See discussions, stats, and author profiles for this publication at: https://www.researchgate.net/publication/273169362

Optimal stomatal behaviour around the world

Article in Nature Climate Change · March 2015 DOI: 10.1038/nclimate2550 CITATIONS READS 52 1,456 54 authors, including: Yan-Shih Lin Belinda E Medlyn French National Institute for Agricultural Res... Western Sydney University 17 PUBLICATIONS 335 CITATIONS 121 PUBLICATIONS 7,601 CITATIONS SEE PROFILE SEE PROFILE Han Wang **Remko Duursma** Western Sydney University Northwest A & F University 88 PUBLICATIONS 2,416 CITATIONS 32 PUBLICATIONS 398 CITATIONS SEE PROFILE SEE PROFILE

Some of the authors of this publication are also working on these related projects:



Do Plants Have Memory of Mechanical Stress - Is It an Epigenetic Phenomenon? View project



Plant drought responses in biodiverse ecosystems View project

All content following this page was uploaded by Derek Eamus on 04 August 2015.

1 Published in Nature Climate Change 5, 459 – 464 (2015)

Optimal stomatal behaviour around the world: synthesis of a global stomatal conductance database

4

Yan-Shih Lin¹, Belinda E. Medlyn¹, Remko A. Duursma², I. Colin Prentice^{1,3}, Owen K. 5 Atkin⁴, Craig V.M. Barton², Jonathan Bennie⁵, Alexandre Bosc^{6,7}, Mark S.J. 6 Broadmeadow⁸, Lucas A. Cernusak⁹, Paolo De Angelis¹⁰, John E. Drake², Derek Eamus¹¹, 7 David S. Ellsworth², Michael Freeman¹², Oula Ghannoum², Teresa E. Gimeno², Qingmin 8 Han¹³, Kouki Hikosaka¹⁴, Lindsay B. Hutley¹⁵, Jeff W. Kelly¹, Kihachiro Kikuzawa¹⁶, Pasi 9 Kolari¹⁷, Kohei Koyama^{16,18}, Jean-Marc Limousin¹⁹, Maj-Lena Linderson²⁰, Markus Löw²¹, 10 Cate Macinins-Ng²², Nicolas K. Martin-StPaul²³, Patrick Meir²⁴, Teis N. Mikkelsen²⁵, 11 Patrick Mitchell²⁶, Jesse B. Nippert²⁷, Yusuke Onoda²⁸, Maarten Op de Beeck²⁹, Victor 12 Resco de Dios³⁰, Ana Rey³¹, Alistair Rogers³², Lucy Rowland²⁴, Samantha A. Setterfield¹⁵, 13 Wei Sun³³, Lasse Tarvainen³⁴, Sabine Tausz-Posch²¹, David T. Tissue², Johan Uddling³⁵, 14 Göran Wallin³⁵, Jeff M. Warren³⁶, Lisa Wingate⁶, Joana Zaragoza-Castells²⁴ 15

16

¹: Department of Biological Sciences, Macquarie University, North Ryde, NSW 2109,
Australia

- ²: Hawkesbury Institute for the Environment, University of Western Sydney, Penrith, New
 South Wales 2751, Australia
- ³: Grantham Institute and Division of Ecology and Evolution, Imperial College, Silwood
 Park Campus, Ascot SL5 7PY, United Kingdom
- ⁴: Division of Plant Sciences, Research School of Biology, The Australian National
 University, Canberra, Australian Capital Territory 0200, Australia
- ⁵: Environment and Sustainability Institute, University of Exeter, Penryn, United Kingdom

- ⁶: Institut National de la Recherche Agronomique, Villenave d'Ornon F-33140, France
- ⁷: Bordeaux Sciences Agro, UMR 1391 ISPA, Gradignan F-33170, France
- 28 ⁸: Climate Change Forest Services, Forestry Commission England, United Kingdom
- ⁹: James Cook University, Cairns, Queensland 4879, Australia
- ¹⁰: Department for Innovation in Biological, Agro-food and Forest systems, University of
- 31 Tuscia, Via San Camillo de Lellis, Viterbo 01100, Italy
- ¹¹: School of Life Sciences, University of Technology, Sydney, New South Wales 2007,
 Australia
- ¹²: Department of Ecology, Swedish University of Agricultural Sciences, UPPSALA 75007,
- 35 Sweden
- 36 ¹³: Hokkaido Research Center, Forestry and Forest Products Research Institute (FFPRI),
- 37 Toyohira, Sapporo, Hokkaido 062-8516, Japan
- ¹⁴: Graduate School of Life Sciences, Tohoku University, Aoba, Sendai 980-8578, Japan
- 39 ¹⁵: Research Institute for Environment and Livelihoods, Charles Darwin University,
- 40 Casuarina, Northern Territory 0810, Australia
- 41 ¹⁶: Department of Environmental Science, Faculty of Bioresources and Environmental
- 42 Sciences, Ishikawa Prefectural University, Ishikawa 921-8836, Japan
- 43 ¹⁷: Department of Physics, University of Helsinki, Finland
- 44 ¹⁸: Department of Life Science and Agriculture, Obihiro University of Agriculture and
- 45 Veterinary Medicine, Obihiro, Hokkaido 080-0834, Japan
- 46 ¹⁹: Department of Biology, University of New Mexico, Albuquerque, NM 87131-0001,

47 United States

- 48 ²⁰: Department of Physical Geography and Ecosystem Science, Lund University, Sweden
- 49 ²¹: Department of Agriculture and Food Systems, University of Melbourne, Creswick,
- 50 Victoria 3363, Australia

- ²²: School of Environment, Unversity of Auckland, Auckland 1142, New Zealand
- ²³: Université Paris-Sud, Laboratoire Ecologie, Systématique et Evolution, UMR8079,
 Orsay F-91405, France
- ²⁴: School of Geosciences, The University of Edinburgh, Edinburgh EH8 9XP, United
 Kingdom
- ²⁵: Center for Ecosystems and Environmental Sustainability, Department of Chemical and
- 57 Biochemical engineering, Technical University of Denmark, DK-4000 Roskilde, Denmark
- ²⁶: CSIRO Ecosystem Sciences, Sandy Bay, Tasmania 7005, Australia
- ²⁷: Division of Biology, Kansas State University, Manhattan, KS 66505, United States
- 60 ²⁸: Division of Environmental Science and Technology, Graduate School of Agriculture,
- 61 Kyoto University, Oiwake, Kitashirakawa, Kyoto 606-8502, Japan
- ²⁹: Research Group Plant and Vegetation Ecology, University of Antwerp, Wilrijk 2610,
 Belgium
- ³⁰: Producció Vegetal i Ciència Forestal, Universitat de Lleida, Lleida 25198, Spain
- ⁶⁵ ³¹: Department of Biogeography and Global Change, MNCN-CSIC, Spanish Scientific
- 66 Council, Madrid 28006, Spain
- 67 ³²: Environmental and Climate Sciences Department, Brookhaven National Laboratory,
- 68 Upton, NY 11973-5000, United States
- 69 ³³: Institute of Grassland Science, Northeast Normal University, Key Laboratory of
- 70 Vegetation Ecology, Changchun, Jilin 130024, China
- ³⁴: Department of Forest Ecology and Management, Swedish University of Agricultural
 Sciences, Umeå 90183, Sweden
- 73 ³⁵: Department of Biological and Environmental Sciences, University of Gothenburg,
- 74 Göteborg 40530, Sweden

³⁶: Environmental Sciences Division, Oak Ridge National Laboratory, Oak Ridge, TN,
USA

77 Main text

78 Stomatal conductance is a key land surface attribute as it links plant water-use and carbon 79 uptake. In this study we synthesised a globally distributed database of stomatal 80 conductance data sets obtained in the field for a wide range of plant functional types (PFTs) and biomes. We employed a model of optimal stomatal conductance¹ to assess differences 81 in stomatal behaviour. We estimated the model slope coefficient, g_1 , which is directly 82 83 related to the marginal carbon cost of water-use, for each dataset. We then tested how g_1 84 varies with climatic factors, including temperature and water availability, and across PFTs. We found that g_1 varied considerably among PFTs, with every savanna trees having 85 the largest g_1 (least conservative water-use), followed by C_3 grasses and crops, angiosperm 86 trees, gymnosperm trees, and C₄ grasses. Amongst angiosperm trees, species with larger 87 88 wood density had a larger marginal carbon cost of water-use, as predicted by the theory underpinning the optimal stomatal model. There was an interactive effect between 89 temperature and moisture availability (on g_1 : for wet environments, g_1 was largest in high 90 91 temperature environments, indicated by high mean annual growing degree days above 0° C (mGDD₀), but it did not vary with mGDD₀ across dry environments. These findings 92 93 provide a robust theoretical framework for understanding and predicting the behaviour of 94 stomatal conductance across biomes and across PFTs that can be applied to regional, continental and global-scale modelling of productivity and ecohydrological processes in a 95 96 future changing climate.

97

98 Earth System Models (ESMs) integrate biogeochemical and biogeophysical land surface
99 processes with physical climate models and have been widely used to demonstrate the

importance of land surface processes in determining climate and to highlight the issue of 100 large uncertainties in quatifying land surface processes^{2, 3, 4, 5}. Within the biogeophysical 101 components of land surface processes, stomatal conductance plays a pivotal role because it 102 103 is a key feedback route for carbon and water exchange between the atmosphere and terrestial vegetation. Stomata are small pores on leaves whose behaviour can be regulated 104 105 by the plant in response to multiple abiotic and biotic factors. Stomatal conductance (g_s) is a major determinant of both transpiration rates and rates of photosynthetic C uptake. . 106 107 Therefore, our ability to model the global carbon and water cycles under future changing climate depends on our ability to predict stomatal behaviour globally¹, an ability that to-108 109 date has remained particularly intractactable . Although there have been previous synthesis studies on plant stomatal conductance and related traits^{6, 7, 8, 9}, a global scale database and 110 associated mechanistic globally applicable model of g_s that would allow prediction of 111 112 stomatal behaviour is lacking.

113

114 For this study, we compiled a unique global database of field measurements of stomatal conductance and photosynthesis suitable for extracting model parameters. We employed a 115 model of optimal stomatal conductance¹ to develop hypotheses for how stomatal behaviour 116 should vary with environmental factors and with plant traits associated with hydraulic 117 function. In the optimal stomatal model, the slope parameter, g_1 , is proportional to the 118 marginal carbon cost of water-use¹, meaning that plants with smaller g_1 values are more 119 conservative with their water-use and have higher water-use-efficiency (and vice versa). 120 Therefore, we hypothesised that variation in g_1 values among climate zones and PFTs 121 122 should reflect differences in the cost of water transport. We proposed that:

123 (1) g_1 values among PFTs should vary according to the cost of stemwood construction, 124 such that C3 herbaceous species should have the largest g_1 (i.e. least conservative wateruse), followed by angiosperm trees and gymnosperm trees. Since the optimal stomatal theory predicts that, for the same marginal water cost, g_1 should be lower by approximately one-half ¹⁰. We therefore predicted that C4 plants would have the smallest g_1 .

128 (2) For trees, the cost of water transport should increase with wood density, due to the 129 higher cost of wood construction¹¹ and the generally smaller hydraulic conductance of 130 sapwoos with large density. Therefore within both angiosperms and gymnosprems, trees 131 with highest wood density should have the smallest g_1 .

132 (3) Moisture stress should increase the cost of water-use to the plant, so plants in dry 133 environments should have a larger marginal cost of water-use and lower g_1 .

134 (4) g_1 values should increase with temperature for two reasons. First, we previously 135 showed that g_1 is approximately proportional to a combination term of the carbon cost of 136 water transport and Γ^* (the CO₂ compensation point in absence of photorespiration)¹. As 137 Γ^* is exponentially dependent on temperature^{1, 12}, g_1 should similarly increase with 138 temperature. Second, the viscosity of water decreases with increasing temperature, making 139 it less costly to transport water leading to a increased g_1^{13} .

140

141 To test these hypotheses, we collated a globally distributed database of g_s and 142 photosynthesis of 56 field studies, covering a wide range of biomes from Arctic tundra, 143 boreal and temperate forest to tropical rainforest (Table S1). We estimated the model coefficient, g_1 , from observations of leaf-level gas exchange (g_s , ratesd of transpiration 144 145 and net photosynthesis, see Methods) and environmental drivers. We used mean annual 146 degree days above 0°C (mGDD₀) and moisture index (MI) derived from observed longterm meteorological data as proxies to quantify the temperature and water availability that 147 are relevant to plant physiological functions for each site¹⁴. The growing degree days 148 above 0°C is an index of the energy available for completion of the annual life cycle and 149

quantifies temperature limitations to carbon assimilation and growth^{15, 16}. Our database covered a range of mGDD₀ from 2.7 to 29.7 °C and a range of MI from 0.17 to 3.26, representing the majority of the climatic space for vegetation covered land surfaces (Fig. 1). We then tested how g_1 varies with MI and mGDD₀ across PFTs and biomes?.

154

155 We found a clear pattern of g_1 variation among different PFTs with evergreen savanna trees having largest g_1 , followed by C_3 grasses and crops, angiosperm trees, gymnosperm 156 157 trees, and C_4 grasses (Table S2 and Fig. 2). For angiosperm trees, g_1 was negatively 158 correlated with wood density, although we did not find any correlation for gymnosperm species (Fig. 3). g_1 significantly increased with both increasing mGDD₀ and MI across the 159 160 entire data set. However, when evaluated as a bivariate relationship (Fig. 2c-d, and Fig. 4a-161 b) we observed that there was an interactive effect between temperature and moisture availability on g_1 : for wet environments, g_1 was largest at sites with high mGDD₀, but it 162 163 varied with mGDD₀ to a much smaller degree across dry environments (Table 1 and Fig. 164 4).

Our results largely supported our hypotheses for how g_1 should vary among PFTs 165 166 (hypothesis 1) and biomes. The variation in g_1 among PFTs is a result of trade-offs among plant functions such as growth, defence and reproduction, through different resource 167 allocation patterns that aim to achieve the optimal cost-to-benefit ratios^{8, 13} Long life-span 168 169 PFTs, such as evergreen gymnosperm and angiosperm trees, must invest more in building supporting and defence structures relative to short life-span PFTs, such as grasses, so that 170 171 they can be sustained over many years of biotic and abiotic stress. Such an investment preference has to come at the cost of reduced growth rates^{17, 18}, meaning reduced the rates 172 of carbon uptake and water loss cost through opening stomata. Therefore we predicted a 173 more conservative water-use strategy in trees (lower g_1) than in C3 grass (higher g_1), and 174

175 this was observed in the database. However, evergreen savanna trees formed an exception 176 with a surprisingly large g_1 , relative to expectations based upon trees wood density and biomes MI. This may result from the fact that these species have several unique hydraulic 177 178 functional traits that may offset the carbon cost of water-use which allow them to have a less conservative water use strategy. These hydraulic functional traits include: deep roots 179 180 to access groundwater, large sapwood area for water transport, narrow but long conduits to reduce the risk of embolism and reduce the cost of conduit wall construction^{19, 20} and drv 181 season declines in LAI to balance increased atmospheric aridity in the dry season. This 182 183 special case of evergreen savanna trees is worthy of further investigation.

184

185 We found a significant relationship between g_1 and wood density among angiosperm trees 186 (Fig. 3; excluding savanna angiosperms) which supported our hypothesis that g_1 is negatively correlated with wood density (hypothesis 2). A larger wood density is 187 188 advantagous for plants that need to avoid hydraulic failure so that they can sustain more negative sapwood water pressures during drought¹⁸. However, such an investment is at the 189 expense of a reduced capacity for stem water storage, reduced sapwood conductivity and 190 the carbon cost of building wood with higher density^{20, 21, 22}, and thus leads to a more 191 conservative water-use-strategy. However, we did not find such a relationship among 192 gymnosperm trees. This lack of correlation may be due to the limited variability in wood 193 194 density in gymnosperms. There are significant differences in the anatomical structure of sapwood between angiosperms and gymnosperms. The majority of angiosperm trees have 195 evolved to separate the water transport structure (i.e. vessels) from the mechanical support 196 structure, while gymnosperm trees do not have such a functional differentiation, as 197 tracheids are used for both water transport and mechanical support^{18, 23}. Therefore, wood 198 density is a good proxy for quantifying the trade-offs between transport and support 199

investments for angiosperm trees but not for gymnosperm trees²³. The distinct differences 200 201 in the water-use strategy between angiosperm trees and gymnosperm trees (Fig. 2) is consistent with a recent observation that angiosperms maintain a much smaller hydraulic 202 safety margin than gymnosperms²⁴, showing that angiosperms allow some loss of 203 hydraulic conductivity - a risky strategy - while gymnosperms minimise lossThis 204 205 evolutionary development confers an advantage to angiosperm trees by allowing them to 206 use water in a less conservative way, thereby increasing their carbon gain relative to 207 gymnosperm trees.

208

Our results only partially supported our hypotheses for how g_1 should vary with moisture 209 210 stress and temperature (hypotheses 3 and 4 as there was an interactive effect between 211 temperature and moisture stress on g_1 . This interactive response between MI and mGDD₀ 212 demonstrates the complexity of how plants co-ordinate their resource allocation strategies 213 along two axes of climatic gradient (Fig. 4). Temperature affects the cost of water transport 214 in such a way that it should be more costly to transport water in a colder environment than 215 in a warmer one. However, lower temperature also comes with water savings as the 216 evaporative demand and photorespiratory cost are lower. The interactive relationship 217 between MI and mGDD₀ suggest that the rate of change in g_1 (i.e. the slope of each 218 exponential curve; Fig. S3) along temperature or water availability gradient is much higher 219 in the wet and warm environments than in dry and cold environments.

220

Our study demonstrated the first mechanistically robust framework that can be applied to various scales for understanding and predicting the behaviour of stomatal conductance across biomes and across PFTs. We analysed a global stomatal behaviour data set along two major climatic axes, providing an analytic framework for understanding how

stomatal behaviour adapts to the environment. Our findings will allow the ESM community to move on from using empirical stomatal models (ref ref) with tuned parameters to using a more robust, theory-derived optimal stomatal model with meaningful parameters. In addition, we provide a valuable stomatal behaviour database that can be used to parameterise g_s among PFTs and which can be applied directly within ESMs for modelling productivity and ecohydrological processes in a future changing climate across regional, continental and global scales.

232

234 Methods

235 Source of data

We synthesised published and unpublished leaf gas exchange data sets for a wide range of 236 237 PFTs and biomes (Table S1). Our database covers 314 species from 56 experiment sites 238 around the world with 17 sites from Australasia, 15 sites from Europe, 14 sites from North 239 America, six sites from Asia, three sites from South America and one site from Africa. Site latitudes range from 42.9°S to 72.3°N although the majority of the sites are within the 240 temperate zone (n=35; latitude range between 23.5° to 55° and between -23.5° and -55°), 241 followed by tropical zone (n=14; latitude range between -23.5° and 23.5°), boreal zone 242 (n=6; latitude range between 55° and 66.5°) and Arctic zone (n=1; latitude range above 243 66.5°). We used MI and mGDD₀ derived from Climate Research Unit data (CRU TS3.1)²⁵ 244 from 1991 to 2010 using a modified version of the STASH model²⁶ at a grid resolution of 245 0.5° . In this derivation, mGDD₀ was calculated as the ratio of the annual sum of 246 247 temperatures above 0°C (growing degree days) to the length of the period with temperatures above 0°C; MI was calculated as the ratio of mean annual precipitation to the 248 249 equilibrium evapo-transpiration (E_{eq}). We estimated E_{eq} from temperature and net radiation 250 (calculated from monthly mean percentage of cloud cover) based on the Priestley-Taylor equation²⁶. The Sea-WiFS fAPAR (fraction absorbed photosynthetically active radiation) 251 product was used to determine areas with green vegetation cover at a grid resolution of 0.5° . 252 The wood density data were obtained from the Global Wood Density Database^{23, 27}. 253

254

255 Data analysis

We used data points measured at a photosynthetic photon flux density (PPFD) > 0 μ mol m⁻² s⁻¹, and only data collected from the top third of the canopy (what would happen if you used data for PAR> 250 μ mol m⁻² s⁻¹ rather than > 0? . Data points with negative photosynthesis rates were excluded. In all cases, species were grown under ambient environmental conditions and were not subjected to any treatments, such as elevated CO_2 , temperature, or drought treatments. We employed an optimal stomatal model¹ as:

$$g_s = g_0 + 1.6 \times (1 + \frac{g_1}{\sqrt{D}}) \frac{A}{C_a}$$

where D is vapour pressure deficit, A is net photosynthesis rate, C_a is CO₂ concentration at 262 leaf surface, and g_0 , g_1 are model coefficients for intercept and slope. We used a non-linear 263 mixed-effect model to estimate the model slope coefficient, g_1 , for each group separately 264 265 for various classification schemes as shown in Fig. 2. In all g_1 estimations, we assumed the 266 intercept coefficient, g_0 , to be zero to avoid strong correlation between g_0 and g_1 which 267 would mask any interesting variation in g_1 . In this model, individual species were assumed 268 to be the random effect to account for the differences in the g_1 slope among species within 269 the same group. To test how g_1 varies with climatic variables (i.e. MI and mGDD₀), we first estimated g_1 for each species using non-linear regression. We then used a linear 270 mixed-effect model to test the relationship between g_1 , MI and mGDD₀. We fitted the 271 272 model as:

$$\log(g_1) \sim MI + mGDD_0 + MI \times mGDD_0$$

assuming PFTs as the random effect to account for the differences in intercept among PFTs. To evaluate the goodness of fit for linear mix-effect model, we calculated both the marginal R^2 to quantify the proportion of variance explained by the fixed factors alone and the conditional R^2 to quantify the proportion of variance explained by both the fixed and random factors as described in Nakagawa and Holger Schielzeth (2013)²⁸. The relationship between g_1 and wood density were tested with a simple linear regression model. All model estimations and statistical analyses were performed within R 3.1.0²⁹.

280	References				
281 282	1.	Medlyn BE, <i>et al.</i> Reconciling the optimal and empirical approaches to modelling stomatal conductance. <i>Global Change Biology</i> 17 , 2134-2144 (2011).			
283 284 285	2.	Cox PM, Betts RA, Jones CD, Spall SA, Totterdell IJ. Acceleration of global warming due to carbon-cycle feedbacks in a coupled climate model. <i>Nature</i> 408 , 184-187 (2000).			
286 287 288 289	3.	Sitch S, <i>et al</i> . Evaluation of ecosystem dynamics, plant geography and terrestrial carbon cycling in the LPJ dynamic global vegetation model. <i>Global Change Biology</i> 9 , 161-185 (2003).			
290 291 292	4.	Cao M, Woodward FI. Dynamic responses of terrestrial ecosystem carbon cycling to global climate change. <i>Nature</i> 393 , 249-252 (1998).			
293 294 295	5.	Friedlingstein P, <i>et al.</i> Climate-carbon cycle feedback analysis: Results from the C4MIP model intercomparison. <i>Journal of Climate</i> 19 , 3337-3353 (2006).			
296 297 298 299 300	6.	Schulze E-D, Kelliher FM, Korner C, Lloyd J, Leuning R. Relationships among maximum stomatal conductance, ecosystem surface conductance, carbon assimilation rate, and plant nitrogen nutrition: a global ecology scaling exercise. <i>Annual Review of Ecology and Systematics</i> , 629-660 (1994).			
301 302 303	7.	Kattge J <i>, et al.</i> TRY – a global database of plant traits. <i>Global Change Biology</i> 17 , 2905- 2935 (2011).			
304 305 306 307	8.	Wright IJ, Falster DS, Pickup M, Westoby M. Cross-species patterns in the coordination between leaf and stem traits, and their implications for plant hydraulics. <i>Physiologia Plantarum</i> 127 , 445-456 (2006).			
308 309 310	9.	Lloyd J, Farquhar G. 13C discrimination during CO2 assimilation by the terrestrial biosphere. <i>Oecologia</i> 99 , 201-215 (1994).			
311 312 313 314	10.	Way DA, Katul GG, Manzoni S, Vico G. Increasing water use efficiency along the C3 to C4 evolutionary pathway: a stomatal optimization perspective. <i>Journal of Experimental Botany</i> , (2014).			
315 316 317 318	11.	Héroult A, Lin Y-S, Bourne A, Medlyn BE, Ellsworth DS. Optimal stomatal conductance in relation to photosynthesis in climatically contrasting Eucalyptus species under drought. <i>Plant, Cell & Environment</i> 36 , 262-274 (2013).			
319					

- Medlyn BE, *et al.* Temperature response of parameters of a biochemically based model of
 photosynthesis. II. A review of experimental data. *Plant Cell and Environment* 25, 1167 1179 (2002).
- Prentice IC, Dong N, Gleason SM, Maire V, Wright IJ. Balancing the costs of carbon gain and water transport: Testing a new theoretical framework for plant functional ecology. *Ecology Letters* 17, 82-91 (2014).

323

327

331

333

336

339

343

349

353

- Harrison SP, Prentice IC, Barboni D, Kohfeld KE, Ni J, Sutra JP. Ecophysiological and
 bioclimatic foundations for a global plant functional classification. *Journal of Vegetation Science* 21, 300-317 (2010).
- 332 15. Woodward FI. *Climate and Plant Distribution* Cambridge University Press (1987).
- 33416.Colin Prentice I, Sykes MT, Cramer W. A simulation model for the transient effects of335climate change on forest landscapes. *Ecological Modelling* **65**, 51-70 (1993).
- 33717.Enquist BJ, West GB, Charnov EL, Brown JH. Allometric scaling of production and life-338history variation in vascular plants. *Nature* **401**, 907-911 (1999).
- Hacke UG, Sperry JS, Pockman WT, Davis SD, McCulloh KA. Trends in wood density and
 structure are linked to prevention of xylem implosion by negative pressure. *Oecologia* 126, 457-461 (2001).
- 34419.Eamus D, O'Grady AP, Hutley L. Dry season conditions determine wet season water use in345the wet-tropical savannas of northern Australia. *Tree Physiology* **20**, 1219-1226 (2000).
- 346
 347 20. Sperry JS, Meinzer FC, McCulloh KA. Safety and efficiency conflicts in hydraulic
 348 architecture: Scaling from tissues to trees. *Plant, Cell and Environment* **31**, 632-645 (2008).
- Meinzer FC, James SA, Goldstein G, Woodruff D. Whole-tree water transport scales with
 sapwood capacitance in tropical forest canopy trees. *Plant, Cell and Environment* 26,
 1147-1155 (2003).
- Bucci SJ, Goldstein G, Meinzer FC, Scholz FG, Franco AC, Bustamante M. Functional
 convergence in hydraulic architecture and water relations of tropical savanna trees: From
 leaf to whole plant. *Tree Physiology* 24, 891-899 (2004).
- Chave J, Coomes D, Jansen S, Lewis SL, Swenson NG, Zanne AE. Towards a worldwide
 wood economics spectrum. *Ecology Letters* 12, 351-366 (2009).
- 360
 361 24. Choat B, *et al.* Global convergence in the vulnerability of forests to drought. *Nature* 491, 752-755 (2012).

363 364 365 366	25.	Harris I, Jones PD, Osborn TJ, Lister DH. Updated high-resolution grids of monthly climatic observations - the CRU TS3.10 Dataset. <i>International Journal of Climatology</i> 34 , 623-642 (2014).
367 368 369	26.	Gallego-Sala A <i>, et al.</i> Bioclimatic envelope model of climate change impacts on blanket peatland distribution in Great Britain. <i>Climate Research</i> 45 , 151-162 (2010).
370 371 372	27.	Zanne AE <i>, et al.</i> Data from: Towards a worldwide wood economics spectrum. Dryad Data Repository (2009).
373 374 375	28.	Nakagawa S, Schielzeth H. A general and simple method for obtaining R2 from generalized linear mixed-effects models. <i>Methods in Ecology and Evolution</i> 4 , 133-142 (2013).
376 377 378	29.	R Core Team. R: A language and environment for statistical computing. R Foundation for Statistical Computing (2014).
379		
380		
381		

382 Acknowledgements

383	This research was supported by the Australian Research Council (ARC MIA Discovery
384	Project 1433500-2012-14). A.R. was financially supported in part by The Next-Generation
385	Ecosystem Experiments (NGEE-Arctic) project that is supported by the Office of
386	Biological and Environmental Research in the Department of Energy, Office of Science,
387	and through the United States Department of Energy contract No. DE-AC02-98CH10886
388	to Brookhaven National Laboratory. M.O.d.B. acknowledges that the Brassica data were
389	obtained within a research project financed by the Belgian Science Policy (OFFQ, contract
390	number SD/AF/02) and coordinated by Dr Karine Vandermeiren at the Open-Top
391	Chamber research facilities of CODA-CERVA (Tervuren, Belgium).
392	
393	Author contributions
394	
395	
396	
397	Competing financial interests

398 The author declear no competing financial interests.

Table 1: Analysis of Variance table for g_1 as a function of MI and mGD
--

Model					
Variables	numDF	denDF	F-value	p-value	Marginal R ²
Intercept	1	97	67.08 <	0.001	0.20
MI	1	97	7.50	0.007	Conditional R ²
mGDD ₀	1	97	11.15	0.001	0.59
MI*mGDD ₀	1	97	1.34	0.250	

401 Figure legends

Figure 1: Climatic space covered by the Stomatal Behaviour Synthesis Database, shown as mean annual degree days above 0°C (mGDD₀; °C) and moisure index (MI). Coloured circles represent climatic space for the database, with different colours indicating different plant functional types. Grey hexagons represent global climatic space for which vegetation is present. The global climatic space data were binned by every 1 °C for mGDD0 and every 0.25 for MI.

408

Figure 2: Mean g_1 values for plant functional types defined by different classification schemes. Each bar represents mean ± SE. Panels (b) (c) and (d) include C₃ species data only.

411

Figure 3: Relationship between g_1 and wood density for angiosperm and gymnosperm 412 trees. Savanna tree species (all angiosperms) are indicated separately. Each data point 413 414 represents mean \pm SE of g_1 for individual species fitted with non-linear regression. A linear regression line was only fitted for angiosperm trees due to limited data for gymnosperm trees. 415 The fitted linear regression relationship between g_1 and wood density for angiosperm trees is: 416 $g_1 = -4.77 \text{*WD} + 6.96 (P = 0.0008, R^2 = 0.23)$. Wood density data were obtained from Global 417 Wood Density Database^{23, 27} and are available for 45 species in the Stomatal Behaviour Synthesis 418 419 Database.

Figure 4: Estimated and predicted g_1 as a function of mGDD₀ and MI. Panels (a) (b) show the relationship between estimated g_1 and (a) mean annual degree days above 0 °C temperature (mGDD₀; °C) and (b) moisture index (MI) at experimental sites among species across different plant functional types (PFTs). Each data point represents mean ± SE of g_1 for individual species

fitted with a non-linear regression. Classification of plant functional types are shown in Figure 2e. Panels (c) and (d) are the predicted g_1 under different ranges of MI and mGDD₀ presented as a partial regression plot. Predictions in (c) and (d) are from linear mixed-effects model for log(g_1) assuming PFTs as a random effect to account for the differences in intercept among PFTs. Colour lines represent the predicted g_1 based on fitted model coefficients (Table S3). Colour dots represent the partial regression predictions at a given fixed MI or mGDD₀ level.

430 Supplementary Materials

Table S1: List of data source.

Data contributor	Location	Species	Reference
Alexandre Bosc	Le Bray, France	Pinus pinaster	Bosc, A. (1999) PhD Thesis.
Alistair Rogers	Barrow, AK, USA	Several Arctic species	Unpublished data.
Ana Rey	Glencorse near Edinburgh, Scotland, UK	Betula pendula	Rey & Jarvis (1998) Tree Physiology.
Belinda Medlyn	Tumbarumba flux tower, Snowy Mts, NSW, Australia	Eucalyptus delegatensis	Medlyn et al. (2007) Tree Physiology.
Cate Macinnis-Ng	Arataki Visitor Centre, Auckland, New Zealand	Agathis australis	Unpublished data
Craig Barton	Glencorse near Edinburgh Scotland	Picea sitchensis	Barton & Jarvis (1999) New Phytologist.
David Ellsworth	Duke Forest, Durham, NC, USA	Pinus taeda	Ellsworth DS (1999) Plant, Cell & Environment.
David Ellsworth	Richmond, Sydney, Australia	Eucalyptus saligna	Unpublished data
David Ellsworth	Richmond, Sydney, Australia	Four Eucalyptus species	Héroult et al. (2013) Plant, Cell & Environment.
David Tissue	Big Bend National Park, Texas, USA	Larrea tridentata	Ogle et al. (2012)
Derek Eamus	Palmerston, NT, Australia	A set of six savanna tree species	Thomas & Eamus (2002) Australian Journal of Botany.
Derek Eamus	Western Sydney, Castlereagh, Australia	Angophora bakeri & Eucalyptus parramattensis	Zeppel et al. (2008) Australian journal of botany.
Harvard forest data archive	Prospect Hill Tract, Harvard Forest, USA	A set of four deciduous angiosperm tree species	Bassow & Bazzaz (1997) Oecologia.
Jean-Marc Limousin	Sevilleta NWR, PJ rainfall manipulation, USA	Juniperus monosperma & Pinus edulis	Limousin et al. (2013) Plant, Cell & Environment.
Jeff Kelly	Daintree forest, Cape Tribulation, QLD, Australia	A set of three tropical rainforest species	Unpublished data
Jeff Warren	ORNL FACE, TN, USA	Liqiudambar styraciflua	Warren et al. (2011) Ecohydrology.
Jesse Nippert	Konza Prairie, KS, USA	A set of C3 and C4 grassland species	Unpublished data
Joana Zaragoza-Castells, Patrick Meir & Owen Atkin	French Guiana	A set of tropical rainforest species	Unpublished data

Joana Zaragoza-Castells,			
Patrick Meir &			
Owen Atkin	Tambopata, Peru	A set of tropical species	Unpublished data
Johan Uddling	Rhinelander, WI, USA	Betula papyrifera & Populus tremuloides	Uddling et al (2009) Tree Physiology
John Drake	Duke Forest, Durham, NC, USA	Pinus taeda	Drake et al. (2011) Global Change Biology
Jonathan Bennie	Agoufou, Hombori, Mali	A set of African savanna tree species	Unpublished data
David Tissue	Narrabri, NSW, Australia	Cotton	Unpublished data
Kohei Koyama & Kihachiro Kikuzawa	Ishikawa, Japan	Fagus crenata	Koyama and Kikuzawa 2012 Ecological Research.
Kouki Hikosaka	Aobayama, Sendai, Japan	A set of nine angiosperm and gymnosperm tree species	Hikosaka and Shigeno (2009) Oecologia.
Kouki Hikosaka	TOEF, Tomakomai, Hokkaido, Japan	Quercus crispula	Hikosaka et al (2007) Tree Physiology.
Lasse Tarvainen & Göran Wallin	Skogaryd, Sweden	Picea abies	Tarvainen et al. (2013) Oecologia.
Lindsay Hutley & Samantha Setterfield	Wildman River, NT, Australia	Alloteropsis semialata & Andropogon gayanus	Unpublished data
Lisa Wingate	Aberfeldy, UK	Picea sitchensis	Wingate et al. (2007) Plant, Cell & Environment.
Lucas Cernusak	Howard Springs, NT, Australia	A set of evergreen savanna tree species	Cernusak et al. (2011) Agriculture & Forest Meteorology.
Lucas Cernusak	Daly River, NT, Australia	A set of evergreen savanna tree species	Cernusak et al. (2011) Agriculture & Forest Meteorology.
Lucas Cernusak	Dry River, NT, Australia	A set of evergreen savanna tree species	Cernusak et al. (2011) Agriculture & Forest Meteorology.
Lucas Cernusak	Adelaide River, NT, Australia	A set of evergreen savanna tree species	Cernusak et al. (2011) Agriculture & Forest Meteorology.
Lucas Cernusak	Sturt Plains, NT, Australia	A set of evergreen savanna tree species	Cernusak et al. (2011) Agriculture & Forest Meteorology.
Lucas Cernusak	Boulia, QLD, Australia	A set of evergreen savanna tree species	Cernusak et al. (2011) Agriculture & Forest Meteorology.
Lucy Rowland & Patrick Meir	Caxiuana, Brazil	Manilkara spp.	Unpublished data
Maj-Lena Linderson & Teis Nørgaard Mikkelsen	Soroe, Denmark	Fagus sylvatica	Linderson et al. (2012) Agriculture & Forest Meteorology

Mark Broadmeadow	Headley S. London, UK	Three Quercus species	Broadmeadow et al. (1999) Water, Air and Soil Pollution.
Markus Löw	Kranzberg forest, Germany	Fagus sylvatica	Op de Beeck et al. (2010) Agriculture & Forest Meteorology.
Michael Freeman	Soroe, Denmark	Fagus sylvatica	Freeman, M. (1998) PhD Thesis.
Nicolas Martin-StPaul	Les Mages, France	Quercus ilex	Martin-StPaul et al. (2012) Functional Plant Biology.
Nicolas Martin-StPaul	Puechabon, France	Quercus ilex	Martin-StPaul et al. (2012) Functional Plant Biology.
Nicolas Martin-StPaul	Vic la Gardiole, France	Quercus ilex	Martin-StPaul et al. (2012) Functional Plant Biology.
Oula Ghannoum	Brian Pastures Res. Stn, Gayndah, QLD, Australia	A set of C4 grasses	Unpublished data
Paolo de Angelis	Montalto di Castro, Italy	Phillyrea angustifolia, Pistacia lentiscus & Quercus ilex	Scarascia-Mugnozza et al. (1996) Plant, Cell & Environment.
Pasi Kolari	Hyytiälä, Finland	Pinus sylvestris	Kolari et al. (2007) Tellus.
Patrick Mitchell	Corrigin Water Reserve, WA, Australia	Eucalyptus capillosa & Eucalyptus salmonophloiia	Mitchell et al. (2009) Agriculture & Forest Meteorology.
Qingmin Han	FFPRI, Tsukuba, Ibaraki, Japan	Chamaecyparis obtusa	Han et al. (2009) Journal of forest research.
Qingmin Han	Mt Fuji, Japan	Pinus densiflora	Han et al. (2003) Tree Physiology.
Maarten Op de Beeck	Tervuren, Belgium	Brassica napus & Brassica oleracea	Op de Beeck et al. (2010) Environmental Pollution.
Sabine Tausz-Posch	AGFACE facility, Horsham, VIC, Australia	Triticum aestivum two varieties	Tausz-Posch et al. (2013) Physiologia Plantarum.
Teresa E. Gimeno	Alto Tajo Natural Park, Guadalajara, Spain	Juniperus thurifera	Gimeno et al. (2012) Tree Physiology.
Victor Resco de Dios	Santa Rita Experimental Range, USA	Eragrostis lehmanniana & Heteropogon contortus	VRD et al. (2012) Prespectives in Plant Ecology, Evolution and Systematics.
Wei Sun	Charleston mesquite site, Tombstone, AZ, USA	A set of mesquite C3 and C4 grass species	Sun et al. (2009) Plant, Cell & Environment.
Wei Sun	San Pedro, Sierra Vista, AZ, USA	A set of riparian C3 and C4 grass species	Sun et al. (2010) Oecologia.
Yusuke Onoda	Hakkoda, Aomori, Japan	Fagus crenata, Lindera umbellata & Magnolia salicifolia	Yasumura et al. (2005) & Onoda unpublished.

Classification scheme	Class	g_1 mean	$g_1 SE$	Number of data points	Number of species
a_Pathway	C4	1.62	0.03	1161	38
	C3	4.16	0.01	14001	276
b_Plantform	Gymno. tree	2.35	0.02	4732	13
	shrub	3.32	0.05	689	15
	Angio. tree	3.97	0.02	6265	203
	Grass	5.25	0.13	304	20
	Savanna tree	5.76	0.22	339	20
	Crop	5.79	0.04	1672	5
c_T region	Arctic	2.22	0.07	162	8
	Boreal	2.19	0.02	917	5
	Temperate	4.31	0.02	11934	75
	Tropical	4.43	0.08	988	189
d_W region	MI < 0.5	3.77	0.03	3328	17
	0.5 <mi<1.0< td=""><td>4.69</td><td>0.04</td><td>1673</td><td>45</td></mi<1.0<>	4.69	0.04	1673	45
	1.0 <mi<1.5< td=""><td>3.87</td><td>0.03</td><td>4313</td><td>29</td></mi<1.5<>	3.87	0.03	4313	29
	MI<1.5	4.02	0.02	4687	186
e_PFTs	C4 grass	1.62	0.03	1161	38
	Ever. gymno. tree	2.35	0.02	4732	13
	Deci. savanna tree	2.98	0.39	30	2
	Shrub	3.32	0.05	689	15
	Ever. angio. tree	3.37	0.03	2828	17
	Trop. Rainforest tree	3.77	0.06	549	167
	Deci. angio. tree	4.64	0.04	2888	19
	C3 grass	5.25	0.13	304	20
	C3 crop	5.79	0.04	1672	5
	Ever. savanna tree	7.18	0.25	309	18

433 Table S2: Estimates of g_1 by different classification schemes.

Table S3: Model coefficients for g_1 **as a function of MI and mGDD**₀**.** The model was fitted

436 with a linear mixed-effects model as $log(g_1) \sim MI + mGDD_0 + MI^*mGDD_0$ using different PFTs

437 as the random effects to account for the differences in intercept among PFTs.

Model				
Variables	mean	SE	DF	
Intercept	0.449	0.289	97	
MI	0.033	0.013	97	
mGDD ₀	0.027	0.192	97	
MI*mGDD ₀	0.014	0.012	97	

440 Supplementary Figure legends

441	Fig. S1: Climatic space covered by the Stomatal Behaviour Synthesis Database. Shown as
442	a combination of mean annual temperature (MAT; °C), mean annual precipitation (MAP; mm),
443	mean annual degree days above $0^{\circ}C$ (mGDD ₀ ; $^{\circ}C$) and moisure index (MI).
444	
445	Fig. S2. Residual plot by PFTs for the model: $log(g_1) \sim MI + mGDD_0 + MI * mGDD_0$. The
446	model was fitted using linear mix-effects model with PFTs as the random effect to account for
447	the differences in intercept among PFTs.
448	
449	Fig. S3. predicted $log(g_1)$ as a function of mGDD ₀ and MI. (a) the predicted $log(g_1)$ under
450	different ranges of MI and mGDD ₀ presented as partial regression plot. Predictions are from
451	linear mixed-effects model for $log(g_1)$ assuming PFTs as a random effect to account for the
452	differences in intercept among PFTs. Colour lines represent the predicted g_1 based on fitted
453	model coefficients (Table S3). Colour dots represent the partial regression predictions at a
454	given fixed MI or mGDD ₀ level.
455	
456	
457	
458	
459	
460	

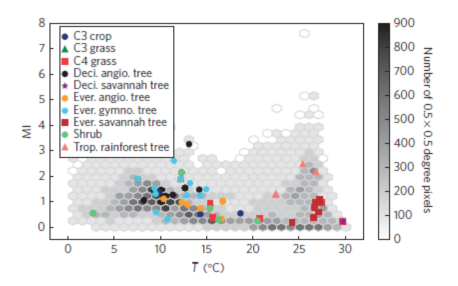


Figure 1 | Climatic space covered by the Stomatal Behaviour Synthesis Database, shown as mean temperature during the period with daily mean temperatures above 0 °C and moisture index. Coloured circles represent climatic space for the database, with different colours indicating different plant functional types. Grey hexagons represent global climatic space for which vegetation is present. The global climatic space data were binned by every 1°C for temperatures above 0 °C (T) and every 0.25 for the moisture index (MI). The grey scale bar indicates the number of 0.5×0.5 degree pixels for a given binned T and MI combination.

461

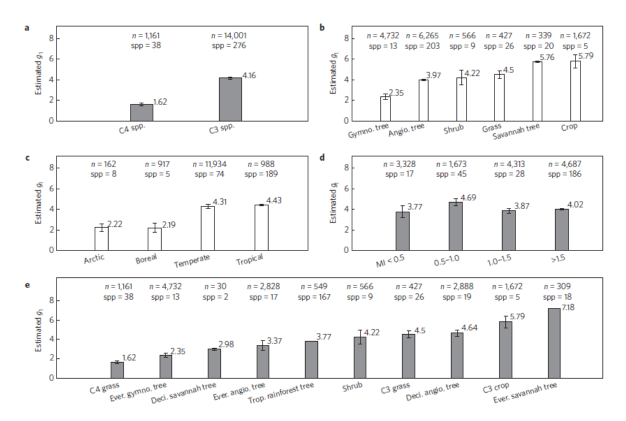


Figure 2 | Mean g_1 values for plant functional types defined by different classification schemes. Each bar represents the mean values \pm 1SE of g_1 from the stomatal model fitted using a nonlinear mixed-effects model assuming species as a random effect. The sample sizes (n) are the number of measurements. In the case of diurnal measurements, measurements might be done on the same leaf but under different environmental conditions. Species number (spp) indicates the number of the species in each group. Panels **b**-**d** include C₃ species data only.

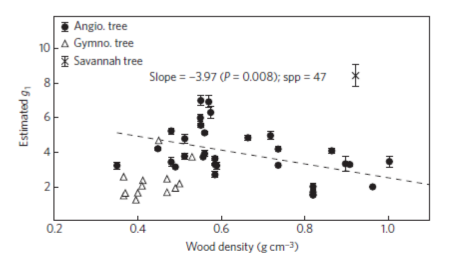


Figure 3 | **Relationship between** g_1 and wood density for angiosperm and gymnosperm trees. Savannah tree species (all of which were angiosperms) are indicated separately. Each data point represents mean \pm 1SE of g_1 for an individual species fitted with a nonlinear regression model. A linear regression line was fitted only for angiosperm trees due to the lack of a significant linear relationship for gymnosperm trees. The fitted linear regression relationship between g_1 and wood density for angiosperm trees is: $g_1 = -3.97^*$ WD + 6.53 (P = 0.0008, $R^2 = 0.21$). Wood density data were obtained from Global Wood Density Database^{2,29} and are available for 47 species in the Stomatal Behaviour Synthesis Database. The wood density database is a collection of published data based on actual measurements.

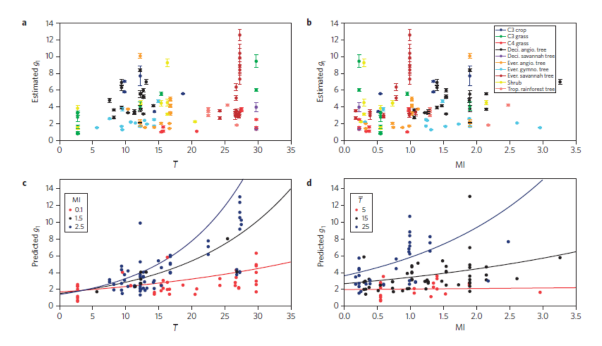


Figure 4 | Estimated and predicted g_1 **as a function of \tilde{T} and MI. a,b**, Relationship between estimated g_1 and mean temperature during the period with daily mean temperatures above 0 °C (\tilde{T} ; °C) (**a**) and moisture index (MI) (**b**) at experimental sites among species across different plant functional types (PFTs). Each data point represents the mean \pm 1SE of g_1 for individual species fitted with a nonlinear regression model. Classification of plant functional types are shown in Fig. 2e. **c**, **d**, Predicted g_1 under different ranges of MI (**c**) and \tilde{T} (**d**) presented as a partial regression plot. Predictions in **c** and **d** are from a weighted linear mixed-effects model for log (g_1) using the inverse of the SE of g_1 as weights to account for the uncertainty of g_1 fitting and assuming PFTs as a random effect to account for the differences in intercept among PFTs. Coloured lines represent the predicted g_1 based on fitted model coefficients (Supplementary Table 5). Coloured dots represent the partial regression predictions at a given fixed MI or \tilde{T} level.