Environmental factors have a greater influence on photosynthetic capacity in C₄ plants than C₄ biochemical subtypes or growth forms

Yuzhen Fan^{1*}, Daniel W. A. Noble², Belinda E. Medlyn³, Russell K. Monson⁴, Rowan F. Sage⁵, Nicholas
G. Smith⁶, Elizabeth A. Ainsworth^{7,8,9}, Florian A. Busch¹⁰, Florence R. Danila¹, Maria Ermakova¹¹,
Patrick Friesen¹², Robert T. Furbank¹, Shu Han Gan¹³, Oula Ghannoum³, Daniel M. Griffith¹⁴, Lianhong
Gu¹⁵, Vinod Jacob³, Jürgen Knauer¹⁶, Andrew D. B. Leakey^{7,8,9}, Shuai Li^{7,8,17}, Danica L. Lombardozzi^{18,19},

- 8 Martha Ludwig²⁰, Varsha S. Pathare⁹, Murilo M. Peixoto^{21,22}, Karine Prado²³, Balasaheb V. Sonawane⁹,
- 9 Christopher J. Still²⁴, Susanne von Caemmerer¹, Russell Woodford¹¹, Danielle A. Way^{1,25,26}*
- 10
- 11 *Authors for correspondence
- 12 Yuzhen Fan (yuzhen.fan@anu.edu.au)
- 13 Danielle A. Way (danielle.way@anu.edu.au)
- 14
- 15 Number of Figures: 7 (+7 in Supplementary)
- 16 Number of Tables: 1 (+1 in Supplementary)
- 17 Number of References: 145
- 18 Number of Pages (main text): 17
- 19 Word count (incl. in-text citations): Total (7,489), Abstract (196), Introduction (1,819), Materials and
- 20 Methods (1,934), Results (1,189), Discussion (2,547)
- 21
- 22 Running title: A data synthesis of $C_4 A/C_i$ curves
- 23

24 Affiliation

- ¹Division of Plant Sciences, Research School of Biology, The Australian National University, Canberra,
 ACT 2601, Australia
- 27 ²Division of Ecology and Evolution, Research School of Biology, The Australian National University,
- 28 Canberra, ACT 2601, Australia
- ³Hawkesbury Institute for the Environment, Western Sydney University, Penrith, NSW 2751, Australia
- ⁴Department of Ecology and Evolutionary Biology, University of Colorado, Boulder, CO 80309, USA
- ⁵Department of Ecology and Evolutionary Biology, University of Toronto, Toronto, ON M5R3C5,
 Canada
- 33 ⁶Department of Biological Sciences, Texas Tech University, Lubbock, TX 79409, USA
- 34 ⁷Center for Advanced Bioenergy and Bioproducts Innovation, University of Illinois at Urbana-
- 35 Champaign, Urbana, IL 61801, USA
- 36 ⁸Carl R. Woese Institute for Genomic Biology, University of Illinois at Urbana-Champaign, Urbana, IL
- 37 61801, USA
- ⁹Departments of Plant Biology and Crop Sciences, University of Illinois Urbana Champaign, Urbana, IL
- **39** 61801, USA
- 40 ¹⁰School of Biosciences and Birmingham Institute of Forest Research, University of Birmingham,
- 41 Birmingham B15 2TT, UK
- 42 ¹¹School of Biological Sciences, Monash University, Melbourne, VIC 3800, Australia
- 43 ¹²BioChambers, Winnipeg, MB R2W 3A8, Canada
- ¹³Department of Ecology, Evolution & Environmental Biology, Columbia University, New York, NY
 10027, USA
- 46 ¹⁴Forest Ecosystems and Society, Oregon State University, Corvallis, OR 97331, USA
- 47 ¹⁵Environmental Sciences Division and Climate Change Science Institute, Oak Ridge National
- 48 Laboratory, Oak Ridge, TN 37831, USA
- ¹⁶School of Life Sciences, Faculty of Science, University of Technology Sydney, Ultimo, NSW 2007,
 Australia
- 51 ¹⁷Guangdong Provincial Key Laboratory of Applied Botany, South China Botanical Garden, Chinese
- 52 Academy of Sciences, Guangzhou, Guangdong 510650, China
- ¹⁸Department of Ecosystem Science & Sustainability, Colorado State University, Fort Collins, CO 80523,
- 54 USA

- ¹⁹Climate and Global Dynamics Laboratory, National Center for Atmospheric Research, Boulder, CO
 80305, USA
- ²⁰School of Molecular Sciences, University of Western Australia, Perth, WA 6009, Australia
- 58 ²¹School of Biological Sciences, Washington State University, Pullman, WA 99164, USA
- 59 ²²Universidade Federal de Goiás, Programa de Pós-graduação em Biodiversidade Vegetal, Instituto de
- 60 Ciências Biológicas, Goiânia, GO 74690-900, Brazil.
- 61 ²³Plant Resilience Institute, Department of Biochemistry and Molecular Biology, Michigan State
- 62 University, East Lansing, MI 48823, USA
- 63 ²⁴Department of Forest Ecosystems and Society, Oregon State University, Corvallis, OR 97331, USA
- 64 ²⁵Department of Biology, The University of Western Ontario, London, ON N6A 3K7, Canada
- 65 ²⁶Nicholas School of the Environment, Duke University, Durham, NC 27708, USA
- 66

67 ORCID

- 68 Yuzhen Fan: 0000-0003-1857-9244
- 69 Daniel W. A. Noble: 0000-0001-9460-8743
- 70 Belinda E. Medlyn: 0000-0001-5728-9827
- 71 Russell K. Monson: 0000-0002-7671-4371
- 72 Rowan F. Sage: 0000-0001-6183-9246
- 73 Nicholas G. Smith: 0000-0001-7048-4387
- 74 Elizabeth A. Ainsworth: 0000-0002-3199-8999
- 75 Florian A. Busch: 0000-0001-6912-0156
- 76 Florence R. Danila: 0000-0002-7352-3852
- 77 Maria Ermakova: 0000-0001-8466-4186
- 78 Patrick Friesen: 0000-0002-1027-4502
- 79 Robert T. Furbank: 0000-0001-8700-6613
- 80 Shu Han Gan: 0000-0002-8699-3669
- 81 **Oula Ghannoum**: 0000-0002-1341-0741
- 82 Daniel M. Griffith: 0000-0001-7463-4004
- 83 Lianhong Gu: 0000-0001-5756-8738
- 84 Vinod Jacob: 0000-0002-6627-5993

- 85 Jürgen Knauer: 0000-0002-4947-7067
- 86 Andrew D. B. Leakey: 0000-0001-6251-024X
- 87 Shuai Li: 0000-0003-2545-7763
- 88 Danica L. Lombardozzi: 0000-0003-3557-7929
- 89 Martha Ludwig: 0000-0002-0324-7602
- 90 Varsha S. Pathare: 0000-0001-6220-7531
- 91 Murilo M. Peixoto: 0000-0002-6242-6870
- 92 Karine Prado: 0000-0002-0502-0380
- 93 Balasaheb V. Sonawane: 0000-0001-6539-5179
- 94 Christopher J. Still: 0000-0002-8295-4494
- 95 Susanne von Caemmerer: 0000-0002-8366-2071
- 96 Russell Woodford: 0000-0002-6766-2274
- 97 Danielle A. Way: 0000-0003-4801-5319

99 Abstract

- Our understanding of how photosynthesis varies among C₄ species and across different growth
 and measurement conditions remains limited.
- We collated 1,696 CO₂ response curves of net CO₂ assimilation rate $(A/C_i \text{ curves})$ from C₄ species grown and measured at various environmental conditions and used these data to estimate the apparent maximum carboxylation activity of phosphoenolpyruvate carboxylase (V_{pmaxA}) and CO₂saturated net photosynthetic rate (A_{max}) , two key parameters describing C₄ photosynthetic capacity. We examined how V_{pmaxA} and A_{max} vary with species-specific traits, growth and measurement conditions.
- We show that V_{pmaxA} and A_{max} do not differ between C₄ biochemical subtypes or growth forms, and highlight that growth temperature and measurement conditions are major factors determining photosynthetic capacity. We found no evidence that common C₄ model species (e.g., maize, sorghum and *Setaria viridis*) differ in photosynthetic capacity from other C₄ species when grown in controlled environments. However, C₄ model species showed up to twice the photosynthetic capacity of other C₄ species when grown in the field.
- Our multivariate model accounts for 47-51% of the variation reported in V_{pmaxA} and A_{max}, and we
 argue that environmental conditions have a greater influence on C₄ photosynthetic capacity than
 inherent biochemical subtypes or growth forms.

117 Keywords

118 A/C_i curve, A_{max} , C₄ photosynthesis, C₄ subtype, environmental response, photosynthesis modelling, V_{pmax}

120 Introduction

121 C_4 photosynthesis evolved at least 66 times in the last 35 Myr in response to inefficiencies in C_3 122 photosynthesis under hot, arid and low CO₂ environments that promote photorespiration (Sage, 2016). In 123 C₄ plants, a CO₂-concentrating mechanism increases CO₂ concentrations at the site of Rubisco (ribulose-124 1,5-bisphosphate carboxylase/oxygenase) in bundle sheath cells, resulting in an increase in its carboxylase 125 activity, a reduction in photorespiration, and an increase in photosynthetic efficiency (Hatch, 1987). 126 Compared to their C_3 counterparts, C_4 plants require one-third of the amount of Rubisco to achieve the 127 same or higher rates of net CO₂ uptake, leading to greater nitrogen-use efficiency (Ghannoum et al., 2011). C₄ plants also have higher water-use efficiency because they can maintain a lower stomatal conductance 128 129 relative to the rate of net photosynthetic CO₂ assimilation (A), which supports greater biomass production 130 when water is limited (Evans & von Caemmerer, 1996; Leegood, 2002; Taylor et al., 2010; Ghannoum et 131 al., 2011). Thus, C₄ plants tend to outperform C₃ plants in hot and dry environments, lending them a 132 competitive advantage in subtropical and semiarid ecosystems. As a result, C₄ plants occupy ~20% of the 133 land surface and contribute up to 23% of global gross primary productivity although they comprise <5% 134 of terrestrial plant species (Still et al., 2003; Luo et al., 2024).

135 Currently, C₄ global primary productivity is typically estimated in Earth System Models using a 136 simplified mechanistic model that uses parameters reflecting photosynthetic capacity (Still et al., 2019; 137 Griffith et al., 2020). The model and its parameter values are largely derived from data collected on a 138 single species, Zea mays (maize) (Collatz et al., 1992). Consequently, the model does not capture 139 substantial variation across C_4 species in the biochemistry underlying CO_2 assimilation (Hatch, 1987), or 140 in their growth forms (Poorter et al., 2009; Liu et al., 2019). The responses of photosynthetic capacity to 141 environmental parameters, such as temperature and irradiance, are also taken from the same study on 142 maize (Collatz et al., 1992). Over the last 30 years, many studies have reported how C₄ photosynthetic 143 parameters vary in response to environmental drivers and described genetic differences within and 144 between species. However, these findings are scattered across publications, except for an analysis by 145 Pignon & Long (2020), and have not been subjected to data synthesis in a manner that could advance the 146 representation of this important plant functional type in Earth System Models.

The most common way to estimate leaf-level photosynthetic capacity is to measure CO₂ response curves of net CO₂ assimilation rate (i.e. A/C_i curves), and assess both the initial slope of the A/C_i curve (a parameter closely related to the apparent maximum activity of phosphoenolpyruvate (PEP) carboxylase, V_{pmaxA}) and the horizontal asymptote of the A/C_i curve (i.e. the rate of CO₂-saturated net photosynthesis, 151 termed A_{max}). The initial slope of a C₄ A/C_i curve is generally insensitive to changes in leaf measurement 152 temperatures (T_{leaf}) for plants grown at moderate temperatures (Long & Woolhouse, 1978; Laisk & 153 Edwards, 1997; Sage, 2002), although it can be reduced in plants grown under chilling conditions, possibly 154 reflecting an increase in the activation energy of PEP carboxylase (PEPc) at low temperatures (Pittermann 155 & Sage, 2001; Kubien & Sage, 2004). In contrast, A_{max} increases with increasing T_{leaf} in C₄ plants grown 156 under moderate temperatures (Sage & Kubien, 2007). As with T_{leaf} , the initial slope of a C₄ A/C_i curve 157 remains largely stable when measured at moderate versus high photosynthetic photon flux density (PPFD) (e.g. 500 vs. 1,500 μ mol photons m⁻² s⁻¹), suggesting that irradiance does not directly change PEPc 158 159 kinetics, and PEP regeneration may not be the limiting step of photosynthesis at low C_i (Pengelly *et al.*, 160 2010). However, A_{max} is sensitive to changes in irradiance and increases when measured under high *PPFD* 161 (Pengelly *et al.*, 2010). This irradiance response of A_{max} may result from the alleviation of chloroplastic 162 electron transport limitation under high PPFD (Ermakova et al., 2019, 2023), thereby enhancing energy 163 availability for PEP or RuBP regeneration, leading to higher overall net photosynthesis (von Caemmerer 164 & Furbank, 2016). Taken together, the responses of photosynthetic capacity to measurement conditions 165 are dynamic, with the initial slope of the A/C_i curve tending to be less affected by T_{leaf} and PPFD than 166 A_{\max} .

167 C₄ photosynthetic capacity can also be altered by growth temperatures. There is evidence that 168 photosynthetic capacity (particularly A_{max}) is reduced in C₄ plants grown at warm to high temperatures 169 (i.e. 30 to 43°C), as a result of lower Rubisco content and/or activity, NADP-malic enzyme activity, 170 cytochrome f content and carbonic anhydrase activity (Pearcy, 1977; Ward, 1987; Dwyer et al., 2007). In 171 contrast, the activity of PEPc (and potentially V_{pmaxA}) may not be affected by high growth temperatures, 172 possibly due to its high thermal stability (Chen et al., 1994; Chinthapalli et al., 2002; Boyd et al., 2015). 173 At low growth temperatures, many chilling-tolerant, cool-adapted C₄ plants show no change in 174 photosynthetic rates or Rubisco concentrations compared to control plants (Pittermann & Sage, 2001; 175 Naidu et al., 2003; Cavaco et al., 2003; Kubien & Sage, 2004), though the content of other photosynthetic 176 enzymes [e.g. pyruvate, phosphate dikinase (PPDK)], can increase (Wang et al., 2008). These findings 177 highlight that while some C_4 plants can adjust their photosynthetic machinery in response to growth 178 temperature, both high and low temperatures can impose biochemical constraints that limit photosynthetic 179 capacity via acclimation and thermal stress.

180 Given that C₄ photosynthesis is thought to be an evolutionary response to high photorespiratory
 181 loads (Sage, 2004), there has been considerable interest in the response of C₄ photosynthesis to variation

182 in CO₂ concentrations. When C₄ plants are grown in controlled environments, A_{max} can increase under 183 sub-ambient CO₂ conditions, reflecting an up-regulation of Rubisco capacity to enhance carbon capture 184 (Ripley et al., 2013; Pinto et al., 2014; Cunniff et al., 2017), but remains largely unchanged under the 185 elevated CO₂ levels projected for the coming century (Leakey, 2009; Heckman et al., 2024). In contrast, 186 while the capacity of PEPc in controlled environment-grown plants can also be increased at glacial CO₂ 187 concentrations, it can be reduced at elevated CO₂ concentrations (Wong, 1979; Ghannoum et al., 2000; 188 Pinto et al., 2014). However, when C₄ plants are grown in the field, changes in growth CO₂ concentrations 189 had little effect on photosynthesis or the activity/capacity of key photosynthetic enzymes including PEPc 190 (Leakey et al., 2006; Leakey, 2009; Markelz et al., 2011). Both sub-ambient and elevated CO₂ 191 concentrations influence stomatal conductance, leading to changes in plant water relations and interactions 192 with drought stress that can be complex in nature and create opportunities for crop improvement 193 (Ghannoum et al., 2000; Markelz et al., 2011; Leakey et al., 2019). Although our understanding of how 194 C₄ plants respond to changes in CO₂ concentration and drought, along with their underlying physiological 195 mechanisms, has advanced in recent decades [e.g. (Tissue et al., 1995; Wang et al., 2020; Ding et al., 196 2022)], this knowledge has yet to be fully integrated into ecosystem models (Cowling et al., 2007; Still et 197 al., 2019).

198 There is some evidence suggesting that the response of C₄ photosynthetic capacity to growth 199 environment depends on the biochemistry, life history and plant functional type of the species (e.g. growth 200 form: monocot versus eudicot) (Liu et al., 2019). Based on the decarboxylases involved, the C4 201 photosynthetic pathway can be categorised into three classical biochemical subtypes: NADP-dependent 202 malic enzyme (NADP-ME subtype), NAD-dependent malic enzyme (NAD-ME subtype) and PEP-203 carboxykinase (PCK subtype) (Hatch, 1987; von Caemmerer & Furbank, 2016). Considerable variation 204 in biochemical components (e.g. enzyme abundance) and photosynthetic capacity has been found in C4 205 plants of different growth forms within and between C₄ biochemical subtypes. For example, the maximum 206 carboxylation activity of PEPc and Rubisco, A_{max} and leakiness (i.e. the CO₂ fraction that leaks out from 207 bundle sheath cells) responded to a short-term increase in T_{leaf} differently in eight C₄ grasses representing 208 the three C₄ biochemical subtypes (Sonawane *et al.*, 2017). In addition, C₄ annual species have been shown to exhibit a greater sensitivity of the chloroplastic electron transport capacity to T_{leaf} than C₄ perennial 209 210 species (Smith & Dukes, 2017). These findings suggest that differences in C₄ biochemistry and growth 211 form could play a role in determining photosynthetic responses to environmental conditions.

212 To understand how C₄ photosynthetic capacity responds to environmental cues—and whether 213 these responses differ systematically among biochemical subtypes and growth forms-data are needed 214 from diverse species grown and measured under contrasting environmental conditions. Unfortunately, 215 while numerous studies have evaluated photosynthetic capacity of C4 plants under various growth and 216 measurement conditions (reference herein), these data are largely siloed in their separate publications, 217 making it challenging to identify broad patterns in photosynthetic traits across C₄ species. In this study, 218 we used published gas-exchange data to explore the drivers of variation in C₄ photosynthetic capacity. 219 Here, A/C_i curves measured from diverse C₄ species across studies were collated to estimate V_{pmaxA} and Amax with a widely used C4 mechanistic model (von Caemmerer & Furbank, 1999; von Caemmerer, 2000, 220 221 2021). We assessed how V_{pmaxA} and A_{max} are affected by species-specific traits (biochemical subtype and 222 growth form) and environmental parameters (growth and measurement conditions). For growth 223 conditions, we considered growth temperature and growth CO₂ concentration. We also considered growth 224 location (whether plants were grown in outdoor fields or in indoor controlled environments; see Materials 225 and Methods for definitions). Compared to outdoor plants, indoor plants may experience limited growth 226 space and different environments (e.g. fewer pests, relatively steady temperature and light conditions), 227 which can affect overall morphology and physiology (Poorter et al., 2016). For measurement conditions, we were interested in how C₄ photosynthetic capacity varies with measurement T_{leaf} and PPFD. An A/C_i 228 229 curve should be measured under light-saturating conditions, such that PPFD is often not considered a key 230 factor in photosynthetic capacity. However, the PPFD required for light saturation of net photosynthesis 231 can vary with growth conditions, and A/C_i curves can also be measured under sub-saturating PPFD to 232 address specific research questions [e.g. (Sonawane et al., 2018)].

- 233 Using these data, we test the following hypotheses:
- 234 (1) V_{pmaxA} and A_{max} are positively correlated;
- 235 (2) C₄ biochemical subtype and growth form alter V_{pmaxA} and A_{max} in a systematic way;
- (3) Indoor plants show a higher photosynthetic capacity than their outdoor counterparts due tomore optimal growth conditions;
- 238 (4) Cool-grown plants have a higher photosynthetic capacity relative to their warm-grown 239 counterparts when measured at a common T_{leaf} ;
- 240 (5) A_{max} increases with increasing T_{leaf} and measurement *PPFD*, but V_{pmaxA} does not;
- 241 (6) Elevated growth CO_2 concentrations have no effect on photosynthetic capacity in C_4 species.

Lastly, we assess whether the photosynthetic capacity of popular C₄ model species—maize, 243 Sorghum bicolor (sorghum), and Setaria viridis—the three most abundant species in our dataset, is similar 244 to that of other C_4 species to assess whether data from these three species can be used to represent the 245 broad range of undomesticated C₄ species found in nature. We hypothesise that:

246 (7) These C₄ model species show higher V_{pmaxA} and A_{max} than other C₄ species in the analysis.

Materials and Methods 247

Data acquisition 248

249 This work was conceived by members of the C₄ Working Group supported by the U.S Geological Survey's 250 Powell Center, with data collaboratively sourced within the group and from researchers closely associated 251 with its members. C₄ A/C_i data were compiled from 52 studies published between 2001 and 2024 (Pittermann & Sage, 2001; Anderson et al., 2001; Cousins & Bloom, 2003; Leakey et al., 2006; Kim et 252 253 al., 2006; Ripley et al., 2007, 2008, 2010, 2013; Nippert et al., 2007; Kakani et al., 2008a,b; Osborne et 254 al., 2008; Horst et al., 2008; Soares et al., 2008; Cunniff et al., 2008; Carmo-Silva et al., 2008; Mantlana 255 et al., 2008; Pengelly et al., 2010; de Souza, 2011; Soares-Cordeiro et al., 2011; Pinto et al., 2011, 2014; 256 Arena et al., 2011; Wang et al., 2012; Bloom et al., 2012; Sun et al., 2012; Vogan & Sage, 2012; Fay et 257 al., 2012; Feng et al., 2012; Chen et al., 2013; Sage et al., 2013; Oakley et al., 2014; Xu et al., 2014; 258 Bissinger et al., 2014; Friesen et al., 2014; Sharwood et al., 2014; Ge et al., 2014; Głowacka et al., 2015; 259 Sonawane et al., 2017, 2018; Peixoto & Sage, 2017; Smith & Dukes, 2018; Li et al., 2019, 2021, 2022; 260 Khoshravesh et al., 2020; Pathare et al., 2020; Danila et al., 2021; Peixoto et al., 2021; Liu et al., 2023; 261 Gan & Sage, 2024; Ermakova et al., 2024) and five unpublished datasets with permission to use. Together, 1,696 C₄ A/C_i curves for 74 species of 12 plant families from 57 unique studies were included in the 262 263 analysis. Raw data files (i.e. direct output files from the measuring instrument) were requested from 264 authors where available. In cases where the raw data files were not available, curves were digitised from 265 published figures as per Pignon & Long (2020). The compiled dataset is provided in Supplementary 266 Dataset S1.

$A/C_{\rm i}$ curve analysis 267

268 To fit the curves and estimate parameters, the 1,696 individual A/C_i curves were consolidated into 543 269 groups. Replicate measurements of the same species and genotype that were measured within five days in a study were considered as one experimental group. This approach allowed us to minimise software crashes due to a lack of model convergence and estimate sampling variance; this grouping approach has been used in other studies (e.g. Wu *et al.*, 2024). The A/C_i data at $C_i < 100 \,\mu$ mol mol⁻¹ (i.e. the initial slope of the A/C_i curve) were used to estimate V_{pmaxA} , according to von Caemmerer (2000):

274

275
$$A = V_{\rm p} - R_{\rm m} = \frac{c_{\rm i} v_{\rm pmaxA}}{c_{\rm i} + \kappa_{\rm p}} - R_{\rm m}$$
Equation (1)

276

where A is the net CO₂ assimilation rate (μ mol CO₂ m⁻² s⁻¹), V_p is the rate of PEP carboxylation (μ mol 277 $CO_2 \text{ m}^{-2} \text{ s}^{-1}$), C_i is the CO_2 concentration in the intercellular space (µmol mol⁻¹ CO₂), K_p is the Michaelis-278 279 Menten constant for CO₂ (µbar; converted to concentrations using a solubility for CO₂ of 0.0334 mol bar-280 ¹ and atmospheric pressure at the measurement site; see von Caemmerer *et al.* (1994)), and R_m is daytime mitochondrial respiration in mesophyll cells (µmol CO2 m⁻² s⁻¹). Under low C_i, A is linearly correlated 281 with the maximum PEPc activity in mesophyll cells, given that the leakage of CO₂ from bundle sheath 282 283 cells is low and is generally ignored (von Caemmerer, 2021). We also assumed that PEP substrate 284 concentrations were saturating under the measurement condition, and A was not limited by PEP regeneration, as required by the C4 model (von Caemmerer, 2000, 2021). Kp was adjusted to account for 285 286 variation in T_{leaf} according to Boyd *et al.* (2015):

287

288
$$K_{\rm p} = K_{\rm p25} \times e^{E_{\rm a}(T_{\rm leaf} - 25)/(298.15R(T_{\rm leaf} + 273.15))}$$
 Equation (2)

289

where K_{p25} is K_p measured at 25°C (assumed to be 80 µbar; Bauwe, 1986; DiMario & Cousins, 2019), E_a is the activation energy of K_p (36.3 kJ mol⁻¹; Boyd *et al.*, 2015), and *R* is the molar gas constant (0.008314 kJ K⁻¹ mol⁻¹). R_m was taken as half the daytime leaf mitochondrial respiration rate (R_{day}) (µmol CO₂ m⁻² s⁻¹) (von Caemmerer, 2000). Given that there are few R_{day} measurements in C₄ plants, R_{day} was assumed to equal leaf dark respiration such that $R_{day} = 1.2 \mu mol CO_2 m^{-2} s^{-1}$ at 25°C (termed as R_{day25}), according to a data synthesis of dark respiration in 39 C₄ species (Fan *et al.*, 2022). R_{day} was adjusted for variation in T_{leaf} using a Q_{10} equation (Atkin & Tjoelker, 2003):

297

298
$$R_{\text{day}} = R_{\text{day25}} \times Q_{10}^{\frac{T_{\text{leaf}} - 25}{10}}$$
 Equation (3)

300 where Q_{10} is 2 (Huntingford *et al.*, 2013).

Although mechanistic models for enzyme- and light-limited C₄ photosynthesis are available, C₄ photosynthesis could be co-limited by Rubisco activity, RuBP regeneration and/or PEP regeneration at high C_i , with all three processes requiring energy from light (von Caemmerer & Furbank, 1999). Thus, as no information was provided on the limitation status of photosynthesis at high C_i by individual studies, we estimated A_{max} using the entire A/C_i curve. The A_{max} was estimated by the horizontal asymptote of a four-parameter non-rectangular hyperbolic function (Leakey *et al.*, 2006; Li *et al.*, 2022):

307

308
$$\theta (A + R_{day})^2 - (\alpha C_i + A_{max})(A + R_{day}) + \alpha C_i A_{max} = 0$$
 Equation (4)

309

310 where θ is the curvature factor of an A/C_i curve (unitless), which was assumed to be 0.7 for C₄ plants 311 (Sonawane *et al.*, 2018; von Caemmerer, 2021), and α is the initial slope of an A/C_i curve (µmol CO₂ m⁻²) 312 s^{-1}). We did not include mesophyll conductance as a term in Eq (4) due to a lack of such data in examined 313 studies. However, by assuming $\theta = 0.7$, we considered a potential drop in the C_i between the intercellular 314 space and the site of Rubisco carboxylation (i.e. bundle sheath cells) (Evans, 1989). All curve fitting was 315 performed using the non-linear least square (nls) function and sampling variances of every fit were 316 extracted using the vcov function in R (version 4.3.2, R Core Team, 2018). The number of individual 317 curves per group (n_{group} , ranging from 1 to 9) was also reported. Sampling variance of parameters (i.e., 318 squared standard error for parameters) was subsequently considered in the statistical models (Nakagawa 319 & Santos, 2012; Noble et al., 2017, 2022).

320 Data classification and summary

321 Among the 543 groups, eight groups contained fewer than two data points at low C_i (i.e. < 100 μ mol mol⁻ ¹), while 131 groups contained fewer than two data points at high C_i (i.e. > 500 μ mol mol⁻¹). As this lack 322 323 of data is likely to influence the accuracy of model fitting using these portions of the A/C_i curve, the 324 corresponding estimated V_{pmaxA} and A_{max} values of these groups were discarded. Given that measurements 325 within a single group were made under the same measurement temperature and irradiance, we provided averaged values of T_{leaf} and *PPFD* per group. We further assessed the estimated parameters of each group: 326 if the curve violated the assumption that A is limited by V_{pmaxA} at low C_i (as indicated by V_{pmaxA} -limited 327 328 photosynthesis being higher than A_{max} -limited photosynthesis at low C_i), this group of A/C_i curves was 329 discarded as per Pignon & Long (2020). This assessment led to 108 additional groups being removed from

the final dataset (Supplementary Dataset S2), many of which included plants subjected to severe abioticstresses such as chilling, drought, and low nutrients.

332 The remaining measurements were classified according to their C4 biochemical subtype (NADP-333 ME, NAD-ME or PCK subtype), growth form (monocot or eudicot) and growth location (indoor or 334 outdoor). Indoor plants refer to those grown in pots within controlled environment chambers or 335 glasshouses. In contrast, outdoor plants are grown directly in the soil or in common gardens without pot 336 restrictions. Experiments conducted at Free-Air Carbon Dioxide Enrichment (FACE) and Lysimeter CO2 337 Gradient (LYCOG) facilities are classified as outdoor-grown, as plants in these settings are planted 338 directly in soil. The growth CO₂ concentration was noted, where the ambient CO₂ level was assumed to 339 be 400 ppm if not given by the study, as 400 ppm represents the average atmospheric CO₂ concentration 340 from 2001 to 2024 (Friedlingstein et al., 2025), which covers the period of publications used in this study, 341 though the experiments themselves may have been conducted one or two years earlier. For indoor-grown 342 plants, mean maximum (T_{max}) and minimum (T_{min}) growth temperatures were the set daytime and 343 nighttime temperatures, respectively. We used the reported measurement T_{leaf} in our analyses, rather than normalising to a rate at a set temperature of 25°C, as we did not want to assume that the V_{pmax} and A_{max} of 344 345 all species and growth environments will have the same sensitivity to short-term changes in T_{leaf} . For 346 outdoor-grown plants, mean T_{max} and T_{min} were the mean maximum and minimum temperatures observed 347 across the experimental periods at the study sites, respectively. Growth PPFD was unavailable in most 348 outdoor studies and was therefore excluded from the analysis. We were unable to consider either 349 fertilisation treatment or water status as potential categories for analysis, as low nutrient treatment and/or 350 water-stressed conditions constituted <5% of the data (Fig. 1D & E). A summary of measurements based 351 on plant traits and growth treatments is given in Figure 1.

Across the dataset, maize, sorghum and *Setaria viridis* were the three most commonly measured species, accounting for 18%, 6% and 4% of the total measurements, respectively. Indeed, maize contributed 43% of the outdoor data and played an important role in the growth location category (Fig. S1). This species skewness is later accounted for in the statistical models (see below).

356 Statistical analysis

To determine how different plant traits and experimental conditions altered V_{pmaxA} and A_{max} , we ran two multivariate linear mixed-effects models with V_{pmaxA} or A_{max} as the dependent variable in each model. These models accounted for the sampling variance of each parameter, with parameters having greater 360 precision being weighted more heavily to overall means. We included three main fixed effect categories 361 in our models: (1) species-specific traits, including C₄ subtype and growth form; (2) environmental growth 362 conditions, such as growth location, temperature and CO₂ concentration; and (3) measurement conditions, 363 such as T_{leaf} and *PPFD*. Given that mean T_{max} and T_{min} were strongly positively correlated with each other 364 (Fig. S2), we used mean T_{max} in the model. A list of fixed effects is given in Table 1.

In addition, we included six a priori interaction terms examining the interactive effects of species-365 366 specific traits, temperatures and irradiance: C₄ subtype \times mean T_{max} , C₄ subtype \times T_{leaf} , C₄ subtype \times *PPFD*, growth form \times mean T_{max} , growth form \times T_{leaf} , and growth form \times *PPFD*. These terms were 367 368 designed to test the question of whether the response of C4 photosynthetic capacity traits to short- and long-term temperatures and measurement irradiance differ among C₄ subtypes or growth forms. We 369 370 confirmed that our V_{pmaxA} and A_{max} data per C₄ subtypes and growth form were distributed across the entire 371 range of mean T_{max} , T_{leaf} and *PPFD* (Fig. S3), such that any significant interaction terms would not be biased due to separated data distributions. Data distribution with respect to mean T_{max} and T_{leaf} is shown 372 in Figure S4. Random effects included in the models were the different studies (indicated by publications), 373 374 grouping of the curves (see above section titled $C_4 A/C_i$ analysis), and species. Given that C_4 model species 375 (maize, sorghum and Setaria viridis) made up a significant proportion of the data, we also conducted a 376 case study examining to what extent the model results obtained from data of all species match with those of the C₄ model species and visualise our results with consideration of C₄ model species (see Results). 377

The multivariate linear mixed-effects models with restricted maximum likelihood estimation were run using the package metafor version 4.6.0 (Viechtbauer, 2010) and plots were made using the package ggplot2 version 3.5.1 (Wickham, 2016) and orchard version 2.0 (Nakagawa *et al.*, 2021, 2023) on R program version 4.4.1 (R Core Team, 2024). Estimated effects were considered significant if P < 0.05 in the models.

383 Results

384 Positive correlations between V_{pmaxA} and A_{max}

In general, V_{pmaxA} and A_{max} were correlated with each other in a logarithmic manner ($R^2 = 0.81$; Fig. 2A). We found that for leaves measured at a moderate T_{leaf} (i.e. < 35°C), changes in V_{pmaxA} were tightly coupled with changes in A_{max} in a nearly linear fashion (Fig. 2A). When leaves were measured at a hotter T_{leaf} (i.e. > 35°C), the coupling between V_{pmaxA} and A_{max} broke down, and an increase in V_{pmaxA} was 389 accompanied by a lesser increase in A_{max} . This result was supported by significantly different slopes (P <

390 0.001; standardised major axis analysis) in the linear correlations between V_{pmaxA} and A_{max} for leaves

391 measured at 25–30°C (slope = 0.49; Fig. 2B), 30–35°C (slope = 0.39; Fig. 2C), and 35–40°C (slope =

392 0.27; Fig. 2D). Together, these findings highlight that at high A_{max} (i.e. high C_i), photosynthesis was less

- 393 dependent on PEP carboxylation, compared to other high-temperature sensitive biochemical processes
- 394 (e.g. Rubisco carboxylation), or photosynthesis was constrained by increased limitations in light and
- bundle sheath [CO₂] at high A_{max} (e.g. affecting RuBP regeneration).

396 Factors influencing V_{pmaxA} and A_{max} are mostly environmental

Multivariate linear mixed-effects models highlighted several factors that significantly affected both V_{pmaxA} and A_{max} (Table 1). C₄ subtype identity and growth form had little influence on values of either V_{pmaxA} or A_{max} (Fig. 3A-D). Note that the number of studies examining eudicots or PCK-type plants was noticeably lower compared to studies focusing on monocots or NADP-ME plants, respectively. This highlights an urgent need to further investigate photosynthetic characteristics in eudicots and PCK-type plants. There is no evidence that V_{pmaxA} and A_{max} vary among families or subfamilies (P = 0.73 and 0.88 for V_{pmaxA} and A_{max} , respectively) or within each biochemical subtype (Fig. S5).

404 We found that both V_{pmaxA} and A_{max} were higher when plants were grown indoors than in the field $(P = 0.005 \text{ and } 0.018 \text{ for } V_{\text{pmaxA}} \text{ and } A_{\text{max}}, \text{ respectively; Fig. 3E&F})$, where averaged mean T_{max} values 405 406 were 28.2 and 25.2°C, respectively, for indoor and field plants. Specifically, the models estimated that 407 V_{pmaxA} of indoor plants was 35% higher than their outdoor counterparts (55.3 \pm 1.7 versus 40.0 \pm 2.1 408 μ mol m⁻² s⁻¹), while A_{max} of indoor-grown plants was 22% higher than those grown outdoors (33.9 ± 0.9 409 versus $26.8 \pm 1.1 \,\mu$ mol m⁻² s⁻¹). It is worth noting that C₄ model species (i.e. maize and sorghum) appear 410 to be at the high-end of the V_{pmaxA} and A_{max} spectrum, particularly for outdoor-grown plants (Fig 3E&F). 411 We thus further examined whether there was an interaction between species group and growth location, and found it to be significant (P < 0.001). The mean values of V_{pmaxA} and A_{max} were significantly higher 412 413 in outdoor-grown C₄ model species compared to non-model species grown outside (P < 0.001; Fig. 4). 414 However, no significant differences were observed between model and non-model species grown indoors. 415 Our results suggest that the lower V_{pmaxA} and A_{max} values of C₄ non-model species grown outdoors 416 compared to their indoor counterparts could have been attributed to the effect of growth location on 417 photosynthetic capacity.

418 We also found that V_{pmaxA} and A_{max} decreased significantly with increasing mean T_{max} (P = 0.001; 419 Fig. 5A&B), with this pattern being stronger for a subset of data measured at a T_{leaf} between 25 and 30°C 420 to account for variation in T_{leaf} in the analysis (Fig. S6). No effect of growth CO₂ concentrations was found 421 on either V_{pmaxA} or A_{max} (Fig. 5C&D). In contrast to the negative response to mean T_{max} , V_{pmaxA} and A_{max} 422 increased significantly with increasing T_{leaf} (P < 0.0001; Fig. 6A&B) and PPFD (P < 0.0001; Fig. 6C&D). 423 These results highlight the different responses of V_{pmaxA} and A_{max} to short-term changes in T_{leaf} and long-424 term acclimation to mean T_{max} (see below). Nonetheless, neither the effect of mean T_{max} nor T_{leaf} on V_{pmaxA} 425 or Amax was explained by the growth location (Fig. 5A&B; Fig. 6A&B). We further examined whether 426 different C₄ subtypes or growth forms varied in their response to mean T_{max} , T_{leaf} and PPFD by considering 427 interaction effects. None of the interaction terms was significant (Table 1), suggesting responses of V_{pmaxA} 428 and A_{max} to growth and measuring temperatures and irradiance are independent of species-specific traits 429 (such as growth form). Overall, the fixed effects considered in our models accounted for 51 and 47% of 430 the variation in V_{pmaxA} and A_{max} , respectively (Table 1).

431 Interactive effects of T_{leaf} and mean T_{max} on photosynthetic capacity

432 The different responses of V_{pmaxA} and A_{max} to changes in long-term mean T_{max} and short-term T_{leaf} pointed 433 towards a possible interaction between these two temperature factors (Fig. 5&6). We thus explored the 434 mean $T_{\rm max} \times T_{\rm leaf}$ interaction and the result showed that the $V_{\rm pmaxA}$ was not affected by the interaction 435 between growth and measurement temperatures (P = 0.724; Table 1), although the interaction term was 436 significant for A_{max} (P = 0.033). We further illustrated the complex responses of V_{pmaxA} and A_{max} to mean T_{max} and T_{leaf} using contour plots (Fig. 7). The contour plots show that at any given mean T_{max} , the 437 438 sensitivity of V_{pmaxA} to changes in T_{leaf} was not influenced by acclimation to different mean T_{max} (Fig. 7A). This pattern also means that V_{pmaxA} measured at the predominant leaf temperatures of warm-grown plants 439 440 is higher compared to V_{pmaxA} measured at the predominant leaf temperatures of plants grown at lower 441 temperatures. A_{max} also increased with increasing T_{leaf} , but the increase in A_{max} per 1°C increase in T_{leaf} 442 was higher in plants grown in cooler environments (i.e. mean $T_{\text{max}} < 25^{\circ}$ C) compared to warmer 443 environments (Fig. 7B). This suggests that A_{max} was more sensitive to changes in T_{leaf} in plants acclimated 444 to lower mean T_{max} . This pattern highlights that while A_{max} is lower in plants grown and measured at 20°C, 445 compared to those grown and measured at 30°C and 38°C, A_{max} in plants grown and measured at 30°C 446 and 38°C is similar (Fig. 7B).

447 Impact of C₄ model species on the analysis

448 Maize, sorghum and *Setaria viridis*, three frequently studied species in C₄ research, dominated our A/C_1 449 dataset (Fig. S1). In fact, data collected from these C4 model species are widely used to model the response 450 of C4 non-model species at the ecosystem level [e.g. in the Community Land Model (Lawrence et al., 451 2019)]. Consequently, we investigated whether V_{pmaxA} and A_{max} of these C₄ model species are 452 representative of the non-model species. We found no significant difference in V_{pmaxA} and A_{max} between 453 the C₄ model and non-model species (P = 0.230 and 0.155 for V_{pmaxA} and A_{max} , respectively; Table 1 – species group). However, it is worth noting that the interaction between species group and growth location 454 455 was significant, due to the lower photosynthetic capacity of non-model species grown outdoors (see above 456 and Fig. 4). Therefore, we suggest that while our results show that C4 model species have similar 457 photosynthetic capacity compared to non-model species, care must be taken when extending this finding 458 to natural outdoor settings, the conditions of most interest to ecologists and Earth System Model 459 modellers. Lastly, our results indicate that our multivariate model was not biased by the weighting of data 460 from C₄ model species.

461 Discussion

462 This study aimed to explore general patterns of C₄ photosynthetic capacity, indicated by V_{pmaxA} and A_{max} , 463 among a wide range of species grown and measured at various conditions. We found that V_{pmaxA} and A_{max} 464 were tightly coupled in a way that reflects the limitation states of C_4 photosynthesis at different C_i and 465 T_{leaf} conditions, supporting Hypothesis 1 (Fig. 2), though this correlation breaks down at higher 466 temperatures. V_{pmaxA} and A_{max} were similar among the three C₄ subtypes and two growth forms, rejecting 467 Hypothesis 2 (Fig. 3A-D), but were influenced by growth and measurement conditions. Indoor plants 468 exhibited higher V_{pmaxA} and A_{max} compared to their outdoor counterparts, supporting Hypothesis 3 (Fig. 469 3E&F). Both V_{pmaxA} and A_{max} decreased with increasing mean growth T_{max} , supporting Hypothesis 4 (Fig. 470 5A&B). Both V_{pmaxA} and A_{max} increased with increasing measurement T_{leaf} and *PPFD*, providing partial 471 support for Hypothesis 5 (Fig. 6). Interestingly, the response of A_{max} to T_{leaf} depended on mean T_{max} , 472 indicating that thermal acclimation alters the thermal sensitivity of photosynthetic capacity in C₄ species 473 (Fig. 7). There was no significant effect of growth CO_2 concentration on photosynthetic capacity, leading 474 us to accept Hypothesis 6, though more data are needed at low and high growth CO₂ concentrations to 475 further confirm this result (Fig. 5C&D). Lastly, we explored whether the more commonly measured C_4

476 model species (maize, sorghum and *Setaria viridis*) were good representatives of a wide range of C₄ plants. 477 We found no evidence of generalised superior photosynthetic capacity in C₄ model species when 478 compared to non-model species (e.g. native C4 grasses and eudicots as a whole), in opposition to 479 Hypothesis 7 (Table 1), although C₄ model species had higher photosynthetic capacity than non-model 480 species when grown outdoors (Fig. 4). Overall, our analysis highlights that C4 photosynthesis is strongly 481 affected by growth and measurement conditions but is largely similar across C₄ species from different 482 biochemical subtypes and growth forms. We discuss the potential reasons and consequences of our 483 findings below.

484 Coupling between V_{pmaxA} and A_{max} reflects photosynthetic limitations

We found a strong positive correlation between estimated V_{pmaxA} and A_{max} in the T_{leaf} range of 25-30°C 485 (Fig. 2B). At T_{leaf} above 30°C, the positive correlation between V_{pmaxA} and A_{max} weakened, with A_{max} 486 487 showing less pronounced change per unit increase in V_{pmaxA} (Fig. 2C-D). This result agrees with findings 488 derived from flux control analysis (von Caemmerer & Furbank, 2016) and data syntheses (Pignon & Long, 489 2020), and highlights differences in the temperature-dependent responses of biochemical processes 490 underpinning V_{pmaxA} and A_{max} , such that A_{max} was less temperature sensitive at high T_{leaf} than V_{pmaxA} . The A_{max} in C₄ plants is primarily determined by the maximum carboxylation activity of Rubisco (V_{cmax}) and 491 492 the rate of RuBP regeneration at high C_i , with a reduction in either of these leading to a lower A_{max} 493 (Furbank et al., 1996; von Caemmerer et al., 1997; von Caemmerer & Furbank, 1999). At high T_{leaf}, 494 Rubisco inactivation due to a decrease in Rubisco activase activity (Hendrickson et al., 2008; Salesse-495 Smith *et al.*, 2018) reduces V_{cmax} and ultimately A_{max} . The RuBP regeneration rate can be limiting when 496 measurement PPFD or the capacity of Calvin cycle enzymes, other than Rubisco, limit net CO₂ 497 assimilation rate at high C_i, or when the thylakoid membrane is damaged at high temperatures (Peixoto & 498 Sage, 2017). The limitation imposed by RuBP regeneration is expected to be pronounced in C₄ plants, 499 which have lower concentrations of RuBP compared to C_3 plants (Arrivault *et al.*, 2019). However, this 500 limitation is likely not related to the abundance and activity of sedoheptulose-1,7-bisphosphatase (a rate-501 limiting enzyme in Calvin cycle) (Ermakova et al. 2022). An RuBP limitation could further constrain 502 photosynthesis as C_i increases, as an increase in C_i promotes Rubisco carboxylase activity and leads to 503 more RuBP being consumed. In our dataset, the majority of A/C_i measurements were done at PPFD > 504 1,500 μ mol photon m⁻² s⁻¹, including those measured at $T_{\text{leaf}} > 35^{\circ}\text{C}$ (Fig. S7). However, photosynthesis in C₄ plants is usually not light-saturated even at 1,500 μ mol photon m⁻² s⁻¹ (Ermakova *et al.*, 2019, 2023). 505

There is evidence that due to adaption to tropical/subtropical climates where high-light environments are common, C₄ plants evolved a series of photosynthetic traits that allow them to grow under high light (Wasilewska-Dębowska *et al.*, 2022). Adaptation to high light conditions may arguably put C₄ plants at a disadvantage in light conditions that are suboptimal for them but saturating for C₃ plants. These findings highlight the complex interplay between temperature, biochemical limitations, and light availability in regulating C₄ photosynthesis, which in part has become a bottleneck for C₄ photosynthesis modelling at the ecosystem level (Knauer *et al.*, 2023).

513 Photosynthetic capacity is similar across biochemical types and growth forms

514 Our results show that variation in C₄ biochemical subtypes and growth forms does not lead to systematic 515 variation in photosynthetic capacity (Fig. 3A-D). In general, popular model species (C₄ NADP-ME type 516 grasses: maize, sorghum and *Setaria viridis*) exhibit similar photosynthetic capacity when compared to 517 non-model C₄ species representing the three C₄ biochemical subtypes (Table 1). However, when growth 518 location is considered, photosynthetic capacity of C₄ model species is higher than that of non-model 519 species when both are grown outdoors (Fig. 4). These results suggest that photosynthetic capacity data 520 collected from model species could overestimate the productivity of non-model species in outdoor 521 conditions. These findings are particularly important for the modelling community, given that the current 522 C₄ vegetation model at the ecosystem level was developed based on parameters measured in maize grown 523 in controlled environments (Collatz et al., 1992; Lawrence et al., 2019), and likely overestimates the 524 productivity of C₄ native and eudicots grown in the field. The lack of a significant effect of C₄ subtypes 525 on photosynthetic capacity further emphasises the biochemical flexibility in the C₄ pathway. There is 526 growing molecular, biochemical and physiological evidence suggesting that the three classical subtypes, 527 originally defined based on early ¹⁴C-labelling studies (Hatch & Slack, 1970; Hatch, 1971), can be 528 biochemically flexible (Furbank, 2011; Bräutigam et al., 2014; Wang et al., 2014; Sales et al., 2021), 529 which in turn diminishes any potential differences in photosynthetic capacity among the subtypes. This 530 biochemical flexibility may be regulated by developmental and environmental variation, further 531 highlighting the role of environmental effects on C₄ photosynthesis (see below).

532 Growth location affects photosynthetic capacity

533 Our results showed that growth and measurement conditions are the major factors affecting photosynthetic 534 capacity in C₄ plants. Indoor plants exhibit consistently higher V_{pmaxA} and A_{max} than their outdoor

counterparts (Fig. 3E&F), despite indoor plants being grown in warmer environments which might 535 536 suppress photosynthetic capacity (Fig. S4A). Plants grown in indoor, controlled environments are usually 537 well-watered and fertilised, and are not challenged with pests and extreme conditions (e.g. heat waves or 538 frost stress) (Poorter et al., 2016). The lower photosynthetic capacity in outdoor non-model species is 539 likely due to the more stressful conditions they experience, which are primarily characterised by lower 540 water and nutrient availability than indoor-grown plants. For example, long-term drought (i.e. >100 days) 541 could cause significant reductions in stomatal conductance, net photosynthetic rate and V_{pmaxA} , with such 542 effects being exacerbated in N-limited C₄ plants (Markelz et al., 2011). Interestingly, the impact of drought 543 on photosynthetic capacity (i.e. A_{max}) was not seen in two well-fertilised C₄ grasses, *Dactyloctenium* 544 aegyptium and Schoenefeldia gracilis (Maroco et al., 2000), suggesting changes in photosynthetic 545 capacity may be associated with interactive effects of drought and N limitation on plants. In contrast, C4 546 model species-particularly maize and sorghum-have been selectively bred for improved drought 547 tolerance (Lopes et al., 2011) and are typically well-watered and well-fertilised in crop fields due to their 548 high agricultural value. These effects may have contributed to higher photosynthetic capacity in C₄ model 549 species grown outdoors (Fig. 4). Further research is needed to dissect the mechanism underpinning the combined effects of drought and N limitation on photosynthesis and to identify why outdoor growth 550 551 conditions affect non-model species more than model C₄ species.

552 Temperature significantly influences photosynthetic capacity

553 We observed that both V_{pmaxA} and A_{max} decreased with increasing mean T_{max} (Fig. 5A&B). This finding is 554 supported by previous work (Berry & Bjorkman, 1980) and aligns with our knowledge of thermal 555 acclimation (Way & Yamori, 2014). Plants grown in warmer conditions often reduce photosynthetic capacity (i.e. $V_{\text{pmax}A}$ and A_{max}), while maintaining comparable (or even higher) A at their growth 556 557 temperatures compared to control plants. This decrease in photosynthetic capacity is likely due to reduced 558 photosynthetic enzyme concentrations. Warmer temperatures allow plants to achieve the same rate of 559 photosynthesis with lower enzyme concentrations because higher temperatures enhance enzyme activity 560 (Yamori et al., 2014; Way & Yamori, 2014). Evidence supports this idea in C₄ plants: Dwyer et al. (2007) 561 compared C₄ Panicum coloratum, Cenchrus ciliaris and Flaveria bidentis grown at moderate and high 562 temperatures and found that warm-grown plants had reduced photosynthetic capacity (i.e. A_{max}), 563 underpinned by lower concentrations of Rubisco and chloroplastic electron transport chain proteins. 564 However, the authors did not find a significant effect of growth temperature on the activity or

565 concentration of PEPc (measured at the respective growth temperatures), suggesting that PEPc may have 566 high thermal stability (Chen et al., 1994; Chinthapalli et al., 2002; Boyd et al., 2015). Thus, it is possible 567 that the decrease in V_{pmaxA} with increasing mean T_{max} observed in our study may be reflective of changes 568 in PEP regeneration via PPDK rather than PEPc capacity *per se*. In *Miscanthus* × *giganteus*, the activity 569 and capacity of PPDK per unit leaf area increases in plants grown at chilling temperature, compared to 570 their warm-grown controls (Wang *et al.*, 2008). Furthermore, whether A/C_i measurements are taken at 571 saturated PPFD may also affect PEP regeneration, which requires two photosynthetically-generated ATP 572 per PEP converted (Hatch, 1987). Although some of the measurements in our dataset were made at PPFD < 1000 μ mol m⁻² s⁻¹, we found no significant effect of lower measurement *PPFD* on the responses of 573 V_{pmaxA} and A_{max} to intrinsic or extrinsic factors (Table 1; Table S1). Further research is needed to explore 574 575 how PEPc and PPDK capacities are coordinated under thermal acclimation.

576 Our results highlight that both V_{pmaxA} and A_{max} increase with measurement T_{leaf} (Fig. 6A&B), which is indicative of enhanced enzymatic activities at higher measurement temperatures. According to 577 578 an *in vitro* study in *Setaria viridis*, the capacities of major C₄ photosynthetic enzymes generally increase 579 with rising T_{leaf} due to a corresponding rise in enzyme activities (Boyd *et al.*, 2015). The carboxylation 580 activities of Rubisco and PEPc increase exponentially between 10°C and 40-45°C, before declining (Chen 581 et al., 1994; Chinthapalli et al., 2002; Boyd et al., 2015). However, the extent to which this temperature 582 response of enzymes holds true in vivo remains uncertain and needs to be confirmed with high-resolution 583 photosynthesis-temperature response curves across a diverse range of C₄ species.

584 Finally, our analysis highlights that the response of A_{max} to T_{leaf} is influenced by mean T_{max} , whereas that of V_{pmaxA} is not (Table 1; Fig. 7). We found that A_{max} increases to a greater extent with 585 586 increasing T_{leaf} in cool-grown plants compared to plants grown in warmer conditions, suggesting that A_{max} 587 of cool-grown plants is more sensitive to changes in T_{leaf} (Fig. 7B). This finding has implications for the 588 photosynthetic performance of C₄ plants under future climates. In a high CO₂ world, C₄ plants will be able 589 to achieve photosynthetic performance closer to their maximum potential (i.e. A_{max}), as higher atmospheric $[CO_2]$ increases C_i and moves plants away from potential V_{pmaxA} limitations of photosynthesis. Our data 590 suggest that A_{max} of C₄ plants grown under future, hotter climates will be less affected by daily temperature 591 592 fluctuations compared to plants grown under the current climate (i.e. A_{max} will be more stable; see circles 593 in Fig. 7). This reduced thermal sensitivity of A_{max} at higher temperatures may help buffer photosynthesis 594 in future climates, where heat waves are predicted to occur more frequently and intensely (Brown, 2020). 595 However, this response could also limit the ability of warm-grown C₄ plants to achieve high A_{max} at the

high T_{leaf} conditions they will experience (Fig. 7), which may suppress CO₂ uptake in future climate conditions. As we saw no effect of elevated growth CO₂ concentrations on either V_{pmaxA} or A_{max} (Fig. 508 5C&D), in line with field studies of C₄ species (Leakey *et al.*, 2006; Leakey, 2009; Markelz *et al.*, 2011), these temperature effects are likely to be stronger controls on C₄ photosynthetic capacity in future climates than predicted increases in CO₂ levels. However, further research is needed to fully understand how climate change affects the photosynthetic physiology of C₄ plants.

602 Future perspectives

603 Looking forward, our analyses highlight areas where more studies are warranted. For example, the 604 physiology of C₄ eudicots and sedges is underrepresented in our dataset, although eudicots and sedges comprise 38% of C₄ plants (Sage, 2017) and thrive in extreme environments [e.g. Haloxylon, a C₄ desert 605 606 shrub (Feng *et al.*, 2023)]. Understanding how C_4 photosynthesis responds to these extreme environments 607 could provide insights into improving how other plants (e.g. C₃ crops) cope with extreme environments. 608 Furthermore, our results show that A_{max} responds differently to growth and measurement temperatures than does V_{pmaxA} . We explored potential biochemical limitations that underpin A_{max} and V_{pmaxA} separately, 609 610 in addition to the response of these limitations to changes in temperature. However, it is unclear to what 611 extent the individual biochemical limitations may interact and affect overall photosynthetic capacity. If 612 one assumes that PEP regeneration is not limited, V_{pmaxA} is largely determined by the property of PEPc 613 (e.g. activation state and activity of the enzyme) and its temperature sensitivity likely reflects the temperature sensitivity of PEPc. In contrast, A_{max} is determined by a variety of biochemical processes (e.g. 614 615 Rubisco carboxylation, rates of electron transport). Therefore, understanding how these processes interact 616 in response to changing temperatures is crucial for fully grasping the overall response of A_{max} .

617 Conclusion

618 C_4 plants play a crucial role in carbon exchange and food security on a global scale. There has been a 619 growing number of studies dissecting the mechanism of C_4 photosynthesis in specific contexts, yet how 620 C_4 photosynthetic capacity responds to differences in biochemical subtypes, plant functional types, and 621 growth and measurement conditions among a wide range of species remains unclear. Our study uncovers 622 broad patterns of photosynthetic capacity from 74 C_4 species and highlights that environmental conditions 623 play a dominant role in determining C_4 photosynthetic capacity. Importantly, we demonstrate that while 624 the current simplified parameterisation of C_4 NADP-ME-type photosynthesis in leaf- and ecosystem-level 625 models likely represents species of all three biochemical types, it overestimates the photosynthetic 626 capacity of C₄ native species in the field conditions. Future research should aim to refine current parameter 627 values by incorporating detailed equations that capture the effects of abiotic factors, such as water and 628 nutrient availability, on photosynthetic capacity (Smith et al., 2019), while also examining their link to 629 the leaf economics spectrum (Monson *et al.*, 2025). Additionally, we compile a C₄ A/C_i database for 630 community use, addressing the shortage of accessible raw data for C₄ gas exchange. For example, the 631 TRY leaf-trait database (http://www.try-db.org), one of the most comprehensive databases of plant 632 physiological trait data in the world, contains leaf photosynthetic point measurements for only 591 C₄ species, out of 69,000 terrestrial species measured, and no C4 photosynthetic response curves (Kattge et 633 634 al., 2011, 2020). Our database opens new avenues for future studies involving big data analysis for C₄ 635 plants, such as simulation modelling.

636 Acknowledgements

637 This work was conducted as part of the C₄ Photosynthesis Working Group supported by the John Wesley 638 Powell Center for Analysis and Synthesis, funded by the US Geological Survey Award Number 20-07-639 0232. Additional support for D.A.W. came from The Australian National University Futures Scheme, and 640 for Y.F. from an Australian Research Council Discovery Project (DP230103122). D.W.A.N. was 641 supported by an ARC Future Fellowship (FT220100276). A.D.B.L. was supported by the DOE Center for 642 Advanced Bioenergy and Bioproducts Innovation (U.S. Department of Energy, Office of Science, 643 Biological and Environmental Research Program under Award Number DE-SC0018420). F.A.B. was 644 supported by the Natural Environment Research Council (NE/W00674X/1). K.P. was supported by U.S. 645 National Science Foundation grants (IOS-2312181, IOS-2406533, IOS-1546838, MCB-1617020; DBI-646 2213983 and OISE-2434687), U.S. Department of Energy, Office of Science, Office of Biological and 647 Environmental Research, Genomic Science Program grants (DE-SC0018277, DE-SC0020366, DE-648 SC0023160, and DE-SC0021286, and DE-SC0008769), and Carnegie Venture Grant (10908). M.L. is 649 supported by an Australian Research Council Discovery Project (DP130102243). Any use of trade, firm, 650 or product names is for descriptive purposes only and does not imply endorsement by the U.S. 651 Government.

652 Conflict of interest

None declared.

654 Author contribution

- 655 D.A.W. conceived the concept for the paper within the discussions provided by the US Geological Survey
- 656 Powell Center C₄ Working Group. Y.F. complied the dataset, coded the scripts and ran the analyses with
- 657 help from D.W.A.N. and B.E.M. N.G.S., E.A.A., F.A.B., F.R.D., M.E., P.F., R.T.F., S.H.G., O.G., L.G.,
- 658 V.J., A.D.B.L., S.L., M.L., V.S.P., M.M.P., B.V.S., S.v.C., R.W. and D.A.W. provided data to this study.
- 659 Y.F. and D.A.W. wrote the first draft, and all authors contributed substantially to improving the draft.

660 Data availability

661 Upon publication, all data will be available as a Supplementary Dataset and downloadable from GitHub.

662 Supplementary material

- 663 Dataset S1 C₄ A/C_i data collated for this study.
- 664 Dataset S2 Estimated apparent V_{pmaxA} and A_{max} used in the analysis.
- 665 Table S1 Results of multivariate linear mixed-effects models for data with $PPFD > 1000 \ \mu mol$ photon 666 m⁻² s⁻¹.
- 667 Figure S1 Species distribution within the categories of growth forms and growth locations.
- 668 Figure S2 Correlation of mean T_{max} and T_{min} .
- 669 Figure S3 Data distribution of V_{pmaxA} and A_{max} in categories of C₄ subtypes and growth forms over T_{leaf} ,
- 670 mean T_{max} and *PPFD*.
- 671 Figure S4 Histograms describing the data distribution across the spectrum of mean T_{max} and T_{leaf} for
- 672 growth location, growth form and important species group.
- Figure S5 V_{pmaxA} and A_{max} plotted by C₄ biochemical subtypes, with symbols highlighting the effect of phylogeny.
- Figure S6 Relationships between V_{pmaxA} and A_{max} with mean T_{max} for measurements done at T_{leaf} between
- 676 25 and 30°C.
- Figure S7 Correlation of mean T_{max} and T_{min} for data with $PPFD > 1,500 \ \mu\text{mol}$ photon m⁻² s⁻¹.

678 Reference

- 679 Anderson LJ, Maherali H, Johnson HB, Polley HW, Jackson RB. 2001. Gas exchange and 680 photosynthetic acclimation over subambient to elevated CO_2 in a C_3 - C_4 grassland. *Global Change* 681 *Biology* **7**: 693–707.
- 682 Arena C, Vitale L, De Santo AV. 2011. Influence of irradiance on photosynthesis and PSII
- photochemical efficiency in maize during short-term exposure at high CO₂ concentration.
 Photosynthetica 49: 267–274.

Arrivault S, Alexandre Moraes T, Obata T, Medeiros DB, Fernie AR, Boulouis A, Ludwig M, Lunn
 JE, Borghi GL, Schlereth A, et al. 2019. Metabolite profiles reveal interspecific variation in operation
 of the Calvin–Benson cycle in both C₄ and C₃ plants. *Journal of Experimental Botany* 70: 1843–1858.

- 688 **Atkin OK, Tjoelker MG**. **2003**. Thermal acclimation and the dynamic response of plant respiration to 689 temperature. *Trends in Plant Science* **8**: 343–351.
- 690 **Bauwe H. 1986**. An efficient method for the determination of K_m values for HCO_3^- of 691 phosphoenolpyruvate carboxylase. *Planta* **169**: 356–360.
- Berry J, Bjorkman O. 1980. Photosynthetic Response and Adaptation to Temperature in Higher
 Plants. Annual Review of Plant Biology 31: 491–543.
- Bissinger K, Khoshravesh R, Kotrade JP, Oakley J, Sage TL, Sage RF, Hartmann HEK, Kadereit
 G. 2014. *Gisekia* (Gisekiaceae): Phylogenetic relationships, biogeography, and ecophysiology of a
 poorly known C₄ lineage in the Caryophyllales. *American Journal of Botany* 101: 499–509.
- Bloom AJ, Asensio JSR, Randall L, Rachmilevitch S, Cousins AB, Carlisle EA. 2012. CO₂
 enrichment inhibits shoot nitrate assimilation in C₃ but not C₄ plants and slows growth under nitrate in
 C₃ plants. *Ecology* 93: 355–367.
- Boyd RA, Gandin A, Cousins AB. 2015. Temperature Responses of C₄ Photosynthesis: Biochemical
 Analysis of Rubisco, Phospho*enol*pyruvate Carboxylase, and Carbonic Anhydrase in *Setaria viridis*.
 Plant Physiology 169: 1850–1861.
- Bräutigam A, Schliesky S, Külahoglu C, Osborne CP, Weber APM. 2014. Towards an integrative
 model of C₄ photosynthetic subtypes: insights from comparative transcriptome analysis of NAD-ME,
 NADP-ME, and PEP-CK C₄ species. *Journal of Experimental Botany* 65: 3579–3593.
- Brown SJ. 2020. Future changes in heatwave severity, duration and frequency due to climate change
 for the most populous cities. *Weather and Climate Extremes* 30: 100278.
- von Caemmerer S. 2000. Modeling C₄ photosynthesis. In: Biochemical models of leaf photosynthesis.
 Collingwood, Australia: CSIRO Publishing, 91–122.
- von Caemmerer S. 2021. Updating the steady-state model of C₄ photosynthesis. *Journal of Experimental Botany* 72: 6003–6017.
- 712 von Caemmerer S, Evans JR, Hudson GS, Andrews TJ. 1994. The kinetics of ribulose-1,5-
- 513 bisphosphate carboxylase/oxygenase in vivo inferred from measurements of photosynthesis in leaves
- of transgenic tobacco. *Planta* **195**: 88–97.

- von Caemmerer S, Furbank RT. 1999. Modeling C₄ Photosynthesis. In: Sage RF, Monson RK, eds.
 C₄ Plant Biology. San Diego: Academic Press, 173–211.
- von Caemmerer S, Furbank RT. 2016. Strategies for improving C₄ photosynthesis. *Current Opinion in Plant Biology* 31: 125–134.
- 719 von Caemmerer S, Millgate A, Farquhar GD, Furbank RT. 1997. Reduction of Ribulose-1,5-
- Bisphosphate Carboxylase/Oxygenase by Antisense RNA in the C₄ Plant *Flaveria bidentis* Leads to
 Reduced Assimilation Rates and Increased Carbon Isotope Discrimination. *Plant Physiology* **113**: 469–
- 722 477.
- Carmo-Silva AE, Powers SJ, Keys AJ, Arrabaça MC, Parry MAJ. 2008. Photorespiration in C₄
 grasses remains slow under drought conditions. *Plant, Cell & Environment* 31: 925–940.
- Cavaco AM, Da Silva AB, Arrabaça MC. 2003. Effects of long-term chilling on growth and
 photosynthesis of the C₄ gramineae *Paspalum dilatatum*. *Physiologia Plantarum* 119: 87–96.
- 727 Chen D-X, Coughenour MB, Knapp AK, Owensby CE. 1994. Mathematical simulation of C₄ grass
 728 photosynthesis in ambient and elevated CO₂. *Ecological Modelling* 73: 63–80.
- Chen J-W, Yang Z-Q, Zhou P, Hai M-R, Tang T-X, Liang Y-L, An T-X. 2013. Biomass accumulation
 and partitioning, photosynthesis, and photosynthetic induction in field-grown maize (*Zea mays* L.) under
 low- and high-nitrogen conditions. *Acta Physiologiae Plantarum* 35: 95–105.
- Chinthapalli B, Raghavendra AS, Rishi AS, Arun G. 2002. Phosphonelopyruvate carboxylase from
 C₄ plants: properties and regulation. *Reviews in Plant Biochemistry and Biotechnology* 1: 143–159.
- Collatz GJ, Ribas-Carbo M, Berry JA. 1992. Coupled Photosynthesis-Stomatal Conductance Model
 for Leaves of C₄ Plants. *Functional Plant Biology* 19: 519–538.
- Cousins AB, Bloom AJ. 2003. Influence of elevated CO₂ and nitrogen nutrition on photosynthesis and
 nitrate photo-assimilation in maize (*Zea mays* L.). *Plant, Cell & Environment* 26: 1525–1530.
- Cowling SA, Jones CD, Cox PM. 2007. Consequences of the evolution of C₄ photosynthesis for
 surface energy and water exchange. *Journal of Geophysical Research: Biogeosciences* 112.
- 740 Cunniff J, Jones G, Charles M, Osborne CP. 2017. Yield responses of wild C₃ and C₄ crop
- 741 progenitors to subambient CO_2 : a test for the role of CO_2 limitation in the origin of agriculture. *Global* 742 *Change Biology* **23**: 380–393.
- Cunniff J, Osborne CP, Ripley BS, Charles M, Jones G. 2008. Response of wild C₄ crop progenitors
 to subambient CO₂ highlights a possible role in the origin of agriculture. *Global Change Biology* 14:
 576–587.
- 746 Danila FR, Thakur V, Chatterjee J, Bala S, Coe RA, Acebron K, Furbank RT, von Caemmerer S,
- 747 Quick WP. 2021. Bundle sheath suberisation is required for C₄ photosynthesis in a *Setaria viridis* 748 mutant. *Communications Biology* 4: 254.
- DiMario RJ, Cousins AB. 2019. A single serine to alanine substitution decreases bicarbonate affinity
 of phosphoenolpyruvate carboxylase in C₄ *Flaveria trinervia*. *Journal of Experimental Botany* 70: 995–
 1004.

- Ding R, Xie J, Mayfield-Jones D, Zhang Y, Kang S, Leakey ADB. 2022. Plasticity in stomatal
 behaviour across a gradient of water supply is consistent among field-grown maize inbred lines with
 varying stomatal patterning. *Plant, Cell & Environment* 45: 2324–2336.
- Dwyer SA, Ghannoum O, Nicotra A, von Caemmerer S. 2007. High temperature acclimation of C₄
 photosynthesis is linked to changes in photosynthetic biochemistry. *Plant, Cell & Environment* 30: 53–
 66.
- Frmakova M, Lopez-Calcagno PE, Furbank RT, Raines CA, von Caemmerer S. 2022. Increased
 sedoheptulose-1,7-bisphosphatase content in Setaria viridis does not affect C₄ photosynthesis. *Plant Physiology* 191: 885–893.
- 761 Ermakova M, Lopez-Calcagno PE, Raines CA, Furbank RT, von Caemmerer S. 2019.
- 762 Overexpression of the Rieske FeS protein of the Cytochrome b6f complex increases C₄ photosynthesis
 763 in Setaria viridis. Communications Biology 2: 1–12.
- 764 Ermakova M, Woodford R, Fitzpatrick D, Nix SJ, Zwahlen SM, Farquhar GD, von Caemmerer S,
- Furbank RT. 2024. Chloroplast NADH dehydrogenase-like complex-mediated cyclic electron flow is the main electron transport route in C₄ bundle sheath cells. *New Phytologist* 243: 2187–2200.
- 767 Ermakova M, Woodford R, Taylor Z, Furbank RT, Belide S, von Caemmerer S. 2023. Faster
- induction of photosynthesis increases biomass and grain yield in glasshouse-grown transgenic
- Sorghum bicolor overexpressing Rieske FeS. *Plant Biotechnology Journal* **21**: 1206–1216.
- **Evans JR. 1989.** Photosynthesis and nitrogen relationships in leaves of C₃ plants. *Oecologia* **78**: 9–19.
- Evans JR, von Caemmerer S. 1996. Carbon dioxide diffusion inside leaves. *Plant Physiology* 110:
 339–346.
- Fan Y, Asao S, Furbank RT, von Caemmerer S, Day DA, Tcherkez G, Sage TL, Sage RF, Atkin
 OK. 2022. The crucial roles of mitochondria in supporting C₄ photosynthesis. *New Phytologist* 233:
 1083–1096.
- Fay PA, Jin VL, Way DA, Potter KN, Gill RA, Jackson RB, Wayne Polley H. 2012. Soil-mediated
 effects of subambient to increased carbon dioxide on grassland productivity. *Nature Climate Change* 2:
 742–746.
- 779 Feng XP, Chen Y, Qi YH, Yu CL, Zheng B-S, Brancourt-Hulmel M, Jiang D-A. 2012. Nitrogen
- enhanced photosynthesis of Miscanthus by increasing stomatal conductance and phosphoenolpyruvate
 carboxylase concentration. *Photosynthetica* 50: 577–586.
- **Feng X, Liu R, Li C, Zhang H, Slot M**. **2023**. Contrasting responses of two C₄ desert shrubs to drought
- but consistent decoupling of photosynthesis and stomatal conductance at high temperature.
 Environmental and Experimental Botany 209: 105295.
 - 785 Friedlingstein P, O'Sullivan M, Jones MW, Andrew RM, Hauck J, Landschützer P, Le Quéré C, Li
 - H, Luijkx IT, Olsen A, et al. 2025. Global Carbon Budget 2024. Earth System Science Data 17: 965–
 1039.

Friesen PC, Peixoto MM, Busch FA, Johnson DC, Sage RF. 2014. Chilling and frost tolerance in
 Miscanthus and *Saccharum* genotypes bred for cool temperate climates. *Journal of Experimental Botany* 65: 3749–3758.

- Furbank RT. 2011. Evolution of the C₄ photosynthetic mechanism: are there really three C₄ acid
 decarboxylation types? *Journal of Experimental Botany* 62: 3103–3108.
- Furbank RT, Chitty JA, von Caemmerer S, Jenkins CLD. 1996. Antisense RNA Inhibition of RbcS
 Gene Expression Reduces Rubisco Level and Photosynthesis in the C₄ Plant *Flaveria bidentis*. *Plant Physiology* 111: 725–734.
- **Gan SH, Sage RF**. **2024**. Stomatal response to VPD in C₄ plants with different biochemical subpathways. *Plant, Cell & Environment* **47**: 3466–3477.
- Ge ZM, Zhang LQ, Yuan L, Zhang C. 2014. Effects of salinity on temperature-dependent
 photosynthetic parameters of a native C₃ and a non-native C₄ marsh grass in the Yangtze Estuary,
 China. *Photosynthetica* 52: 484–492.
- 801 **Ghannoum O, Caemmerer SV, Ziska LH, Conroy JP**. **2000**. The growth response of C₄ plants to 802 rising atmospheric CO₂ partial pressure: a reassessment. *Plant, Cell & Environment* **23**: 931–942.
- Ghannoum O, Evans JR, von Caemmerer S. 2011. Nitrogen and water use efficiency of C₄ plants. In:
 Raghavendra AS, Sage RF, eds. C₄ photosynthesis and related CO₂ concentrating mechanisms.
 Dordrecht: Springer, 129–146.
- 806 Głowacka K, Jørgensen U, Kjeldsen JB, Kørup K, Spitz I, Sacks EJ, Long SP. 2015. Can the
 807 exceptional chilling tolerance of C₄ photosynthesis found in *Miscanthus* × *giganteus* be exceeded?
 808 Screening of a novel *Miscanthus* Japanese germplasm collection. *Annals of Botany* 115: 981–990.
- 809 Griffith DM, Osborne CP, Edwards EJ, Bachle S, Beerling DJ, Bond WJ, Gallaher TJ, Helliker BR,
- Lehmann CER, Leatherman L, et al. 2020. Lineage-based functional types: characterising functional
 diversity to enhance the representation of ecological behaviour in Land Surface Models. New
 Phytologist 228: 15–23.
- Hatch MD. 1971. The C₄-pathway of photosynthesis: evidence for an intermediate pool of carbon
 dioxide and the identity of the donor C₄-dicarboxylic acid. *Biochemical Journal* 125: 425–432.
- Hatch MD. 1987. C₄ photosynthesis: a unique elend of modified biochemistry, anatomy and
 ultrastructure. *Biochimica et Biophysica Acta* 895: 81–106.
- Hatch MD, Slack CR. 1970. Photosynthetic CO₂-fixation pathways. *Annual Review of Plant Physiology*21: 141–162.
- Heckman RW, Pereira CG, Aspinwall MJ, Juenger TE. 2024. Physiological responses of C₄
 perennial bioenergy grasses to climate change: causes, consequences, and constraints. *Annual Review of Plant Biology* 75: 737–769.
- Hendrickson L, Sharwood R, Ludwig M, Whitney SM, Badger MR, von Caemmerer S. 2008. The
 effects of Rubisco activase on C₄ photosynthesis and metabolism at high temperature. *Journal of Experimental Botany* 59: 1789–1798.

- Horst RJ, Engelsdorf T, Sonnewald U, Voll LM. 2008. Infection of maize leaves with Ustilago maydis
 prevents establishment of C₄ photosynthesis. Journal of Plant Physiology 165: 19–28.
- 827 Huntingford C, Zelazowski P, Galbraith D, Mercado LM, Sitch S, Fisher R, Lomas M, Walker AP,
- Jones CD, Booth BB. 2013. Simulated resilience of tropical rainforests to CO₂-induced climate change. *Nature Geoscience* 6: 268.
- 830 **Kakani VG, Boote KJ, Reddy KR, Lang DJ**. **2008a**. Response of bahiagrass carbon assimilation and 831 photosystem activity to below optimum temperatures. *Functional Plant Biology* **35**: 1243.
- Kakani VG, Surabhi GK, Reddy KR. 2008b. Photosynthesis and fluorescence responses of C₄ plant
 Andropogon gerardii acclimated to temperature and carbon dioxide. *Photosynthetica* 46: 420–430.
- Kattge J, Bönisch G, Díaz S, Lavorel S, Prentice IC, Leadley P, Tautenhahn S, Werner GDA,
 Aakala T, Abedi M, et al. 2020. TRY plant trait database enhanced coverage and open access.
 Global Change Biology 26: 119–188.
- Kattge J, Díaz S, Lavorel S, Prentice IC, Leadley P, Bönisch G, Garnier E, Westoby M, Reich PB,
 Wright IJ, et al. 2011. TRY a global database of plant traits. *Global Change Biology* 17: 2905–2935.
- Khoshravesh R, Stata M, Busch FA, Saladié M, Castelli JM, Dakin N, Hattersley PW, Macfarlane
 TD, Sage RF, Ludwig M, *et al.* 2020. The evolutionary origin of C₄ photosynthesis in the grass subtribe
- 841 Neurachninae. *Plant Physiology* **182**: 566–583.
- Kim S-H, Sicher RC, Bae H, Gitz DC, Baker JT, Timlin DJ, Reddy VR. 2006. Canopy
 photosynthesis, evapotranspiration, leaf nitrogen, and transcription profiles of maize in response to CO₂
 enrichment. *Global Change Biology* 12: 588–600.
- Knauer J, Cuntz M, Smith B, Canadell JG, Medlyn BE, Bennett AC, Caldararu S, Haverd V. 2023.
 Higher global gross primary productivity under future climate with more advanced representations of
 photosynthesis. *Science Advances* 9: eadh9444.
- 848 **Kubien DS, Sage RF**. **2004**. Low-temperature photosynthetic performance of a C₄ grass and a co-849 occurring C₃ grass native to high latitudes. *Plant, Cell & Environment* **27**: 907–916.
- Laisk A, Edwards GE. 1997. CO₂ and Temperature-dependent Induction in C₄ Photosynthesis: an
 Approach to the Hierarchy of Rate-limiting Processes. *Functional Plant Biology* 24: 505–516.
- 852 Lawrence DM, Fisher RA, Koven CD, Oleson KW, Swenson SC, Bonan G, Collier N, Ghimire B,
- van Kampenhout L, Kennedy D, et al. 2019. The Community Land Model Version 5: Description of
 New Features, Benchmarking, and Impact of Forcing Uncertainty. *Journal of Advances in Modeling Earth Systems* 11: 4245–4287.
- Leakey ADB. 2009. Rising atmospheric carbon dioxide concentration and the future of C₄ crops for
 food and fuel. *Proceedings of the Royal Society B: Biological Sciences* 276: 2333–2343.
- Leakey ADB, Ferguson JN, Pignon CP, Wu A, Jin Z, Hammer GL, Lobell DB. 2019. Water Use
 Efficiency as a Constraint and Target for Improving the Resilience and Productivity of C3 and C4
 Crops. Annual Review of Plant Biology 70: 781–808.

- Leakey ADB, Uribelarrea M, Ainsworth EA, Naidu SL, Rogers A, Ort DR, Long SP. 2006.
- 862 Photosynthesis, productivity, and yield of maize are not affected by open-air elevation of CO₂ 863 concentration in the absence of drought. *Plant Physiology* **140**: 779–790.
- **Leegood RC**. **2002**. C₄ photosynthesis: principles of CO₂ concentration and prospects for its introduction into C₃ plants. *Journal of Experimental Botany* **53**: 581–590.
- Li S, Courbet G, Ourry A, Ainsworth EA. 2019. Elevated ozone concentration reduces photosynthetic
 carbon gain but does not alter leaf structural traits, nutrient composition or biomass in switchgrass.
 Plants 8: 85.
- Li S, Moller CA, Mitchell NG, Lee D, Ainsworth EA. 2021. Bioenergy sorghum maintains
 photosynthetic capacity in elevated ozone concentrations. *Plant, Cell & Environment* 44: 729–746.
- Li S, Moller CA, Mitchell NG, Lee D, Sacks EJ, Ainsworth EA. 2022. Testing unified theories for
 ozone response in C₄ species. *Global Change Biology* 28: 3379–3393.
- Liu X, Qiao Y, Zhou W, Dong W, Gu L. 2023. Determinants of photochemical characteristics of the
 photosynthetic electron transport chain of maize. *Frontiers in Plant Science* 14: 1279963.
- Liu H, Taylor SH, Xu Q, Lin Y, Hou H, Wu G, Ye Q. 2019. Life history is a key factor explaining functional trait diversity among subtropical grasses, and its influence differs between C_3 and C_4 species. *Journal of Experimental Botany* **70**: 1567–1580.
- Long SP, Woolhouse HW. 1978. The Responses of Net Photosynthesis to Vapour Pressure Deficit
 and CO₂ Concentration in *Spartina townsendii* (sensu lato), a C₄ Species from a Cool Temperate
 Climate. *Journal of Experimental Botany* 29: 567–577.
- Lopes MS, Araus JL, van Heerden PDR, Foyer CH. 2011. Enhancing drought tolerance in C₄ crops.
 Journal of Experimental Botany 62: 3135–3153.
- Luo X, Zhou H, Satriawan TW, Tian J, Zhao R, Keenan TF, Griffith DM, Sitch S, Smith NG, Still
 CJ. 2024. Mapping the global distribution of C₄ vegetation using observations and optimality theory.
 Nature Communications 15: 1219.
- Mantlana KB, Arneth A, Veenendaal EM, Wohland P, Wolski P, Kolle O, Wagner M, Lloyd J. 2008.
 Photosynthetic properties of C₄ plants growing in an African savanna/wetland mosaic. *Journal of Experimental Botany* 59: 3941–3952.
- Markelz RJC, Strellner RS, Leakey ADB. 2011. Impairment of C₄ photosynthesis by drought is
 exacerbated by limiting nitrogen and ameliorated by elevated [CO₂] in maize. *Journal of Experimental Botany* 62: 3235–3246.
- Maroco JP, Pereira JS, Manuela Chaves M. 2000. Growth, photosynthesis and water-use efficiency
 of two C₄ Sahelian grasses subjected to water deficits. *Journal of Arid Environments* 45: 119–137.

Monson RK, Li S, Ainsworth EA, Fan Y, Hodge JG, Knapp AK, Leakey ADB, Lombardozzi D,
 Reed SC, Sage RF, et al. 2025. C₄ photosynthesis, trait spectra, and the fast-efficient phenotype. New
 Phytologist 246: 879–893.

- 897 Naidu SL, Moose SP, AL-Shoaibi AK, Raines CA, Long SP. 2003. Cold tolerance of C₄
- photosynthesis in *Miscanthus* × *giganteus*: adaptation in amounts and sequence of C₄ photosynthetic
 enzymes. *Plant Physiology* 132: 1688–1697.
- Nakagawa S, Lagisz M, O'Dea RE, Pottier P, Rutkowska J, Senior AM, Yang Y, Noble DWA. 2023.
 orchaRd 2.0: An R package for visualising meta-analyses with orchard plots. *Methods in Ecology and Evolution* 14: 2003–2010.
- Nakagawa S, Lagisz M, O'Dea RE, Rutkowska J, Yang Y, Noble DWA, Senior AM. 2021. The
 orchard plot: Cultivating a forest plot for use in ecology, evolution, and beyond. *Research Synthesis Methods* 12: 4–12.
- 906 Nakagawa S, Santos ESA. 2012. Methodological issues and advances in biological meta-analysis.
 907 *Evolutionary Ecology* 26: 1253–1274.
- Nippert JB, Fay PA, Knapp AK. 2007. Photosynthetic traits in C₃ and C₄ grassland species in
 mesocosm and field environments. *Environmental and Experimental Botany* 60: 412–420.
- 910 **Noble DWA, Lagisz M, O'dea RE, Nakagawa S**. **2017**. Nonindependence and sensitivity analyses in ecological and evolutionary meta-analyses. *Molecular Ecology* **26**: 2410–2425.
- Noble DWA, Pottier P, Lagisz M, Burke S, Drobniak SM, O'Dea RE, Nakagawa S. 2022. Meta analytic approaches and effect sizes to account for 'nuisance heterogeneity' in comparative physiology.
- 914 Journal of Experimental Biology **225**: jeb243225.
- Oakley JC, Sultmanis S, Stinson CR, Sage TL, Sage RF. 2014. Comparative studies of C₃ and C₄
 Atriplex hybrids in the genomics era: physiological assessments. Journal of Experimental Botany 65: 3637–3647.
- Osborne CP, Wythe EJ, Ibrahim DG, Gilbert ME, Ripley BS. 2008. Low temperature effects on leaf
 physiology and survivorship in the C₃ and C₄ subspecies of *Alloteropsis semialata*. *Journal of Experimental Botany* 59: 1743–1754.
- Pathare VS, Koteyeva N, Cousins AB. 2020. Increased adaxial stomatal density is associated with
 greater mesophyll surface area exposed to intercellular air spaces and mesophyll conductance in
 diverse C₄ grasses. *New Phytologist* 225: 169–182.
- Pearcy RW. 1977. Acclimation of photosynthetic and respiratory carbon dioxide exchange to growth
 temperature in *Atriplex lentiformis* (Torr.) Wats. *Plant Physiology* 59: 795–799.
- Peixoto M de M, Sage RF. 2017. Comparative photosynthetic responses in upland and lowland
 sugarcane cultivars grown in cool and warm conditions. *Brazilian Journal of Botany* 40: 829–839.
- Peixoto MM, Sage TL, Busch FA, Pacheco HDN, Moraes MG, Portes TA, Almeida RA, Graciano Ribeiro D, Sage RF. 2021. Elevated efficiency of C₃ photosynthesis in bamboo grasses: A possible
 consequence of enhanced refixation of photorespired CO₂. *GCB Bioenergy* 13: 941–954.
- Pengelly JJL, Sirault XRR, Tazoe Y, Evans JR, Furbank RT, von Caemmerer S. 2010. Growth of
 the C₄ dicot *Flaveria bidentis*: photosynthetic acclimation to low light through shifts in leaf anatomy and
 biochemistry. *Journal of Experimental Botany* 61: 4109–4122.

- Pignon CP, Long SP. 2020. Retrospective analysis of biochemical limitations to photosynthesis in 49
 species: C₄ crops appear still adapted to pre-industrial atmospheric [CO₂]. *Plant, Cell & Environment* 43: 2606–2622.
- Pinto H, Sharwood RE, Tissue DT, Ghannoum O. 2014. Photosynthesis of C₃, C₃-C₄, and C₄
 grasses at glacial CO₂. *Journal of Experimental Botany* 65: 3669–3681.
- Pinto H, Tissue DT, Ghannoum O. 2011. Panicum milioides (C₃-C₄) does not have improved water or
 nitrogen economies relative to C₃ and C₄ congeners exposed to industrial-age climate change. Journal
 of Experimental Botany 62: 3223–3234.
- 942 Pittermann J, Sage RF. 2001. The response of the high altitude C₄ grass *Muhlenbergia montana*943 (Nutt.) A.S. Hitchc. to long- and short-term chilling. *Journal of Experimental Botany* 52: 829–838.
- Poorter H, Fiorani F, Pieruschka R, Wojciechowski T, van der Putten WH, Kleyer M, Schurr U,
 Postma J. 2016. Pampered inside, pestered outside? Differences and similarities between plants
 growing in controlled conditions and in the field. *New Phytologist* 212: 838–855.
- Poorter H, Niinemets Ü, Poorter L, Wright IJ, Villar R. 2009. Causes and consequences of variation
 in leaf mass per area (LMA): a meta-analysis. *New Phytologist* 182: 565–588.
- 949 **R Core Team**. **2024**. A language and environment for statistical computing. In: R foundation for 950 statistical computing. https://www.r-project.org/: R v.4.4.1.
- Ripley BS, Abraham TI, Osborne CP. 2008. Consequences of C₄ photosynthesis for the partitioning
 of growth: a test using C₃ and C₄ subspecies of *Alloteropsis semialata* under nitrogen-limitation. *Journal* of *Experimental Botany* 59: 1705–1714.
- **Ripley BS, Cunniff J, Osborne CP**. **2013**. Photosynthetic acclimation and resource use by the C_3 and C_4 subspecies of *Alloteropsis semialata* in low CO_2 atmospheres. *Global Change Biology* **19**: 900–910.
- Ripley B, Frole K, Gilbert M. 2010. Differences in drought sensitivities and photosynthetic limitations
 between co-occurring C₃ and C₄ (NADP-ME) Panicoid grasses. *Annals of Botany* 105: 493–503.
- Ripley BS, Gilbert ME, Ibrahim DG, Osborne CP. 2007. Drought constraints on C₄ photosynthesis:
 stomatal and metabolic limitations in C₃ and C₄ subspecies of *Alloteropsis semialata*. *Journal of Experimental Botany* 58: 1351–1363.
- 961 **Sage RF**. **2002**. Variation in the k_{cat} of Rubisco in C₃ and C₄ plants and some implications for 962 photosynthetic performance at high and low temperature. *Journal of Experimental Botany* **53**: 609–620.
- 963 Sage RF. 2004. The evolution of C₄ photosynthesis. *New Phytologist* 161: 341–370.
- Sage RF. 2016. Tracking the evolutionary rise of C₄ metabolism. *Journal of Experimental Botany* 67:
 2919–2922.
- Sage RF. 2017. A portrait of the C₄ photosynthetic family on the 50th anniversary of its discovery:
 species number, evolutionary lineages, and Hall of Fame. *Journal of Experimental Botany* 68: e11–e28.
- Sage RF, Kubien DS. 2007. The temperature response of C₃ and C₄ photosynthesis. *Plant, Cell & Environment* 30: 1086–1106.

- Sage RF, Peixoto MM, Sage TL. 2013. Photosynthesis in Sugarcane. In: Sugarcane: Physiology,
 Biochemistry, and Functional Biology. 121–154.
- Sales CRG, Wang Y, Evers JB, Kromdijk J. 2021. Improving C₄ photosynthesis to increase
 productivity under optimal and suboptimal conditions. *Journal of Experimental Botany* 72: 5942–5960.
- 974 Salesse-Smith CE, Sharwood RE, Busch FA, Kromdijk J, Bardal V, Stern DB. 2018.
- 975 Overexpression of Rubisco subunits with RAF1 increases Rubisco content in maize. *Nature Plants* 4:
 976 802–810.
- 977 **Sharwood RE, Sonawane BV, Ghannoum O. 2014**. Photosynthetic flexibility in maize exposed to salinity and shade. *Journal of Experimental Botany* **65**: 3715–3724.
- Smith NG, Dukes JS. 2017. Short-term acclimation to warmer temperatures accelerates leaf carbon
 exchange processes across plant types. *Global Change Biology* 23: 4840–4853.
- Smith NG, Dukes JS. 2018. Drivers of leaf carbon exchange capacity across biomes at the continental
 scale. *Ecology* 99: 1610–1620.
- Smith NG, Keenan TF, Colin Prentice I, Wang H, Wright IJ, Niinemets Ü, Crous KY, Domingues
 TF, Guerrieri R, Yoko Ishida F, et al. 2019. Global photosynthetic capacity is optimized to the
 environment. *Ecology Letters* 22: 506–517.
- Soares AS, Driscoll SP, Olmos E, Harbinson J, Arrabaça MC, Foyer CH. 2008. Adaxial/abaxial
 specification in the regulation of photosynthesis and stomatal opening with respect to light orientation
 and growth with CO₂ enrichment in the C₄ species *Paspalum dilatatum*. *New Phytologist* **177**: 186–198.
- Soares-Cordeiro AS, Driscoll SP, Arrabaça MC, Foyer CH. 2011. Dorsoventral variations in dark
 chilling effects on photosynthesis and stomatal function in *Paspalum dilatatum* leaves. *Journal of Experimental Botany* 62: 687–699.
- Sonawane BV, Sharwood RE, von Caemmerer S, Whitney SM, Ghannoum O. 2017. Short-term
 thermal photosynthetic responses of C₄ grasses are independent of the biochemical subtype. *Journal of Experimental Botany* 68: 5583–5597.
- Sonawane BV, Sharwood RE, Whitney S, Ghannoum O. 2018. Shade compromises the
 photosynthetic efficiency of NADP-ME less than that of PEP-CK and NAD-ME C₄ grasses. *Journal of Experimental Botany* 69: 3053–3068.
- de Souza AP. 2011. Photosynthetic mechanisms and source-sink relationship in sugarcane grown in
 elevated CO₂, University of Sao Paulo.
- Still CJ, Berry JA, Collatz GJ, DeFries RS. 2003. Global distribution of C₃ and C₄ vegetation: carbon
 cycle implication. *Global Biogeochemical Cycles* 17: 1–14.
- Still CJ, Cotton JM, Griffith DM. 2019. Assessing earth system model predictions of C₄ grass cover in
 North America: From the glacial era to the end of this century. *Global Ecology and Biogeography* 28:
 145–157.
- Sun W, Ubierna N, Ma J-Y, Cousins AB. 2012. The influence of light quality on C₄ photosynthesis
 under steady-state conditions in *Zea mays* and *Miscanthus × giganteus*: changes in rates of

- photosynthesis but not the efficiency of the CO₂ concentrating mechanism. *Plant, Cell & Environment* **35**: 982–993.
- Taylor SH, Hulme SP, Rees M, Ripley BS, Ian Woodward F, Osborne CP. 2010. Ecophysiological
 traits in C₃ and C₄ grasses: a phylogenetically controlled screening experiment. *New Phytologist* 185:
 780–791.
- 1012 **Tissue DT, Griffin KL, Thomas RB, Strain BR**. **1995**. Effects of low and elevated CO_2 on C_3 and C_4 1013 annuals. *Oecologia* **101**: 21–28.
- 1014 Viechtbauer W. 2010. Conducting Meta-Analyses in R with the metafor Package. *Journal of Statistical* 1015 Software 36: 1–48.
- 1016Vogan PJ, Sage RF. 2012. Effects of low atmospheric CO_2 and elevated temperature during growth on1017the gas exchange responses of C_3 , C_3 -- C_4 intermediate, and C_4 species from three evolutionary1018lineages of C_4 photosynthesis. *Oecologia* 169: 341-352.
- Wang Y, Bräutigam A, Weber APM, Zhu X-G. 2014. Three distinct biochemical subtypes of C₄
 photosynthesis? A modelling analysis. *Journal of Experimental Botany* 65: 3567–3578.
- Wang F, Gao J, Yong JWH, Wang Q, Ma J, He X. 2020. Higher atmospheric CO₂ levels favor C₃
 plants over C₄ plants in utilizing ammonium as a nitrogen source. *Frontiers in Plant Science* 11:
 537443.
- Wang Z, Kang S, Jensen CR, Liu F. 2012. Alternate partial root-zone irrigation reduces bundle-sheath
 cell leakage to CO₂ and enhances photosynthetic capacity in maize leaves. *Journal of Experimental Botany* 63: 1145–1153.
- Wang D, Portis AR Jr, Moose SP, Long SP. 2008. Cool C₄ photosynthesis: Pyruvate P_i dikinase
 expression and activity corresponds to the exceptional cold tolerance of carbon assimilation in
 Miscanthus × giganteus. Plant Physiology 148: 557–567.
- 1030 **Ward DA**. **1987**. The temperature acclimation of photosynthetic responses to CO_2 in *Zea mays* and its 1031 relationship to the activities of photosynthetic enzymes and the CO_2 -concentrating mechanism of C_4 1032 photosynthesis. *Plant, Cell & Environment* **10**: 407–41.
- Wasilewska-Dębowska W, Zienkiewicz M, Drozak A. 2022. How light reactions of photosynthesis in
 C₄ plants are optimized and protected under high light conditions. *International Journal of Molecular Sciences* 23: 3626.
- Way DA, Yamori W. 2014. Thermal acclimation of photosynthesis: on the importance of adjusting our
 definitions and accounting for thermal acclimation of respiration. *Photosynthesis Research* 119: 89–
 100.
- 1039 Wickham H. 2016. ggplot2: Elegant Graphics for Data Analysis. New York: Springer-Verlag.
- 1040 **Wong SC**. **1979**. Elevated atmospheric partial pressure of CO_2 and plant growth. I. Interactions of 1041 nitrogen nutrition and photosynthetic capacity in C_3 and C_4 plants. *Oecologia* **44**: 68–74.

1042 Wu A, Truong SH, McCormick R, van Oosterom EJ, Messina CD, Cooper M, Hammer GL. 2024.

1043 Contrasting leaf-scale photosynthetic low-light response and its temperature dependency are key to 1044 differences in crop-scale radiation use efficiency. *New Phytologist* **241**: 2435–2447.

- 1045 Xu Z, Shimizu H, Ito S, Yagasaki Y, Zou C, Zhou G, Zheng Y. 2014. Effects of elevated CO₂,
- warming and precipitation change on plant growth, photosynthesis and peroxidation in dominant
 species from North China grassland. *Planta* 239: 421–435.
- **Yamori W, Hikosaka K, Way DA**. **2014**. Temperature response of photosynthesis in C₃, C₄, and CAM plants: temperature acclimation and temperature adaptation. *Photosynthesis Research* **119**: 101–117.

1052 Tables

Table 1. Results of multivariate linear mixed-effects models testing the response of V_{pmaxA} and A_{max} against fixed effects of speciesspecific traits and experimental conditions (see main text for more details). Effect size denotes the number of studies, while the sample size is the number of V_{pmaxA} or A_{max} data points. Model's marginal R^2 reflects the variance explained by fixed effects only, whereas conditional R^2 considers the variance explained by both fixed and random effects. Bold text indicates statistical significance. Significant codes: *, P < 0.05; **, P < 0.01; ***, P < 0.001. Continuous variables (i.e. mean T_{max} , growth CO₂, T_{leaf} and *PPFD*) were mean-centred before the analysis. SE = standard error. Lower and high CI: lower and upper 95% confidence intervals.

Fixed effects	V _{pmaxA}					A _{max}				
	Estimate	SE	Lower CI	Higher CI	P value	Estimate	SE	Lower CI	Higher CI	P value
C ₄ subtype	-1.299	7.716	-16.470	13.873	0.969	-1.339	3.363	-7.960	5.282	0.727
Growth form	0.270	8.272	-15.996	16.535	0.196	-15.091	11.082	-37.636	7.455	0.295
Growth location	-15.190	5.154	-25.747	-4.634	0.005**	-6.676	2.651	-12.295	-1.057	0.018*
Mean T _{max}	-2.549	1.362	-5.228	0.130	0.001**	0.367	0.754	-1.117	1.851	0.001**
Growth CO ₂	-0.018	0.012	-0.042	0.006	0.138	-0.003	0.009	-0.021	0.016	0.761
T_{leaf}	3.580	0.849	1.910	5.250	< 0.0001***	2.406	0.476	1.468	3.343	< 0.0001***
PPFD	0.014	0.002	0.010	0.018	< 0.0001***	0.010	0.001	0.008	0.013	< 0.0001***
C ₄ subtype × T_{leaf}	0.544	0.519	-0.477	1.565	0.504	0.187	0.261	-0.327	0.700	0.761
C_4 subtype × Mean T_{max}	-0.413	2.268	-4.873	4.047	0.840	-0.943	1.188	-3.283	1.396	0.642
C_4 subtype × <i>PPFD</i>	-0.001	0.005	-0.010	0.008	0.674	0.002	0.003	-0.003	0.007	0.750
Growth form \times <i>T</i> _{leaf}	-1.412	0.982	-3.342	0.518	0.146	3.590	2.519	-1.368	8.549	0.219
Growth form \times Mean T_{max}	3.477	2.132	-0.890	7.845	0.099	0.148	1.207	-2.410	2.707	0.781
Growth form \times <i>PPFD</i>	0.004	0.008	0.631	-0.012	0.645	-0.023	0.019	-0.066	0.007	0.143
$T_{\text{leaf}} \times \text{Mean } T_{\text{max}}$	0.013	0.036	-0.058	0.083	0.724	-0.043	0.020	-0.082	-0.004	0.033*
Species group	8.209	6.740	-5.385	21.802	0.230	3.467	2.381	-1.376	8.311	0.155
Overall model statistics	VpmaxA					Amax				
Effect size/sample size	49/401					37/297				

Marginal R^2	0.51	0.47
Conditional R ²	0.83	0.75



Figure 1. A summary of data points (by groups) based on (A) growth forms, (B) locations, (C) CO₂
treatments, (D) fertilisation treatments and (E) watering status. Decreased and elevated CO₂ treatments
consist of measurements made on plants with a growth CO₂ concentration of < 400 ppm and > 400 ppm,
respectively. See Materials and Methods for more information.



1068

Figure 2. Correlation of A_{max} versus V_{pmaxA} estimated from data collected from a range of species measured at various growth and measuring conditions. Point shapes reflect important species groups and colours denote measurement temperature (T_{leaf}). (A) All available A_{max} - V_{pmaxA} data; (B–D) data subsets measured at 25–30°C, 30–35°C, and 35–40°C, respectively. Equation, adjusted R^2 values and P values of individual fits are shown in each panel.



1075

1076 Figure 3. V_{pmaxA} and A_{max} plotted by C₄ biochemical subtypes (A and B, respectively), growth forms (C 1077 and D, respectively) and growth locations (E and F, respectively). Data presented here are a mixture of measurements done at various growth and measuring temperatures and irradiance. Raw data points are 1078 1079 plotted as coloured symbols, with different symbol shapes reflecting important species groups. Model-1080 predicted marginal means and intervals are shown as a horizontal line. On this line, black circles indicate model-predicted meta-analytic means of V_{pmaxA} or A_{max} , thick bars are 95% confidence intervals and thin 1081 1082 bars are 95% prediction intervals. On the right-hand side of each panel, the number of unique species per 1083 category (n.spp) and the number of individual studies (e.size) are indicated. P values are indicated in each panel (see Table 1). Significant codes: *, P < 0.05; **, P < 0.01; ***, P < 0.001. 1084 1085



Figure 4. Comparison between V_{pmaxA} and A_{max} (mean \pm SE) with growth location coloured by species group (C₄ non-model species versus model species). A linear mixed-effect model was conducted to examine V_{pmaxA} and A_{max} between two species groups at a location, or within a species group at both locations. Statistical results of comparisons are denoted with horizontal lines (i.e. the two bars at the beginning and the end of a horizontal line are compared) and asterisks indicate statistical significance (*, P < 0.05; **, P < 0.01; ***, P < 0.001).



1094

Figure 5. Relationships between V_{pmaxA} and A_{max} with mean T_{max} (A and B, respectively) and growth CO₂ levels (C and D, respectively). Data points in (A) and (B) are coloured in gradient by the measurement temperature (T_{leaf}), and the shapes of data points in all panels reflect the growth location (i.e. indoor vs. outdoor). Solid lines represent model-predicted values of V_{pmaxA} or A_{max} at a given mean T_{max} or CO₂ level (equations are shown at the top of each panel), dashed lines indicate 95% confidence intervals, and dotted lines show 95% prediction intervals. *P* values of the multivariate linear mixed-effects models are indicated. Significant codes: *, *P* < 0.05; **, *P* < 0.01; ***, *P* < 0.001.



1102

Figure 6. Relationships between V_{pmaxA} and A_{max} with T_{leaf} (A and B, respectively) and *PPFD* (C and D, respectively). Data points in (A) and (B) are coloured by the mean maximum growth temperature (mean T_{max}), and the shapes of data points in all panels reflect the growth location (i.e. indoor vs. outdoor). Solid lines represent model-predicted values of V_{pmaxA} or A_{max} at a given T_{leaf} or *PPFD* (equations are shown at the top of each panel), dashed lines indicate 95% confidence intervals, and dotted lines show 95% prediction intervals. *P* values of the multivariate linear mixed-effects models are indicated. Significant codes: *, *P* < 0.05; **, *P* < 0.01; ***, *P* < 0.001.



Figure 7. Contour plots illustrating the model-predicted responses of V_{pmaxA} and A_{max} (μ mol m⁻² s⁻¹) to T_{leaf} and mean T_{max} . *P* values of linear mixed-effects model testing the interaction between T_{leaf} and mean T_{max} are indicated (see Table 1). Black circle represent comparisons of V_{pmaxA} and A_{max} at the same mean T_{max} and T_{leaf} of 20, 30 and 38°C, with the colour gradient reflecting changes in V_{pmaxA} and A_{max} (see the colour legend on the right-hand side of each panel). Significant codes: *, *P* < 0.05; **, *P* < 0.01; ***, *P* < 0.001.

1117 Supporting Information

1118 Table S1. Results of multivariate linear mixed-effects models testing the response of V_{pmaxA} and A_{max}

1119 against species-specific traits and experimental conditions for data with $PPFD > 1000 \ \mu mol \ photon \ m^{-1}$

1120 2 s⁻¹. See Table 1 and main text for more details. Effect size denotes the number of studies, while the

1121 sample size is the number of V_{pmaxA} or A_{max} data points. Model's marginal R^2 reflects the variance 1122 explained by fixed factors only, whereas conditional R^2 considers the variance explained by both fixed 1123 and random factors. Bold text indicates statistical significance. Significant codes: *, P < 0.05; **, P <1124 0.01; ***, P < 0.001.

	V _{pmaxA}	A_{\max}			
Fixed effects					
C ₄ subtype	0.764	0.864			
Growth form	0.294	0.760			
Growth location	0.013*	0.030*			
Mean T _{max}	0.002**	0.003**			
Growth CO ₂	0.097	0.542			
$T_{ m leaf}$	< 0.0001***	< 0.0001***			
PPFD	0.119	0.061			
C ₄ subtype \times <i>T</i> _{leaf}	0.795	0.702			
C ₄ subtype × Mean T_{max}	0.960	0.744			
C ₄ subtype \times <i>PPFD</i>	0.704	0.795			
Growth form \times <i>T</i> _{leaf}	0.219	0.263			
Growth form \times Mean T_{max}	0.625	0.692			
Growth form \times <i>PPFD</i>	0.097	0.521			
$T_{\text{leaf}} \times \text{Mean } T_{\text{max}}$	0.278	0.002**			
Species group	0.179	0.162			
Overall model statistics					
Effect size/sample size	43/317	34/261			
Marginal R ²	0.50	0.41			
Conditional R ²	0.80	0.72			

1125



Figure S1. Species distribution within the categories of (A) growth forms and (B) growth locations. The top three measured species (maize, sorghum and *Setaria viridis*) are labelled explicitly, whereas all other species are shown as "Others". Detailed information on species is available in Supplementary Datasets S1 and S2.



1134Figure S2. Correlation of mean T_{max} and T_{min} (°C). The size of the dots reflects the number of A/C_i curve1135groups. See Supplementary Dataset S1 for more details.



1137

1138 Figure S3. Data distribution of V_{pmaxA} and A_{max} in categories of C₄ subtypes and growth forms over T_{leaf} , 1139 mean T_{max} and *PPFD*. Linear regression is fitted to each category within each panel, with equations, 1140 adjusted R^2 and P values being shown.



Figure S4. Histograms describing the data distribution across the spectrum of mean T_{max} and T_{leaf} for growth location (A & B), growth form (C & D) and important species group (E & F). Dash lines indicate the average values of mean T_{max} and T_{leaf} across plants grouped by different traits.



1147

1148 Figure S5. V_{pmaxA} and A_{max} plotted by C₄ biochemical subtypes (A and B, respectively). Data presented here are a mixture of measurements done at various growth and measuring temperatures and irradiance. 1149 1150 Raw data points are plotted as coloured symbols, with different symbol shapes reflecting lineages. Modelpredicted marginal means and intervals are shown as a horizontal line. On this line, black circles indicate 1151 1152 model-predicted meta-analytic means of V_{pmaxA} or A_{max} , thick bars are 95% confidence intervals and thin 1153 bars are 95% prediction intervals. On the right-hand side of each panel, the number of unique species per 1154 category (n.spp) and the number of individual studies (e.size) are indicated. P values of a multivariate 1155 mixed effect model indicating the effect of lineages on V_{pmaxA} and A_{max} are shown on each panel. 1156 Significant codes: *, *P* < 0.05; **, *P* < 0.01; ***, *P* < 0.001.



Figure S6. Relationships between V_{pmaxA} and A_{max} with mean T_{max} (A and B, respectively) for measurements done at T_{leaf} between 25 and 30°C. Data presented here are a mixture of measurements done at various *PPFD*. Data points are coloured in gradient by T_{leaf} , the shapes of data points in all panels reflect the growth location (i.e. indoor vs. outdoor). Solid lines represent model-predicted values of V_{pmaxA} or A_{max} at a given T_{leaf} or *PPFD* (equations are shown at the top of each panel), dashed lines indicate 95% confidence intervals, and dotted lines show 95% prediction intervals. *P* values of the multivariate linear mixed-effects models are indicated. Significant codes: *, P < 0.05; **, P < 0.01; ***, P < 0.001.



Figure S7. Correlation of A_{max} versus V_{pmaxA} estimated from data with $PPFD > 1,500 \ \mu\text{mol}$ photon m⁻² s⁻ ¹, coloured by $T_{\text{leaf.}}$