Lett., 20, 1097-1106, https://doi.org/10.1111/ele.12804, 2017.
- Yang, H., Ciais, P., Wigneron, J.-P., Chave, J., Cartus, O., Chen, X., Fan, L., Green, J. K., Huang, Y., Joetzjer,
  E., Kay, H., Makowski, D., Maignan, F., Santoro, M., Tao, S., Liu, L., & Yao, Y.: Climatic and biotic
  factors influencing regional declines and recovery of tropical forest biomass from the 2015/16 El Niño.
  Proc. Natl Acad. Sci. USA, 119(26): e2101388119. 2022.
- Yang, J. T., Preiser, A. L., Li, Z., Weise, S. E., and Sharkey, T. D.: Triose phosphate use limitation of
  photosynthesis: short-term and long-term effects, Planta, 243, 687-698, https://doi.org/10.1007/s00425015-2436-8, 2016.
- Yang, X., Tang, J., Mustard, J. F., Lee, J.-E., Rossini, M., Joiner, J., Munger, J. W., Kornfeld, A., and
  Richardson, A. D.: Solar-induced chlorophyll fluorescence that correlates with canopy photosynthesis
  on diurnal and seasonal scales in a temperate deciduous forest, Geophys. Res. Lett., 42, 2977-2987,
  https://doi.org/10.1002/2015gl063201, 2015.
- Yang, X., Chen, X., Ren, J., Yuan, W., Liu, L., Liu, J., Chen, D., Xiao, Y., Song, Q., Du, Y., Wu, S., Fan, L.,
  Dai, X., Wang, Y., and Su, Y.: A gridded dataset of a leaf-age-dependent leaf area index seasonality
  product over tropical and subtropical evergreen broadleaved forests, Earth Syst. Sci. Data, 15, 26012622, https://doi.org/10.5194/essd-15-2601-2023, 2023.
- Yang, X., Wu, J., Chen, X., Ciais, P., Maignan, F., Yuan, W., Piao, S., Yang, S., Gong, F., Su, Y., Dai, Y., Liu,
  L., Zhang, H., Bonal, D., Liu, H., Chen, G., Lu, H., Wu, S., Fan, L., Gentine, P., and Wright, S. J.: A
  comprehensive framework for seasonal controls of leaf abscission and productivity in evergreen
  broadleaved tropical and subtropical forests, Innovation (New York, N.Y.), 2, 100154,
  https://doi.org/10.1016/j.xinn.2021.100154, 2021.
- Yuan, W., Zheng, Y., Piao, S., Ciais, P., Lombardozzi, D., Wang, Y., Ryu, Y., Chen, G., Dong, W., Hu, Z., Jain,
  A. K., Jiang, C., Kato, E., Li, S., Lienert, S., Liu, S., Nabel, J. E. M. S., Qin, Z., Quine, T., Sitch, S.,
  Smith, W. K., Wang, F., Wu, C., Xiao, Z., and Yang, S.: Increased atmospheric vapor pressure deficit
  reduces global vegetation growth, Sci. Adv., 5, eaax1396, https://doi.org/10.1126/sciadv.aax1396, 2019.
- 916 Zhao, Y., Chen, X., Smallman, T. L., Flack-Prain, S., Milodowski, D. T., and Williams, M.: Characterizing
- 917 the error and bias of remotely sensed LAI Products: An Example for Tropical and Subtropical Evergreen
  918 Forests in South China, Remote Sens., 12, 3122, https://doi.org/10.3390/rs12193122, 2020.
- Zhou, H., Xu, M., Pan, H., and Yu, X.: Leaf-age effects on temperature responses of photosynthesis and
   respiration of an alpine oak, Quercus aquifolioides, in southwestern China, Tree Physiol., 35, 1236-1248,
   https://doi.org/10.1093/treephys/tpv101, 2015.

# 925Table 1. Data sources for mapping the $V_{c,max25}$ of young leaves across tropical and subtropical evergreen926broadleaved forests

Table

Data name (Abbr.)	Source	Usage	Spatial resolution	Temporal resolution	Temporal coverage
Temperature (T <sub>mean</sub> )	ERA5-Land	Calculate the $K_C, K_0, \Gamma^*,$ and $R_d$ for $A_n$	0.1°×0.1°	Monthly	2001.1-2018.12
Shortwave solar radiation (SW)	BESS	Calculate the $J_e$ for $A_n$	0.05°×0.05°	Monthly	2001.1-2018.12
Vapor pressure deficit (VPD)	ERA5-Land	Calculate the $C_i$ for $A_n$	0.1°×0.1°	Monthly	2001.1-2018.12
Sun induced chlorophyll fluorescence (RTSIF)	TROPOMI SIF	RTSIF- derived GPP	0.05°×0.05°	Monthly	2001.1-2018.12
Gross primary production retrieved from OCO-2 Solar induced chlorophyll fluorescence (GOSIF)	GOSIF	GOSIF- derived GPP	0.05°×0.05°	Monthly	2001.1-2018.12
Gross primary production from eddy covariance flux tower measurements (FLUXCOM)	FLUXCOM	FLUXCOM GPP	0.5°×0.5°	Monthly	2001.1-2013.12
Leaf-age-dependent leaf area index (Lad-LAI)	Yang et al., 2023	Dissolved V <sub>c,max25</sub> from GOSIF- derived GPP	0.25°×0.25°	Monthly	2001.1-2018.12

Figures



932 Figure 1. Tropical and subtropical evergreen broadleaved forests (TEFs) and *in situ* observation sites. The

- 933 TEFs is determined as those labeled as evergreen broadleaf forest (EBF) from the MODIS land cover maps
- at a  $0.05^{\circ}$  spatial resolution. The red dots are *in situ* observation sites of  $V_{c,max25}$ .
- 935

936



**Figure 2.** Procedures for mapping the  $V_{c,max25}$  of young leaves using a neighbor-based approach. 938



**Figure 3.** Validations of simulated seasonal  $V_{c,max25}$  for all canopy leaves and young leaves with *in situ* observations. The green lines and green dots are the seasonal  $V_{c,max25}$  of young leaf simulated from RTSIF derived GPP and *in situ* observations, respectively. The black dotted line and red dots are the  $V_{c,max25}$  of canopy-averaged leaves from the simulations and *in situ* observations, respectively. Simulated  $V_{c,max25}$  of denoted as the  $V_{c,max25}$  of young leaf simulated from RTSIF-derived GPP by using the new proposed method.



947 Figure 4. Comparisons of the  $V_{c,max25}$  of young leaves simulated from RTSIF-derived GPP against that 948 dissolved from GOSIF-derived GPP.

949



**Figure 5.** The root mean square error (RMSE) and correlation coefficient (R) between the  $V_{c,max25}$  of young leaves derived from RTSIF-derived GPP and that dissolved from GOSIF-derived GPP.





**Figure 6.** Comparison of sub-regions of the  $V_{c,max25}$  of young leaves (**a-c**) with those of climatic factors classified by the K-means clustering analysis (**d-f**) analyzed by Chen et al. (2021), and those of the Leaf-agedependent leaf area index product (L ad L AL) (**g** i) developed by Yang et al. (2023)

957 dependent leaf area index product (Lad-LAI) (g-i) developed by Yang et al. (2023).

958



959

**Figure 7.** Spatial maps of the correlation coefficient (R) between the monthly simulated  $V_{c,max25}$  and the Leaf-

961 age-dependent leaf area index seasonality product (Lad-LAI) developed by Yang et al. (2023).

962



965Figure 8. Seasonality of the simulated  $V_{c,max25}$  of young leaves in comparison with the Leaf-age-dependent966leaf area index product (Lad-LAI) developed by Yang et al. (2023).





**Figure 9.** Spatial maps of correlation coefficient (R) between the SIF-simulated monthly  $V_{c,max25}$  and vapor pressure deficit (VPD, **a-c**), air temperature (T<sub>mean</sub>, **d-f**), and downward shortwave solar radiation (SW, **g-i**).



**Figure 10.** Climatic drivers of spatial variations in average  $V_{c,max25}$  of young leaves across the TEFs (**a**) and three major tropical forests regions (**b-d**). Contributions ( $\emptyset$ ) of three climate factors to the multiple-yearaverage  $V_{c,max25}$  using the random forest (RF) algorithm. R<sup>2</sup> represents the coefficient of determination between simulated- and observed-  $V_{c,max25}$ . RMSE indicates the root mean standard error. Partial dependence plots (PDP) of the relationships between three climate drivers [T<sub>mean</sub> (K), SW (W m<sup>-2</sup>), VPD (hPa)] and  $V_{c,max25}$ . Relations for each pixel are displayed in black lines and relations on regional average are shown in red lines.

980



**Figure 11.** Scatter plots between the simulated  $V_{c,max25}$  of young leaves simulated using the 2×2 (0.25° resolution) and 4×4 (0.5° resolution) neighboring pixels in the above-mentioned ten clustered sub-regions.



**Figure 12.** Seasonality of the simulated  $V_{c,max25}$  of young leaf derived from RTSIF-, GOSIF- and FLUXCOM GPP in the ten clustered sub-regions.



**Figure 13.** Comparison of  $V_{c,max25}$  derived from three GPP products (i.e., GPP<sub>RTSIF</sub>, GPP<sub>GOSIF</sub>, GPP<sub>FLUXCOM</sub>)

and *in situ* observations. (a) Correlation coefficients (R); (b) Root mean square error (RMSE).