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43

44 ABSTRACT

45 The conventional representation of Plant Functional Types (PFTs) in Dynamic Global 46 Vegetation Models (DGVMs) is increasingly recognized as simplistic and lacking in 47 predictive power. Key ecophysiological traits, including photosynthetic parameters, 48 are typically assigned single values for each PFT while the substantial trait variation 49 within PFTs is neglected. This includes continuous variation in response to 50 environmental factors, and differences linked to spatial and temporal niche 51 differentiation within communities. A much stronger empirical basis is required for 52 the treatment of continuous plant functional trait variation in DGVMs. We analyse 53 431 sets of measurements of leaf and plant traits, including photosynthetic 54 measurements, on evergreen angiosperm trees in tropical moist forests of Australia 55 and China. Confining attention to tropical moist forests, our analysis identifies trait 56 differences that are linked to vegetation dynamic roles. Coordination theory predicts 57 that Rubisco- and electron-transport limited rates of photosynthesis are co-limiting 58 under field conditions. The least-cost hypothesis predicts that air-to-leaf CO₂ 59 drawdown minimizes the combined costs per unit carbon assimilation of maintaining 60 carboxylation and transpiration capacities. Aspects of these predictions are supported 61 for within-community trait variation linked to canopy position, just as they are for 62 variation along spatial environmental gradients. Trait differences among plant species 63 occupying different structural and temporal niches may provide a basis for the 64 ecophysiological representation of vegetation dynamics in next-generation DGVMs.

65

66 Keywords: plant traits, photosynthesis, vegetation dynamics, tropical forests, DGVMs

67 Introduction

68 The development of Dynamic Global Vegetation Models (DGVMs) from the 69 earliest stages has emphasized the role of the distribution of different types of plants 70 and vegetation in predicting the exchanges of carbon between the atmosphere and the 71 land biota (Prentice et al., 2007; Prentice and Cowling, 2013). Plant Functional Type 72 (PFT) classifications can be traced back to the Raunkiær's (1934) 'life form' 73 classification, based on plant traits that ensure persistence through seasons 74 unfavourable for growth (Harrison et al., 2010). After several decades during which 75 plant functional geography was neglected, new PFT classifications appeared during 76 the 1980s (Box, 1981; Woodward, 1987) with a view to the development of DGVMs 77 - which began in earnest during the late 1980s. The PFT concept has received 78 significant attention since then (Prentice et al., 1992; Díaz and Cabido, 1997; Lavorel 79 and Garnier, 2002; Wright et al., 2004; Prentice et al., 2007; Harrison et al., 2010; 80 Fyllas et al., 2012). It has become widely accepted that PFT classifications for 81 modelling purposes ideally should reflect aspects of trait diversity that can predict 82 plant responses to the physical environment.

83 PFTs adopted in DGVMs today are commonly defined in terms of up to five 84 qualitative traits: (a) life form, (b) leaf type, (c) phenological type, (d) photosynthetic 85 pathway and (e) climatic range defined in terms of broad climatic classes such as 86 'boreal' and 'tropical'. This conventional approach to PFT classification has manifold 87 limitations (Prentice and Cowling, 2013). For example, life-forms are often 88 incompletely defined in functional terms. Informal and potentially ambiguous terms 89 such as 'shrub' have been used in place of Raunkiær's explicit functional categories. 90 Leaf-type definitions usually ignore the huge variations in leaf size and shape among 91 'broad-leaved' plants. Even the distinction between broad and needle-leaved trees is 92 often effectively an imperfect surrogate for the important distinction in hydraulic 93 architecture between angiosperms and gymnosperms – the latter, in fact, including 94 many species with broad leaves. Thermal climate categories may make sense if they 95 are recognized as a surrogate for different cold-tolerance mechanisms in 96 phanerophytes (Prentice et al., 1992). Harrison et al. (2010) provided a more recent 97 compilation of experimental data on cold tolerance. However, such categories are often used without clear definitions. They may stand in for a continuum of 98 99 physiological differences between plants adapted or acclimated to different seasonal

100 temperature regimes, rather than representing qualitative differences among distinct 101 types of plant. Moreover, thermal climate categories may artificially restrict modelled 102 PFT distributions within confined areas even if the absence of a given PFT from a 103 wider area could be due to competitive exclusion by other types. Ideally, shifts of 104 dominance in models should not be imposed in this way, but should emerge naturally 105 through competitive advantage (Fisher et al., 2015).

106 A distinct aspect of plant functional classification pertains to species' 'roles' 107 in vegetation dynamics. Classifications of tree species according to shade tolerance 108 (Whitmore, 1982), growth characteristics: maximum height and growth rate (Shugart, 109 1984; Swaine and Whitmore, 1988) and successional stage (pioneer versus climax) 110 Swaine and Whitmore (1988) have been a mainstay of regionally specific 'gap 111 models' designed to predict forest dynamics under constant or changing 112 environmental conditions (Botkin et al., 1972; Shugart, 1984; Denslow, 1987; 113 Prentice and Leemans, 1990; Prentice et al., 1993; Turner, 2001) but are not treated 114 by most DGVMs. Exceptions are those models with individual-based dynamical 115 cores, such as LPJ-GUESS (Smith et al., 2001), Hybrid (Friend et al., 1993), ED 116 (Moorcroft et al., 2001; Medvigy et al., 2009), aDGVM2 (Langan et al., 2017), and 117 models that make use of the Perfect Plasticity Approximation (PPA). PPA is a 118 mathematical approach designed to represent the essentials of forest dynamics 119 without simulating individual trees explicitly (Purves et al., 2008; Fyllas et al., 2014; 120 Fisher et al., 2015).

121 A general critique of the use of PFTs for modelling purposes has emerged 122 with the development of new dynamic vegetation models based on continuous trait 123 variation (Pavlick et al., 2013; Scheiter et al., 2013; Verheijen et al., 2013; Fyllas et 124 al., 2014; van Bodegom et al., 2014; Sakschewski et al., 2015), raising a question as 125 to whether distinct PFTs are necessary for modelling vegetation. In our view there is a 126 clear-cut case for retaining the well-understood distinctions among photosynthetic 127 pathways, and there may also be good reasons also to retain life-form distinctions - at 128 least at the highest level of Raunkiær's classification. However, most quantitative 129 traits show continuous adaptive variation along environmental gradients (Meng et al., 130 2015), indicating that the conventional approach of assigning fixed values of leaf-131 level traits such as carboxylation capacity (V_{cmax}) and nitrogen content per unit leaf area (N_{area}) to PFTs does not adequately describe the plasticity of such traits within 132 133 species (phenotypic plasticity). Even biophysical traits such as leaf mass per area 134 (LMA) and leaf dry-matter content, which are typically less plastic than metabolic 135 traits (Meng et al., 2015; Dong et al., 2017), show systematic, quantitative variations 136 along environmental gradients, partly as a consequence of species turnover within 137 PFTs and thus not necessarily the replacement of one PFT by another. Faced with 138 continuous trait variation (due to species or genotypic turnover and/or phenotypic 139 plasticity), models can either treat it as continuous – as the LPJ DGVM (Sitch et al., 140 2003) does for photosynthetic traits, following the approach developed by Haxeltine 141 and Prentice (1996) - or subdivide the continuum into arbitrary sections. However, 142 problems such as unrealistically abrupt modelled vegetation transitions can arise if the 143 subdivision of the continuum is too coarse, suggesting that a continuous 144 representation will be more useful.

145 This paper describes an empirical analysis that is oriented towards the 146 improvement of DGVMs. Our primary focus is on the largely neglected 'dynamical' 147 aspect of PFT classification. We adopt the fourfold scheme introduced by Shugart 148 (1984) as an initial scheme to classify species' dynamic roles. We recognize that this 149 classification represents a subdivision of two orthogonal continua of variation: shade 150 tolerance (requiring, versus not requiring, a gap for regeneration) and size at maturity 151 (producing, versus not producing, a gap upon mortality). We focus on functional trait 152 variations within tropical moist forests, which harbour enormous tree species diversity 153 and contain species that exhibit all combinations of these traits (Turner, 2001). We 154 build on a previous analysis by Fyllas et al. (2012), who showed that quantitative traits including foliar δ^{13} C discrimination, LMA and nutrients including N and P 155 156 could be used to discriminate PFTs with distinct dynamic characteristics in 157 Amazonian rain forests. Our analysis focuses on east Asian (SW China) and northern 158 Australian tropical rain forests, and extends the approach of Fyllas et al. (2012) to 159 include field photosynthetic measurements.

160 **Theory and Principles**

161 Recent empirical analyses aiming to inform the development of 'next-162 generation' DGVMs have focused on the predictability of key quantitative traits as a 163 function of environmental variation (Yang et al., 2018). The 'least-cost' (Prentice et 164 al., 2014) and 'coordination' (Maire et al., 2012) hypotheses together suggest a degree 165 of predictability for the air-to-leaf CO₂ drawdown (χ , the ratio of leaf-internal to ambient CO₂) (Prentice et al., 2014; Wang et al., 2017), V_{cmax} and the electrontransport capacity J_{max} (Togashi et al., 2017), and N_{area} (Dong et al., 2017) across environments and clades. Both hypotheses have a much longer pedigree than indicated by the recent references cited here, but systematic testing of these hypotheses has only been undertaken quite recently.

171 The least-cost hypothesis proposes that at the leaf level, plants should respond 172 to differences in the relative costs (per unit of assimilation achieved) of maintaining 173 the biochemical capacity for photosynthesis versus the structural capacity for 174 transpiration by making an optimal investment 'decision' that minimizes the total 175 carbon cost of maintaining both essential functions. This hypothesis can be shown to 176 lead to an optimum value of χ that depends predictably on temperature, vapour 177 pressure deficit and atmospheric pressure (Prentice et al., 2014; Wang et al., 2017). 178 The mathematical expression of this optimum value includes a parameter that is 179 influenced by low plant-available moisture, and therefore by soil moisture and rooting 180 characteristics (Zhou et al., 2013). This optimum has the same mathematical form as 181 that predicted approximately by the Cowan-Farquhar optimality criterion for electron-182 transport limited photosynthesis. This form is known to provide good predictions of 183 stomatal behaviour under a range of conditions (Medlyn et al., 2011; Lin et al., 2015; 184 Dewar et al., 2018). The least-cost hypothesis however is more explicit than the 185 Cowan-Farquhar criterion in that it 'unpacks' water transport and biochemical costs, 186 and assigns to each of them an explicit ecophysiological meaning.

187 The coordination hypothesis indicates that under typical daytime conditions, 188 the Rubisco-limited and electron transport-limited rates of photosynthesis should be 189 approximately equal (Chen et al., 1993; Haxeltine and Prentice, 1996; Maire et al., 190 2012). This represents the optimal disposition of resources between light capture and 191 carbon fixation. It leads to the prediction that the outer-canopy V_{cmax} measured at the 192 prevailing growth temperature should be determined by χ (higher values of one 193 quantity are consistent with lower values of the other), temperature (higher V_{cmax} is 194 required to achieve a given assimilation rate at higher temperatures), and incident 195 photosynthetically active radiation (PAR) (productive investment in V_{cmax} is directly 196 proportional to the available PAR) (Dong et al., 2017). Narea is generally found to be 197 roughly linearly related to Rubisco content, and thus to $V_{\rm cmax}$ at standard temperature. 198 However, leaf N also has structural and defensive components that are roughly 199 proportional to LMA and represent a largely independent source of variation in N_{area}

200 (Dong et al., 2017). So far, these predictions have been supported for seasonal 201 variations within individual plants (Togashi et al., 2017), and for spatial variations 202 along environmental gradients (Prentice et al., 2014; Dong et al., 2017). Here we 203 extend their application to biotically conditioned, microenvironmental variation 204 within forest environments. The framework provided by the least-cost and 205 coordination hypotheses suggest moreover that shade tolerance, and stem properties 206 such as height and wood density, should also be related to leaf metabolic and 207 structural traits, as proposed by Whitehead et al. (1984) and many later commentators.

208 The least-cost and coordination hypotheses are optimality concepts, whose 209 rationale depends on the heuristic principle that natural selection is expected to have 210 eliminated all trait combinations that fall short of optimality according to some 211 specified criterion. Another optimality concept lies behind the Leaf Economics 212 Spectrum, LES (Wright et al., 2004). Fundamentally, the LES represents a universal 213 negative correlation between LMA and leaf life-span (Lloyd et al., 2013), which can 214 be considered to arise from a trade-off because (a) carbon available for investment in 215 leaves is limited and (b) long-lived leaves need to be thicker and/or tougher than 216 short-lived leaves in order to avoid high risks of predation by herbivores and other 217 kinds of mechanical damage. Thus leaves can be short-lived and flimsy or long-lived 218 and thick and/or tough, or somewhere in between. In contrast, short-lived leaves with 219 high LMA would be uneconomic, while long-lived leaves with low LMA would be 220 unviable.

221 In this study we consider four groups of leaf and stem traits. The first group 222 consists of leaf metabolic traits: V_{cmax} , J_{max} and leaf dark respiration (R_{dark}), which has 223 been found to correlate with V_{cmax} (Atkin et al., 2000; Weerasinghe et al., 2014). The 224 second group contains the leaf structural/chemical traits N_{area} , P_{area} and LMA. As 225 previously noted, Narea has a metabolic component as well as a structural component, 226 and the same may be true for Parea (Evans, 1989; Reich et al., 1997; Fyllas et al., 227 2009). But increasing evidence points to the dominance of the structural component 228 when large sets of species are considered (Dong et al., 2017; Yang et al., 2018). The 229 third group, represented principally by wood density (WD), stands in for plant 230 hydraulics: denser wood tends to have lower permeability to water (Sperry, 2003; Lin 231 et al., 2015). Wood density also directly influences plant growth because volume 232 growth is necessarily slower, for a given photosynthetic output, in trees with dense 233 wood. Relatively more carbon also needs to be allocated to high-density wood, at the

234 expense of allocation to leaves and fine roots. Although the correlation between WD 235 and more directly instrumental traits for plant hydraulics, such as the Huber value (the 236 ratio of cross-sectional sapwood area to subtended leaf area) (Togashi et al., 2015), 237 vessel density and calibre, and permeability (Reid et al., 2005) is imperfect, these last 238 traits are much more time-consuming to measure than WD and thus comparatively 239 under-represented in available data sets. We anticipate that plants with a lavish water 240 use strategy should present high conductivity and low-density wood, while plants 241 adapted to environments with long droughts, or vulnerable to water use competition, 242 should tend to adopt a conservative water-use strategy and to have high-density wood. 243 The fourth group reflects the ability to compete with other species for light, expressed 244 as potential maximum height (H_{max}) : taller plants are able to harvest more light while 245 shorter plants are often more shade-tolerant (Turner, 2001). This group includes χ , 246 which has been reported to show a negative relationship with tree height (Koch et al., 247 2015). The reason χ belongs with H_{max} and not with V_{cmax} in this analysis is because 248 we have intentionally restricted the climatic range to moist forests, so that the 249 variation in χ is mainly related to plant strategy rather than to aridity or temperature. 250 H_{max} is considered to be a species characteristic and although it may not be reached at 251 every site, its use as an indicator of plant strategy is likely better than using actual 252 observed height, which varies among individuals and over time.

253 Leaf-level measurements were conducted in tropical forests of Queensland, Australia and Yunnan, China, and combined with (a) published and unpublished 254 255 datasets on H_{max} , WD, and (b) expert knowledge of the species' vegetation dynamical 256 roles (climax, subcanopy, large and small pioneers). Thus we are able to present what 257 is (to our knowledge) the first study to analyse key biochemical rates in the context of 258 tree species' contrasting dynamic roles, and the first empirical trait-based analysis to 259 include measured biochemical rates in a PFT classification with the aim to inform 260 progress in DGVM development. The objective of this work was to quantify trait 261 variation within these forests that can be linked to dynamical roles, but also 262 specifically to test the following predictions: (i) when moisture supply and low 263 temperatures are not limiting factors, photosynthetic capacity should be governed by incident PAR (a prediction of the coordination hypothesis); (ii) χ should be lower at 264 265 high H_{max} (a prediction of the least-cost hypothesis); and (iii) pioneer species should

tend to have low WD, an expectation from the theory of forest dynamics (Shugart,1984).

268 Materials and methods

269 Study Sites

270 Our analysis includes material from 232 evergreen angiosperm tree species 271 (431 leaf samples) from moist tropical forests of Queensland, northern Australia 272 (Robson Creek on the inland Atherton Tablelands to Cape Tribulation near the Pacific 273 coast) and Yunnan, southwestern China (the Xishuangbanna region in southern 274 Yunnan, near the land borders with Myanmar, Thailand and Laos). Field campaigns 275 conducted in Queensland and Yunnan yielded data on 191 species. Data from these 276 campaigns were combined with data on a further 41 Queensland species from field 277 studies carried out by the TROpical Biomes In Transition (TROBIT) network 278 (Bloomfield et al., 2014). Climates covered by the sampled areas range in mean 279 annual precipitation (MAP) from 1427 to 5143 mm (Liddell, 2013b, a; Harris et al., 280 2014; Hutchinson, 2014c) and in mean annual temperature (MAT) between 19.0 and 281 24.4 °C (Liddell, 2013b, a; Harris et al., 2014; Hutchinson, 2014a, b). Both 282 Oueensland and Yunnan have a marked wet season and 'dry' (drier) season, and 283 although the range of MAP values is considerable, all the sites correspond to climates 284 where moisture is unlikely to be a limiting factor for forest development. The drier 285 season at the Queensland and Yunnan sites lasts four to five months, but there is still 286 typically 100-300 mm precipitation per month. Gridded climatological data at 0.01° 287 resolution for 1971-2000 on MAP, MAT, annual Moisture Index (MI, the ratio of 288 precipitation to equilibrium evaporation) and mean monthly photosynthetic active 289 radiation (mPAR) were acquired for the Australian sites at www.tern.org.au. 290 Climatological data for the Chinese sites were derived from records at 1814 291 meteorological stations (740 stations have observations from 1971-2000, the rest from 292 1981-1990: China Meteorological Administration, unpublished data), interpolated to a 293 0.01° grid using a three dimensional thin-plate spline (ANUSPLIN version 4.36, 294 Hancock and Hutchinson, 2006). Fig. 1 and Table 1 provide further details on sites 295 and climates.

296 Gas exchange measurements and photosynthetic variables

297 We used a portable infrared gas analyser (IRGA) system (LI-6400; Li-Cor, 298 Inc., Lincoln, NB, USA) to perform leaf gas-exchange measurements. Sunlit terminal 299 branches from the top one-third of the canopy were collected and immediately re-cut 300 under water. One of the youngest fully expanded leaves, attached to the branch, was 301 sealed in the leaf chamber. Measurements in the field were taken with relative 302 humidity and chamber block temperature close to those of the ambient air at the time of measurement. The rate of airflow was held constant at 500 μ mol s⁻¹, but 303 exceptionally the flow was reduced (to a minimum of 250 μ mol s⁻¹) under very low 304 305 stomatal conductance.

306 We obtained 130 A-c_i curves from 41 species from Robson Creek (RCR1 and 307 *RCR2*) in both the dry and the wet season. The CO₂ mixing ratios for the A- c_i curves 308 proceeded stepwise down from 400 to 35 and up to 2000 μ mol mol⁻¹. Prior to the 309 measurements, we tested plants to determine appropriate light-saturation levels. The 310 photosynthetic photon flux density (PPFD) adopted for measurement ranged between 1500 and 1800 μ mol m⁻² s⁻¹. After measuring the A-c_i curves over about 35 minutes, 311 312 light was set to zero for five minutes before measuring respiration. This was a time-313 saving compromise to allow four or five replicate curves per machine per day, based 314 on our experience that stable results are commonly obtained after five minutes. 315 Moreover, this quick estimate should be superior to the common practice of deriving 316 R_{dark} as one of the parameters in a curve-fitting routine. Following the protocol of 317 Domingues et al. (2010), we discarded 37 of a total 167 A- c_i curves in which stomatal 318 conductance (g_s) declined to very low levels, adversely affecting the calculation of 319 $V_{\rm cmax}$. These procedures were very similar to the ones applied to the 125 A- c_i curves 320 obtained from 26 species in the TROBIT sites (CTR2, KBL1, KBL3 and KCR) during 321 the wet season, further described in Bloomfield et al. (2014).

We sampled 114 leaves of 91 species in Yunnan (*Y1X*, *Y2U*, *Y3M1*, *Y3M2*, *Y4L*) in the dry season. Data for 62 leaves of 16 species were also obtained from Cape Tribulation in the dry season (*CTR1*:Weerasinghe et al., 2014). We used the same sampling methods for Yunnan and for Cape Tribulation. PPFD was held constant at 1800 µmol m⁻² s⁻¹. For each leaf, we first set the CO₂ mixing ratio to 400 µmol mol⁻¹ to obtain the rate of photosynthesis under light saturation (*A*_{sat}). Measurement was taken under stable g_s (> 0.5 µmol m⁻² s⁻¹), CO₂ and leaf-to-air vapour pressure deficit. The next step was to increase the CO₂ mixing ratio to 2000 μ mol mol⁻¹ in order to register the rate of photosynthesis under light and CO₂ saturation (A_{max}). R_{dark} was not measured in Yunnan. For R_{dark} in *CTR1*, the leaf was wrapped in foil sheets after A_{sat} and A_{max} measurements. There was a waiting period of at least 30 minutes of darkness before taking R_{dark} values.

334 Values of V_{cmax} and J_{max} were fitted using the Farquhar et al. (1980) model. 335 The assumption of unlimited mesophyll conductance (Miyazawa and Kikuzawa, 336 2006; Lin et al., 2013) remains the standard implementation of the Farquhar model 337 although it is recognized to be an approximation that results in an overestimation of 338 $V_{\rm cmax}$ and $J_{\rm max}$. Hence all of the values estimated are 'apparent' $V_{\rm cmax}$ and $J_{\rm max}$ values, 339 as in most of the ecophysiological literature. In cases where $A-c_i$ curves were not 340 measured, we estimated V_{cmax} from A_{sat} by the so-called one-point method, which 341 inverts the equation for Rubisco-limited photosynthesis taking into account the 342 measured c_i and leaf temperature by applying the temperature dependencies of the 343 Michaelis-Menten coefficients of Rubisco for carboxykation (K_C) and oxygenation 344 (K_0) and the photorespiratory compensation point (Γ^*) from Bernacchi et al. (2001). 345 The one-point method relies on the assumption that light-saturated photosynthesis 346 measured on field-grown plants is Rubisco-limited, which has been found to be true 347 in almost all cases (De Kauwe et al., 2016). J_{max} was estimated from A_{max} on the 348 assumption that high CO₂ forces the leaves into electron-transport limitation 349 (Bernacchi et al., 2003). Triose phosphate utilization limitation was not considered, as 350 it would be unlikely to occur at our field temperatures > 22 °C (Sharkey et al., 2007; 351 Lombardozzi et al., 2018)

352 Nutrient analyses

353 After completion of the leaf gas-exchange measurements, the leaf was retained 354 to determine leaf area, dry mass, and mass-based N and P concentrations (N_{mass} and $P_{\text{mass.}}$ mg g⁻¹). Leaves were sealed in plastic bags containing moist tissue paper to 355 356 prevent wilting. Leaf area was determined using a 600 dot/inch flatbed top-357 illuminated optical scanner and Image J software (http://imagej.nih.gov/ij/). Leaves 358 were dried in a portable desiccator for 48 hours for preservation until the end of the 359 campaign and subsequently oven-dried in the laboratory for 24 hours at 70°C. Then 360 the dry weight was determined (Mettler-Toledo Ltd, Port Melbourne, Victoria,

Australia). LMA (g m⁻²) was calculated from leaf area and dry mass. N_{mass} and P_{mass} 361 362 were obtained by Kjeldahl acid digestion of the same leaves (Allen et al., 1974). The 363 leaf material was digested using 98% sulphuric acid and 30% hydrogen peroxide. 364 Digested material was analyzed for N and P using a flow injection analyser system (LaChat QuikChem 8500 Series 2, Lachat Instruments, Milwaukee, WI, USA). Narea 365 and P_{area} (mg m⁻²) were calculated as products of LMA and N_{mass} or P_{mass} . TROBIT 366 367 nutrient analysis was performed using similar methods but different equipment, as 368 described in Bloomfield et al. (2014).

369 Wood density and tree height

Twenty-year series of wood density (*WD*), tree height (*H*), and tree diameter at breast height (*D*) were obtained from Bradford et al. (2014a) (n = 138). Maximum tree height (H_{max}) was estimated using the derivative of the Mitscherlich function relating diameter and height (Li et al., 2014):

374
$$dH/dD = a \exp(-aD/H_{max}) = a (1 - H/H_{max})$$
(1)

where *a* is the initial slope of the relationship between height and diameter. A typical range of *a* in the literature is 116 ± 4.35 .

377 Dynamic roles of species

Australian species (n = 61) were assigned to dynamic roles with the help of the database published by Bradford et al. (2014b) and expert knowledge by MB. Chinese species (n = 85) were assigned to dynamic roles based on expert knowledge by ZH. These 'expert' classifications (A1) were compared with a quantitative traitbased classification (A2) as described in the next section. Both classification approaches were implemented according to the Shugart (1984) framework, which can also be related to those of Denslow (1987), Turner (2001) and Fyllas et al. (2012):

385

386 (1) Requires a gap, and produces a gap. These are long-lived pioneers that reach the387 canopy. Shade intolerant.

388 (2) Does not require a gap, but produces a gap. These are long-lived climax species389 that reach the canopy and grow large. Shade tolerant.

390 (3) Requires a gap, but does not produce a gap. These are short-lived pioneers that391 never grow large. Shade intolerant.

392 (4) Does not require a gap, and does not produce a gap. These are sub-canopy species.393 Shade tolerant.

394

The geographic distribution of expert assessment of dynamic roles per number of
species and per number of leaves is shown in Table 2. This dataset includes 262
observations.

398 Statistical analyses

399 All statistics were performed in R (R Core Team, 2012). For graphing we used the 400 ggplot2 package (Wickham, 2010). Moisture index was represented in Fig. 1 as its square root, a transformation appropriate to precipitation values (M.F. Hutchinson, 401 402 personal communication, 2011), which approximately normalizes the distribution of 403 values and thus contains the large spread of values at the high end. V_{cmax} , J_{max} , R_{dark} , 404 LMA, Narea, Parea, Hmax and WD data were log10-transformed, unless otherwise 405 indicated, achieving an approximately normal distribution of values. χ was logit-406 transformed as this variable is bounded between 0 and 1, and the logit transformation 407 results in approximately linear relationships between the transformed ratio and 408 environmental predictors, including temperature (Wang et al., 2017). Ordinary least-409 squares linear regression was used to test relationships between plant traits and 410 climate variables. Pairwise combinations of quantitative traits were tested for 411 significant relationships across all data, and within groups corresponding to high and 412 low MI, high and low mPAR, and high and low MAT. Slopes and elevations of 413 regressions were compared using standardized major axis regression with the smatr 414 package (Warton et al., 2006). The package vegan (Oksanen et al., 2015) was used 415 to assess multivariate trait variation, using the following methods:

416 417 Principal component analysis (PCA) of nine plant traits (V_{cmax}, J_{max}, R_{dark}, LMA, N_{area}, P_{area}, H_{max}, WD and χ);

418 419 • Redundancy analysis (RDA) of the same nine traits, constrained by three climate variables (MI, mPAR and MAT);

- 420
- RDA of the same nine traits, constrained by dynamic roles (as factors); and

421 • RDA of the same nine traits constrained simultaneously by both climate and
422 dynamic roles.

423 PCA was used to identify patterns of covariation among traits irrespective of their 424 dynamic or environmental correlates, and RDA to analyse multivariate trait 425 relationships to predictors. Note that PCA is an exploratory method with no 426 associated formal test of significance. By contrast, the significance of trait-427 environment relationships identified by RDA can be assessed approximately in a 428 similar way to generalized linear models (Ter Braak and Prentice, 1988). The K-429 means (R Core Team, 2012) clustering method was used to create four groups of 430 species based on the nine plant traits (A2: Dynamic roles based on quantitative 431 assessment). K-means clustering was performed with the number of iterations set to 432 100 and bootstrapped with 10,000 repetitions. RDA and bivariate correlations were 433 used to compare classifications A2 and A1. The dataset used for PCA and RDA 434 analysis consisted of 130 observations with information for all traits, climate variables 435 and dynamic roles. All RDA visualizations here follow the response-variable focused 436 'Type 2 scaling' (Oksanen et al., 2015), such that the angles between pairs of vectors 437 as plotted approximate their pairwise correlations. For PCA and RDA input data 438 where direct measurements of R_{dark} were not available, R_{dark} (n = 58) was estimated 439 from A_{sat} following Prentice et al. (2014) using the approximation $R_{\text{dark}} \approx 0.01 V_{\text{cmax}}$ 440 (De Kauwe et al., 2016).

441 Research data

Robson Creek (*RCR1* and *RCR2*) data can be requested at <u>www.tern.org.au</u>
(Prentice et al., 2013). Access to TROBIT data (*CTR2*, *KBL1*, *KBL3* and *KCR*) and
Cape Tribulation 1 (*CTR1*) are described in Bloomfield et al. (2014) and Weerasinghe
et al. (2014) respectively. For Yunnan data (*Y1X*, *Y2U*, *Y3M1*, *Y3M2*, *Y4L*), refer to
Wang et al. (2018).

447

448 **Results**

449 Trait values and dimensions of variation

450 Average values of the metabolic traits V_{cmax} , J_{max} , and R_{dark} were 52.0, 82.0, 451 and 0.63 µmol m⁻² s⁻¹ respectively. The corresponding ranges were 4.2 to 148.9, 14.0 to 203.6, and near zero to 3.70 μ mol m⁻² s⁻¹. Average values of the chemical/structural traits LMA, N_{area} and P_{area} were 110.9 × 10³, 0.19 × 10³ and 0.013 g m⁻² with ranges of 12.04 to 610.3 × 10³ (LMA), near zero to 1.49 g m⁻² (N_{area}), and near zero to 0.06 mg m⁻² (P_{area}). Average values of χ , H_{max} and WD were 0.71, 26.3 m and 0.55 g cm⁻³ with ranges of 0.39 to 0.94, 1.3 to 54.5 m, and 0.33 to 0.98 g cm⁻³ respectively.

458 Four orthogonal dimensions of trait variation were identified corresponding to 459 the metabolic, chemical/structural, hydraulic and height trait groups described above 460 (Fig. 2, Table 3). The metabolic traits V_{cmax} , J_{max} and R_{dark} varied continuously and in close correlation with one another. V_{cmax} and γ were negatively correlated, but the 461 462 correlation was weak (not shown: slope = -1.85, intercept = 1.32, R² = 0.13, p < 463 0.05). Table 3 makes it clear that variation in χ in this data set does not belong to the metabolic dimension. The chemical/structural traits LMA, Narea and Parea were 464 positively correlated with one another (p < 0.05), although the pairwise relationship of 465 466 P_{area} to LMA was weaker than that of N_{area} to LMA. The strong correlation between 467 LMA and N_{area} suggests that much of the N content in the leaves is structural rather 468 than metabolic (see also Yang et al., 2018). A similar result was obtained when mass-469 rather than area-based nutrient values were used in the PCA (not shown). The third 470 dimension was mostly represented by variation in WD, with some contribution from 471 P_{area} . Finally H_{max} and γ had a non-significant negative pairwise relationship and were 472 associated with the fourth dimension, suggesting a trade-off between water loss and 473 the length of the water-transport pathway. These dimensions of trait variation are 474 broadly in agreement with those described by Baraloto et al. (2010), Fyllas et al. 475 (2012) and Reich (2014).

476 **Contribution of climate variables to trait variation**

High and low values for MI, mPAR and MAT were defined as values above or below the mean value of the climate variable. Regression slopes between V_{cmax} and J_{max} for both high MI and low MI groups were close together but statistically distinct (p < 0.05, Fig. 3), and the same was true for high *versus* low mPAR and MAT groups (Fig. 3). Regressions for V_{cmax} *versus* N_{area} were not significant within these climatic groups (Fig. 3) but the relationship was significant, albeit weak, when all of the data were considered together (not shown: slope = 0.69, R² = 0.10, p < 0.05). The weakness of this relationship corroborates our previous assessment of leaf N content as being primarily structural rather than metabolic. High MI and low mPAR were associated with high V_{cmax} , J_{max} , and N_{area} (Fig. 3). These variables were also associated with R_{dark} and LMA. The remaining traits χ , P_{area} , H_{max} and WD were very scattered against MI and mPAR (no significant relationship). All traits had high and low values spanning the full range of MAT.

490 The clustered vectors for metabolic traits, MI and MAT in Fig. 4 indicate that 491 higher moisture and air temperature favour species with higher metabolic rates (Fig. 492 4). The RDA constrained by climate variables explained 35% of trait variation with 493 19% and 11% on axes 1 and 2 respectively (p < 0.05, Fig. 4). This represents an 494 unexpectedly large fraction of the trait variation, considering the modest range in 495 MAT (19 to 24°C), mPAR (27.4 to 30.5 mol m⁻²) and MI (0.9 to 2.5; i.e. typical 496 values for non-drought-stressed conditions) among these tropical moist forest sites. 497 No individually significant trait-climate variable relationship was found. An 498 association of high metabolic rates with aridity (Prentice et al., 2011) has been found 499 when considering longer climate gradients, but this is not apparent over the more 500 limited climatic range sampled here.

501 Contribution of dynamic roles to trait variation

The assignment of the four groups obtained by *K*-means clustering to dynamic roles was based on the degree of correspondence between the mean values of plant traits for each group and the classification by Shugart (1984). H_{max} determined whether a cluster was labeled as climax or large pioneer, or small pioneer or subcanopy. Higher values of photosynthetic traits defined tall trees as climax, as opposed to large pioneer, and small trees as small pioneer, as opposed to subcanopy.

Expert (A1) and quantitative (A2) role definitions explained 23% and 55% of total plant trait variance, respectively (Fig. 5). With respect to patterns, the RDA results obtained with the two classifications are quite similar to one another, which is expected as the clustering was performed using the same trait data represented in the RDA. However, the quantitative role definitions explained substantially more variance than the expert definitions. The major common patterns shown in the two RDA plots are as follows:

516 (1) The metabolic traits V_{cmax} , J_{max} , R_{dark} and the structural-chemical traits N_{area} and

517 LMA tend to be higher in climax species than in the other groups.

- 518 (2) P_{area} tends to be greater in subcanopy species than in the other groups.
- 519 (3) WD tends to be smaller in pioneer species than in the other groups.
- 520

521 These distinctions are supported, and further information provided, by the summary 522 statistics for trait variation within each group (Fig. 6). Climax species consistently 523 have the highest values of V_{cmax} , J_{max} , R_{dark} , LMA and N_{area} . High WD, consistent with 524 slow growth, characterized the subcanopy species. The χ ratio was lowest in climax 525 species and highest in subcanopy species. The scaling slopes of the bivariate 526 relationships between V_{cmax} and J_{max} , and between V_{cmax} and N_{area} , were largely similar 527 within each group, whether the roles were defined quantitatively or by expert 528 assessment (Fig. 7).

529 Partitioning trait variance to climate variables versus dynamic roles

530 RDA constrained by the two sets of predictors (climate and dynamic roles) 531 both separately and collectively provides the necessary information to partition the 532 total explained variation into the unique contributions of each set and a combined 533 contribution associated with covariation of the two sets, via the Legendre variation 534 partitioning method (Legendre and Anderson, 1999; Peres-Neto et al., 2006; Meng et 535 al., 2015; Yang et al., 2018). Based on the quantitative assessment of dynamic roles 536 (A2), RDA constrained by both sets of predictors explained 61% of trait variation, 537 which could be partitioned as follows: 26% from dynamic roles alone, 6% from 538 climate alone, and 29% from the combination. The corresponding figures based on 539 expert assessment were as follows: 43% of trait variation explained, composed of 8% 540 from dynamic roles alone, 20% from climate alone, and 15% from the combination.

Although significant trait variation was linked to climate, individual traitclimate relationships were weak and patterns that have been observed across a wider range of climates, such as the widely reported increase of N_{area} with aridity, were not present. This pattern is to be expected considering that semi-arid and arid ecosystems are not considered. Variance partitioning showed that between 8 and 26% of trait variation (depending on the source of information on dynamics roles) could not be attributed to the temperature and moisture regime, but could be related uniquely tospecies' dynamic roles.

549 Unexplained trait variance

550 Unexplained variance amounted to 57% and 39% for the expert and 551 quantitative assessments, respectively. In principle unexplained variance might be 552 related to a variety of factors including the season of measurement, forest age and 553 aspects of soil fertility. However, dividing the data according to wet-season (CTR2, 554 KBL1, KBL3, KCR, RCRs) versus dry-season (CTR1, RCRw, Y1X, Y2U, Y3M1, Y3M2, 555 Y4L) measurements yielded patterns similar to those found in the full data set. No data 556 on forest age were available. No correlations were found between trait values and soil 557 total N, soil total P and cation exchange capacity (Table 1).

558

559 **Discussion**

560 This study provides support for the idea that forest dynamic roles, as described 561 by Shugart (1984), might be systematically related to the biophysical and 562 ecophysiological traits used in DGVMs. Our analysis explores plant trait diversity and 563 plasticity with a view to more realistic modelling of plant and vegetation processes, 564 whether for local or global model applications (Fyllas et al., 2009; 2012; Quesada et 565 al., 2012). Expert classification of dynamic roles in forests is notoriously difficult 566 because it requires observation over many decades. Our quantitative analyses suggest 567 a possible alternative approach to classification based on trait measurements at one 568 point in time. Moreover, our results have supported certain specific predictions of the 569 least-cost and coordination hypotheses, which are key to explaining species strategies, 570 community assembly and ecosystem structure and function (Reich, 2014). They 571 collectively hold the promise of providing general, testable trait-environment 572 relationships that could reduce the excessive number of parameters required by most 573 DGVMs (Prentice et al., 2015).

574 **Dynamic roles and the coordination hypothesis**

575 Our results support a core prediction of the coordination hypothesis for 576 forests: that J_{max} and V_{cmax} should be higher under high illumination and lower in the 577 shade, as seen both in the vertical gradient of light-saturated assimilation rates in 578 dense canopies (Chen et al., 1993) and more generally, in the solar radiation gradient 579 across canopies situated in diverse environments (Maire et al., 2012). With respect to 580 dynamic roles, outer-canopy climax species are expected to receive most PAR and 581 therefore should have the highest photosynthetic capacity, while subcanopy species 582 should have the lowest. Pioneer (gap-requiring) species would be expected to have 583 intermediate photosynthetic capacity and this too is consistent with our findings. 584 Additionally, the widely reported conservative ratio of J_{max} and V_{cmax} seems to be 585 maintained, both within and across dynamic roles. The association of R_{dark} with V_{cmax} 586 and J_{max} was also found to be strong, with R_{dark} maintaining a near constant ratio for 587 leaves whether in sun-exposed or shade conditions, as previously reported (e.g. by 588 Hirose and Werger, 1987; Weerasinghe et al., 2014; Atkin et al., 2015).

589 The observed relationships among N_{area} , P_{area} and LMA, and the weaker 590 correlations of these traits with primary metabolic traits, reflect the fact that a 591 substantial part of the N and P content of leaves is not directly tied to photosynthetic 592 functions (Dong et al., 2017). Although strong linear relationships between V_{cmax} (at a 593 reference temperature) and N_{area} seem to be widely expected, they are not always 594 found (Prentice et al., 2014; Togashi et al., 2017), perhaps due to the overpowering 595 effect of variation in structural and/or defensive components of leaf N. The 596 photosynthetic component of N_{area} however is expected to be proportional to incident PAR. This expectation is supported by the analysis of Dong et al. (2017), and by our 597 598 finding of highest Narea among climax species.

599 Dynamic roles and the least-cost hypothesis

600 It has been reported that γ declines with tree height. This too is a prediction of 601 the least-cost hypothesis (Prentice et al., 2014), as the cost of maintaining the water 602 transport pathway increases at with height. Therefore, tall trees - and the top stratum 603 of leaves in a tall tree, as noted by Koch et al. (2015) – may be expected to aim for a 604 lower optimum γ by investing more in the maintenance of biochemical capacity and 605 less in the maintenance of transport capacity. Even if the path-length effect on stem 606 hydraulic conductance is fully compensated by xylem tapering (Tyree and Ewers, 1991; Enquist and Bentley, 2012; Olson et al., 2014) as often seems to be the case, it 607 608 is still more expensive in terms of sapwood respiration to maintain a tall stem as

609 opposed to a short stem (Prentice et al., 2014). This prediction is supported by the low 610 χ found here in climax species. However, surprisingly, large pioneer species (with 611 H_{max} equivalent to climax species) did not show this adaptation. Subcanopy species 612 did nonetheless show high χ , consistent with their short stature. Given a reduced χ , the 613 coordination hypothesis then predicts that V_{cmax} should be increased. This mechanism 614 may additionally contribute to the high V_{cmax} found in climax species and the low 615 V_{cmax} in subcanopy species.

616 **Dynamic roles, the leaf economics spectrum and the theory of forest dynamics**

617 A third group of predictions broadly supported by our results comes from the 618 framework presented in Shugart (1984), Turner (2001) and Fyllas et al. (2012). This 619 approach considers two main axes of ecological specialization, one reflecting canopy 620 position and access to light, the other life span and growth rate. The main advantage 621 for large pioneers in rapidly achieving tall stature is to shade lower canopies nearby, 622 while obtaining rapid access to full sunlight. Compared to climax species, large 623 pioneers adopt a less conservative strategy regarding water use, and are likely to have 624 a shorter lifespan (Shugart, 1984; Reich, 2014). One way to achieve fast growth is to 625 invest in low-density conducting tissues, which implies lower WD. Subcanopy 626 species by contrast are necessarily shade-tolerant and often have traits associated with 627 slow growth. Our results support the existence of this tradeoff, with subcanopy trees 628 having generally high WD (a trait often accompanied by a high density of short and 629 narrow vessels: Reich, 2014).

630 According to the LES, across species globally, high LMA is linked to 631 longevity of individual leaves; and it has generally been found that LMA varies as 632 much or more within communities as with environmental gradients (Wright et al., 633 2004). Our data do not allow us to address the LMA-lifespan linkage directly. 634 However, they do show that LMA varies systematically across the dynamic roles, 635 being greatest in climax species and associated with high V_{cmax} and least in subcanopy species where it is associated with low V_{cmax} . These findings suggest a more nuanced 636 637 interpretation of the variation in LMA among dynamic roles. Namely: that thick, 638 high-LMA leaves are a pre-requisite for a leaf to attain V_{cmax} commensurate with high 639 levels of PAR at the top of a canopy (Niinemets and Tenhunen, 1997), while thin, 640 low-LMA leaves provide optimum light capture for the least investment in leaves -a good strategy for subcanopy species. Fast-growing pioneer species with their high
water-use strategy also require a low investment in leaf structure, developing thin,
low-LMA leaves in order to obtain a quicker return on investment (Turner, 2001).
The downside is that these leaves are likely to be more exposed to herbivory losses,
while the low-density stems are subject to the risks of cavitation and embolism,
shortening their life expectancy (Enquist and Bentley, 2012).

647 Implications for modelling

648 DGVMs based on continuous trait variation have been developed in response 649 to the growing realization that PFTs, as conventionally defined, do not adequately 650 describe the genotypic or phenotypic plasticity of plant traits in the real world. The 651 existence of systematic, adaptive trait variation in forests, within a climate range 652 where neither moisture nor low temperature is limiting, provides further support for 653 the conclusion (e.g. Meng et al., 2015) that models should not be based on fixed, 654 PFT-specific values for many quantitative traits. In general, consideration of the 655 adaptive function of trait differences among dynamic roles should contribute to 656 reducing the multiplicity of uncertain parameters, and simultaneously increase the 657 realism, of next-generation DGVMs. DGVMs in general (including recent trait-based 658 vegetation models, with the exception of the model of Fyllas et al., 2014 developed 659 for Amazonian forests) have paid minimal attention to the co-existing functional 660 diversity of traits present in communities where climate variation is small but tree 661 species diversity is large, including tropical forests. Our results suggest that the 662 framework provided by optimality concepts (the coordination and least-cost 663 hypotheses) could be combined with classical forest dynamics theory, which 664 differentiates complementary survival strategies for tree species in a highly 665 competitive environment, to yield successful predictions that would allow vegetation 666 dynamics to be represented more faithfully in DGVMs. The combination of these 667 different research strands can be achieved by extending existing predictions about 668 trait-environment relationships based on optimality considerations to cover biotically 669 induced microhabitat variation within complex plant communities.

We therefore suggest that the ecophysiological correlates of species dynamical roles be further analysed in other tropical and extratropical forests, as part of the empirical research required to establish a firmer foundation for next-generation vegetation models. Moreover, we look forward to the widespread use of adaptive
schemes in which trait combinations, such as those characterizing species' dynamical
roles, emerge naturally from the competition among plants.

676

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700 Tables

Table 1. Climate averages (MAT = mean annual temperature, MI = Moisture Index,
mPAR = mean monthly photosynthetic active radiation), geographic location,
elevation above sea level and soil properties (CEC = cation exchange capacity, TN =
total soil nitrogen, TP = total soil phosphorus) of the study sites in north-east
Australia (*CTR1*, *CTR2*, *KBL1*, *KBL3*, *KCR*, *RCRs*, *RCRw*) and south-east China
(*Y1X*, *Y2U*, *Y3M1*, *Y3M2*, *Y4L*).

SITE	LON	LAT	Altitude (m)	MAT (°C)	MAP (mm)	mPAR (mol/m²)	МІ	CEC(cmol/kg)	TN (%)	TP (%)
CTR1	145.45	-16.10	64	24.4	5143	27.5	2.54	11.8	0.64	0.023
CTR2	145.45	-16.10	90	24.4	5143	27.5	2.54	11.8	0.02	0.011
KBL1	145.54	-17.76	761	20.4	1976	28.2	1.39	10.83	0.08	0.030
KBL3	145.54	-17.69	1055	19.0	1726	28.3	1.22	11.11	0.08	0.030
KCR	145.60	-17.11	813	19.6	2541	27.9	1.82	9.81	0.01	0.006
RCRs	145.63	-17.12	700	19.4	2246	27.9	1.29	4.3	0.18	0.019
RCRw	145.63	-17.12	700	19.4	2246	27.9	1.29	4.3	0.18	0.019
Y1X	101.27	21.92	502	21.7	1427	30.1	0.94	8.68	0.08	0.044
Y2U	101.24	21.98	1075	19.7	1562	30.6	1.03	6.09	0.08	0.044
Y3M1	101.58	21.61	668	19.6	1662	29.8	1.14	10.21	0.08	0.050
Y3M2	101.58	21.62	828	20.5	1604	29.9	1.07	10.21	0.08	0.050
Y4L	101.58	21.62	1034	20.5	1604	30.1	1.06	10.21	0.08	0.050

707

Table 2. Geographic distribution of expert assessment of dynamic roles per number of
species and per number of leaves for the study sites in north-east Australia (*CTR1*, *CTR2*, *KBL1*, *KBL3*, *KCR*, *RCRs*, *RCRw*) and south-east China (*Y1X*, *Y2U*, *Y3M1*, *Y3M2*, *Y4L*). This dataset includes 262 observations.

SITE	Number of species					Number of leaves				
	climax	large pioneer	Small pioneer	subcanopy		climax	large pioneer	Small pioneer	subcanopy	
CTR1	7	4		1		27	16		4	
CTR2	4	1				11	5			
KBL1	2	4				7	25			
KBL3	3	4				17	13			
KCR	3	5				9	17			
RCRs	7	16	3	4		9	52	7	4	
RCRw		8	1				26	4		
Y1X		2					2			
Y2U		3		1			3		1	
Y3M1				1					2	
Y3M2				1					1	
Y4L										

	PC1	PC2	PC3	PC4
$\log V_{cmax}$	-0.48	-0.27	0.21	0
log J _{max}	-0.46	-0.25	0.18	0.04
log R _{dark}	-0.45	-0.16	0.27	-0.01
log LMA	-0.32	0.55	-0.21	0.03
log N _{area}	-0.37	0.45	-0.24	-0.12
log P _{area}	-0.08	0.54	0.45	0.03
log WD	0.15	0.1	0.68	-0.16
$\log H_{max}$	-0.2	-0.15	-0.24	-0.74
logit χ	0.23	0.11	0.16	-0.64
	PC1	PC2	PC3	PC4
Standard				
deviation	1.84	1.26	1.09	0.98
Proportion				
of Variance	0.37	0.18	0.13	0.11
Cumulative				
Proportion	0.37	0.55	0.68	0.79

Table 3. Principal Component Analysis loadings for nine traits. The highest
correlations (absolute magnitudes > 0.45) are indicated in bold.





Fig. 1. Mean annual temperature (MAT, °C), the square root of Moisture Index (MI,
ratio of precipitation to equilibrium evapotranspiration) and mean monthly
photosynthetic active radiation (mPAR, mol m⁻²) for northern Australia (*CTR1*, *CTR2*, *KBL1*, *KBL3*, *KCR*, *RCRs*, *RCRw*) and southwestern China (*Y1X*, *Y2U*, *Y3M1*, *Y3M2*, *Y4L*).



Fig. 2. Principal component analysis (PCA) of nine traits in northern Australia and southwestern China (n = 130). Blue dotted lines and names extend backwards from the plane of the paper; and black lines and names protrude forwards towards the observer.





Fig. 3. Bivariate relationships of $log_{10}V_{cmax}$ versus $log_{10}J_{max}$ and $log_{10}V_{cmax}$ versus log_{10}N_{area}, within groups defined by high and low values of climate variables (3A and 3B: MI; 3C and 3D: mPAR; 3E and 3F: MAT) (n = 431). Only significant linear regressions (p < 0.05) are shown.





Fig. 4. Redundancy analysis (RDA) of nine traits constrained by climate variables

737 Mean annual temperature (MAT, °C), the square root of Moisture Index (MI, ratio of

738 precipitation to equilibrium evapotranspiration) and mean monthly photosynthetic

active radiation (mPAR, mol m⁻²) (n = 130, p < 0.05). Dynamic roles do not

participate in this RDA calculation and are shown for visual comparison only.



Fig. 5. Redundancy analysis (RDA) of nine traits constrained by dynamic roles,

defined by quantitative (5A) *versus* expert (5B) assessment (n = 130). Ellipses

represent 95% confidence intervals around the centroid of each group.



Fig. 6. Box plots showing means and standard deviation of nine traits according to the

four dynamic roles based on quantitative versus expert assessment (n = 130, p < 130

752 0.05). 'Expert' group averages of LMA, *N*_{area} land *P*_{area} are not significantly different

- (ANOVA). Dynamic roles of each trait sharing the same letter (Tukey *post hoc* test)
- are not significantly different.





Fig. 7. Bivariate relationships of V_{cmax} versus J_{max} , V_{cmax} versus N_{area} (upper panels)

and *N_{area} versus* LMA (lower panels) within dynamic role groups, according to

- 758 quantitative (left, n = 130) versus expert (right, n = 262) assessment. Significant linear
- regressions between \log_{10} -transformed variables are shown (p < 0.05).

761 **References**

- Allen, S.E., Grimshaw, H., Parkinson, J.A., Quarmby, C., 1974. Chemical analysis ofecological materials. Blackwell Scientific Publications.
- 764 Atkin, O.K., Bloomfield, K.J., Reich, P.B., Tjoelker, M.G., Asner, G.P., Bonal, D.,
- 765 Bönisch, G., Bradford, M.G., Cernusak, L.A., Cosio, E.G., Creek, D., Crous, K.Y., 766 Domingues, T.F., Dukes, J.S., Egerton, J.J.G., Evans, J.R., Farquhar, G.D., Fyllas, N.M.,
- 767 Gauthier, P.P.G., Gloor, E., Gimeno, T.E., Griffin, K.L., Guerrieri, R., Heskel, M.A.,
- 768 Huntingford, C., Ishida, F.Y., Kattge, J., Lambers, H., Liddell, M.J., Lloyd, J., Lusk,
- 769 C.H., Martin, R.E., Maksimov, A.P., Maximov, T.C., Malhi, Y., Medlyn, B.E., Meir, P.,
- 770 Mercado, L.M., Mirotchnick, N., Ng, D., Niinemets, Ü., O'Sullivan, O.S., Phillips, O.L.,
- Poorter, L., Poot, P., Prentice, I.C., Salinas, N., Rowland, L.M., Ryan, M.G., Sitch, S.,
 Slot, M., Smith, N.G., Turnbull, M.H., VanderWel, M.C., Valladares, F., Veneklaas,
- E.J., Weerasinghe, L.K., Wirth, C., Wright, I.J., Wythers, K.R., Xiang, J., Xiang, S.,
 Zaragoza-Castells, J., 2015. Global variability in leaf respiration in relation to
 climate, plant functional types and leaf traits. New Phytologist 206, 614–636.
- Atkin, O.K., Holly, C., Ball, M.C., 2000. Acclimation of snow gum (*Eucalyptus pauciflora*) leaf respiration to seasonal and diurnal variations in temperature:
 the importance of changes in the capacity and temperature sensitivity of respiration. Plant, Cell & Environment 23, 15-26.
- Baraloto, C., Timothy Paine, C., Poorter, L., Beauchene, J., Bonal, D., Domenach,
 A.M., Hérault, B., Patino, S., Roggy, J.C., Chave, J., 2010. Decoupled leaf and stem
 economics in rain forest trees. Ecology Letters 13, 1338-1347.
- Bernacchi, C.J., Pimentel, C., Long, S.P., 2003. In vivo temperature response
 functions of parameters required to model RuBP-limited photosynthesis. Plant,
 Cell & Environment 26, 1419-1430.
- Bernacchi, C.J., Singsaas, E.L., Pimentel, C., Portis Jr, A.R., Long, S.P., 2001.
 Improved temperature response functions for models of Rubisco-limited
 photosynthesis. Plant, Cell & Environment 24, 253-259.
- 789 Bloomfield, K.J., Domingues, T.F., Saiz, G., Bird, M.I., Crayn, D.M., Ford, A., Metcalfe,
- D.J., Farquhar, G.D., Lloyd, J., 2014. Contrasting photosynthetic characteristics of
 forest vs. savanna species (far North Queensland, Australia). Biogeosciences 11,
- 792 7331-7347, https://doi.org/7310.5194/bg-7311-7331-2014.
- 793 Botkin, D.B., Janak, J.F., Wallis, J.R., 1972. Some ecological consequences of a 794 computer model of forest growth. The Journal of Ecology, 849-872.
- Box, E.O., 1981. Predicting physiognomic vegetation types with climate variables.Vegetatio 45, 127-139.
- Bradford, M.G., Metcalfe, D.J., Ford, A., Liddell, M.J., McKeown, A., 2014a.
 Floristics, stand structure and aboveground biomass of a 25-ha rainforest plot in
- the wet tropics of Australia. Journal of Tropical Forest Science 26, 543-553.
- Bradford, M.G., Murphy, H.T., Ford, A.J., Hogan, D.L., Metcalfe, D.J., 2014b. Longterm stem inventory data from tropical rain forest plots in Australia. Ecology 95,
 2362-2000.
- Calvin, M., Benson, A.A., 1948. The path of carbon in photosynthesis. US AtomicEnergy Commission, Technical Information Division.
- 805 Chen, J.-L., Reynolds, J., Harley, P., Tenhunen, J., 1993. Coordination theory of leaf 806 nitrogen distribution in a canopy. Oecologia 93, 63-69.
- 807 De Kauwe, M.G., Lin, Y.-S., Wright, I.J., Medlyn, B.E., Crous, K.Y., Ellsworth, D.S.,
- 808 Maire, V., Prentice, I.C., Atkin, O.K., Rogers, A., Niinemets, Ü., Serbin, S.P., Meir, P.,
- 809 Uddling, J., Togashi, H.F., Tarvainen, L., Weerasinghe, L.K., Evans, B.J., Ishida, F.Y.,

- 810 Domingues, T.F., 2016. A test of the 'one-point method' for estimating maximum
- 811 carboxylation capacity from field-measured, light-saturated photosynthesis. New
 812 Phytologist 210, 1130-1144.
- B13 Denslow, J.S., 1987. Tropical rainforest gaps and tree species diversity. Annual
 Review of Ecology and Systematics 18, 431-451.
- 815 Dewar, R., Mauranen, A., Mäkelä, A., Hölttä, T., Medlyn, B., Vesala, T., 2018. New
- 816 insights into the covariation of stomatal, mesophyll and hydraulic conductances
- 817 from optimization models incorporating nonstomatal limitations to 818 photosynthesis. New Phytologist 217, 571-585.
- B19 Díaz, S., Cabido, M., 1997. Plant functional types and ecosystem function in
 relation to global change. Journal of Vegetation Science 8, 463-474.
- 821 Domingues, T.F., Meir, P., Feldpausch, T.R., Saiz, G., Veenendaal, E.M., Schrodt, F.,
- Bird, M., Djagbletey, G., Hien, F., Compaore, H., Diallo, A., Grace, J., Lloyd, J.O.N.,
 2010. Co-limitation of photosynthetic capacity by nitrogen and phosphorus in
- West Africa woodlands. Plant, Cell & Environment 33, 959-980.
- B25 Dong, N., Prentice, I.C., Evans, B.J., Caddy-Retalic, S., Lowe, A.J., Wright, I.J., 2017.
 B26 Leaf nitrogen from first principles: field evidence for adaptive variation with
 B27 climate. Biogeosciences 14, 481-495.
- Enquist, B.J., Bentley, L.P., 2012. Land plants: new theoretical directions and
 empirical prospects. Metabolic Ecology: A Scaling Approach. Hoboken, NJ: Wiley-
- 830 Blackwell, 164-187.
- Evans, J., 1989. Photosynthesis and nitrogen relationships in leaves of C3 plants.Oecologia 78, 9-19.
- Farquhar, G.D., von Caemmerer, S., Berry, J.A., 1980. A biochemical model of photosynthetic CO2 assimilation in leaves of C3 species. Planta 149, 78-90.
- Fisher, R., Muszala, S., Verteinstein, M., Lawrence, P., Xu, C., McDowell, N., Knox,
 R., Koven, C., Holm, J., Rogers, B., 2015. Taking off the training wheels: the
 properties of a dynamic vegetation model without climate envelopes,
 CLM4.5(ED) Geoscientific Model Development 8, 3593–3619.
- Friend, A., Schugart, H., Running, S., 1993. A physiology-based gap model offorest dynamics. Ecology 74, 792-797.
- 841 Fyllas, N., Gloor, E., Mercado, L., Sitch, S., Quesada, C., Domingues, T., Galbraith, D.,
- Torre-Lezama, A., Vilanova, E., Ramírez-Angulo, H., 2014. Analysing Amazonian
 forest productivity using a new individual and trait-based model (TFS v. 1).
- 844 Geoscientific Model Development 7, 1251-1269.
- 845 Fyllas, N.M., Patino, S., Baker, T., Bielefeld Nardoto, G., Martinelli, L., Quesada, C.,
- Paiva, R., Schwarz, M., Horna, V., Mercado, L., 2009. Basin-wide variations in
 foliar properties of Amazonian forest: phylogeny, soils and climate.
 Biogeosciences 6, 2677-2708.
- 849 Fyllas, N.M., Quesada, C.A., Lloyd, J., 2012. Deriving Plant Functional Types for
- Amazonian forests for use in vegetation dynamics models. Perspectives in Plant
 Ecology, Evolution and Systematics 14, 97-110.
- Hancock, P., Hutchinson, M., 2006. Spatial interpolation of large climate data sets
 using bivariate thin plate smoothing splines. Environmental Modelling &
 Software 21, 1684-1694.
- 855 Harris, I., Jones, P.D., Osborn, T.J., Lister, D.H., 2014. Updated high-resolution
- 856 grids of monthly climatic observations the CRU TS3.10 Dataset. International
- 857 Journal of Climatology 34, 623-642.

- Harrison, S.P., Prentice, I.C., Barboni, D., Kohfeld, K.E., Ni, J., Sutra, J.P., 2010.
 Ecophysiological and bioclimatic foundations for a global plant functional
 classification. Journal of Vegetation Science 21, 300-317.
- Haxeltine, A., Prentice, I.C., 1996. A General Model for the Light-Use Efficiency of
 Primary Production. Functional Ecology 10, 551-561.
- Hirose, T., Werger, M.J., 1987. Nitrogen use efficiency in instantaneous and daily
 photosynthesis of leaves in the canopy of a Solidago altissima stand. Physiologia
- 865 Plantarum 70, 215-222.
- Hutchinson, M., 2014a. Daily maximum precipitation: ANUClimate 1.0, 0.01
 degree, Australian Coverage, 1970-2012.
- Hutchinson, M., 2014b. Daily minimum temperature: ANUClimate 1.0, 0.01
 degree, Australian Coverage, 1970-2012.
- Hutchinson, M., 2014c. Daily precipitation: ANUClimate 1.0, 0.01 degree,
 Australian Coverage, 1970-2012.
- 872 Koch, G.W., Sillett, S.C., Antoine, M.E., Williams, C.B., 2015. Growth maximization
- trumps maintenance of leaf conductance in the tallest angiosperm. Oecologia177, 321-331.
- 875 Köppen, W.P., 1931. Grundriss der klimakunde. Walter de Gruyter Berlin.
- Kortschak, H.P., Hartt, C.E., Burr, G.O., 1965. Carbon dioxide fixation in sugarcaneleaves. Plant Physiology 40, 209.
- Langan, L., Higgins, S. I. and Scheiter, S., 2017. Climate biomes, pedo biomes
 or pyro biomes: which world view explains the tropical forest savanna
 boundary in South America? Journal of Biogeography, 44: 2319-2330.
- Lavorel, S., Garnier, E., 2002. Predicting changes in community composition and
 ecosystem functioning from plant traits: revisiting the Holy Grail. Functional
 Ecology 16, 545-556.
- Legendre, P., Anderson, M.J., 1999. Distance based redundancy analysis: testing
 multispecies responses in multifactorial ecological experiments. Ecological
 Monographs 69, 1-24.
- Li, G., Harrison, S.P., Prentice, I.C., Falster, D., 2014. Simulation of tree-ring widths
 with a model for primary production, carbon allocation, and growth.
 Biogeosciences 11, 6711-6724.
- Liddell, M., 2013a. Cape Tribulation OzFlux tower site OzFlux: Australian and
 New Zealand Flux Research and Monitoring TERN hdl: 102.100.100/14242.
- Liddell, M., 2013b. Robson Creek OzFlux tower site OzFlux: Australian and New
 Zealand Flux Research and Monitoring. TERN hdl: 102.100.100/14243.
- Lin, Y.-S., Medlyn, B.E., De Kauwe, M.G., Ellsworth, D.S., 2013. Biochemical photosynthetic responses to temperature: how do interspecific differences compare with seasonal shifts? Tree Physiology 33, 793-806.
- Lin, Y.-S., Medlyn, B.E., Duursma, R.A., Prentice, I.C., Wang, H., Baig, S., Eamus, D.,
- de Dios, V.R., Mitchell, P., Ellsworth, D.S., de Beeck, M.O., Wallin, G., Uddling, J.,
- 899 Tarvainen, L., Linderson, M.-L., Cernusak, L.A., Nippert, J.B., Ocheltree, T.W.,
- 900 Tissue, D.T., Martin-StPaul, N.K., Rogers, A., Warren, J.M., De Angelis, P., Hikosaka,
- 901 K., Han, Q., Onoda, Y., Gimeno, T.E., Barton, C.V.M., Bennie, J., Bonal, D., Bosc, A.,
- Low, M., Macinins-Ng, C., Rey, A., Rowland, L., Setterfield, S.A., Tausz-Posch, S.,
 Zaragoza-Castells, J., Broadmeadow, M.S.J., Drake, J.E., Freeman, M., Ghannoum,
- 903 Zaragoza-Castells, J., Broadmeadow, M.S.J., Drake, J.E., Freeman, M., Ghannoum, 904 O., Hutley, L.B., Kelly, J.W., Kikuzawa, K., Kolari, P., Koyama, K., Limousin, J.-M.,
- 905 Meir, P., Lola da Costa, A.C., Mikkelsen, T.N., Salinas, N., Sun, W., Wingate, L., 2015.

- 906 Optimal stomatal behaviour around the world. Nature Climate Change 5, 459-907 464.
- 908 Lloyd, J., Bloomfield, K., Domingues, T.F., Farquhar, G.D., 2013. Photosynthetically
- 909 relevant foliar traits correlating better on a mass vs an area basis: of
 910 ecophysiological relevance or just a case of mathematical imperatives and
 911 statistical quicksand? New Phytologist 199, 311-321.
- 912 Lombardozzi, D.L., Smith, N.G., Cheng, S.J., Dukes, J.S., Sharkey, T.D., Rogers, A.,
 913 Fisher, R. and Bonan, G.B., 2018. Triose phosphate limitation in photosynthesis
- 914 models reduces leaf photosynthesis and global terrestrial carbon storage.
 915 Environmental Research Letters 13(7), 074025.
- 916 Maire, V., Martre, P., Kattge, J., Gastal, F., Esser, G., Fontaine, S., Soussana, J.-F.,
- 2012. The coordination of leaf photosynthesis links C and N fluxes in C3 plantspecies. PLoS ONE 7, e38345.
- 919 Medlyn, B.E., Duursma, R.A., Eamus, D., Ellsworth, D.S., Prentice, I.C., Barton,
- 920 C.V.M., Crous, K.Y., De Angelis, P., Freeman, M., Wingate, L., 2011. Reconciling the
- 921 optimal and empirical approaches to modelling stomatal conductance. Global922 Change Biology 17, 2134-2144.
- Medvigy, D., Wofsy, S.C., Munger, J.W., Hollinger, D.Y., Moorcroft, P.R., 2009.
 Mechanistic scaling of ecosystem function and dynamics in space and time:
 Ecosystem Demography model version 2. Journal of Geophysical Research:
 Biogeosciences 114, 0148-0227.
- Meng, T., Wang, H., Harrison, S., Prentice, I., Ni, J., Wang, G., 2015. Responses of
 leaf traits to climatic gradients: adaptive variation versus competitional shifts.
 Biogeosciences 12, 5339-5352.
- 930 Miyazawa, Y., Kikuzawa, K., 2006. Physiological basis of seasonal trend in leaf
- 931 photosynthesis of five evergreen broad-leaved species in a temperate deciduous932 forest. Tree Physiology 26, 249-256.
- Moorcroft, P.R., Hurtt, G.C., Pacala, S.W., 2001. A method for scaling vegetation
 dynamics: The Ecosystem Demography Model (ED). Ecological Monographs 71,
 557-586.
- Niinemets, Ü., Tenhunen, J.D., 1997. A model separating leaf structural and
 physiological effects on carbon gain along light gradients for the shade-tolerant
 species Acer saccharum. Plant, Cell & Environment 20, 845-866.
- 939 Oksanen, J., Blanchet, F.G., Kindt, R., Legendre, P., Minchin, P.R., O'Hara, R.,
- Simpson, G.L., Solymos, P., Stevens, M.H.H., Wagner, H., 2015. Package 'vegan'.
 Community ecology package, version, 2.2-1.
- 942 Olson, M.E., Anfodillo, T., Rosell, J.A., Petit, G., Crivellaro, A., Isnard, S., León -
- 943 Gómez, C., Alvarado Cárdenas, L.O., Castorena, M., 2014. Universal hydraulics of
 944 the flowering plants: vessel diameter scales with stem length across angiosperm
- 945 lineages, habits and climates. Ecology Letters 17, 988-997.
- 946 Pavlick, R., Drewry, D.T., Bohn, K., Reu, B., Kleidon, A., 2013. The Jena Diversity-
- 947 Dynamic Global Vegetation Model (JeDi-DGVM): a diverse approach to 948 representing terrestrial biogeography and biogeochemistry based on plant 949 functional trade-offs. Biogeosciences 10, 4137-4177.
- 950 Peres-Neto, P.R., Legendre, P., Dray, S., Borcard, D., 2006. Variation partitioning
- 951 of species data matrices: estimation and comparison of fractions. Ecology 87,952 2614-2625.
- 953 Prentice, I., Leemans, R., 1990. Pattern and process and the dynamics of forest 054 ctructure: a cimulation approach. The Journal of Ecology 240, 255
- structure: a simulation approach. The Journal of Ecology, 340-355.

- 955 Prentice, I.C., Bondeau, A., Cramer, W., Harrison, S.P., Hickler, T., Lucht, W., Sitch,
- 956 S., Smith, B., Sykes, M.T., 2007. Dynamic global vegetation modeling: quantifying
- 957 terrestrial ecosystem responses to large-scale environmental change, Terrestrial
- 958 ecosystems in a changing world. In: Canadell J.G., Pataki D.E., Pitelka L.F. (eds)
 959 Terrestrial Ecosystems in a Changing World. Global Change The IGBP Series.
 960 Springer, Berlin, Heidelberg, pp. 175-192.
- 961 Prentice, I.C., Cowling, S.A., 2013. Dynamic global vegetation models, in: Levin,
 962 S.A., Academic Press (Ed.), Encyclopedia of Biodiversity, 2nd edition ed, pp. 607963 689.
- Prentice, I.C., Cramer, W., Harrison, S.P., Leemans, R., Monserud, R.A., Solomon,
 A.M., 1992. A global biome model based on plant physiology and dominance, soil
 properties and climate. Journal of Biogeography 19, 117-134.
- Prentice, I.C., Dong, N., Gleason, S.M., Maire, V., Wright, I.J., 2014. Balancing the
 costs of carbon gain and water transport: testing a new theoretical framework
 for plant functional ecology. Ecology Letters 17, 82-91.
- 970 Prentice, I.C., Liang, X., Medlyn, B.E., Wang, Y.P., 2015. Reliable, robust and
 971 realistic: the three R's of next-generation land-surface modelling. Atmospheric
 972 Chemistry and Physics 15, 5987-6005.
- 973 Prentice, I.C., Liddell, M., Furstenau Togashi, H., Atkin, O., Weerasinghe, L., 2013.
- 974 Leaf Level Physiology, Chemistry and Structural Traits, Far North Queensland
 975 SuperSite, Robson Creek, 2012. TERN Australian SuperSite Network.
 976 http://portal.tern.org.au/leaf-level-physiology-creek-2012 12.
- 977 Prentice, I.C., Meng, T., Wang, H., Harrison, S.P., Ni, J., Wang, G., 2011. Evidence of
 978 a universal scaling relationship for leaf CO₂ drawdown along an aridity gradient.
 979 New Phytologist 190, 169-180.
- Prentice, I.C., Sykes, M.T., Cramer, W., 1993. A simulation model for the transient
 effects of climate change on forest landscapes. Ecological Modelling 65, 51-70.
- Purves, D.W., Lichstein, J.W., Strigul, N., Pacala, S.W., 2008. Predicting and
 understanding forest dynamics using a simple tractable model. Proceedings of
 the National Academy of Sciences 105, 17018.
- 985 Quesada, C.A., Phillips, O.L., Schwarz, M., Czimczik, C.I., Baker, T.R., Patino, S.,
- Fyllas, N.M., Hodnett, M.G., Herrera, R., Almeida, S., Alvarez Dávila, E., Arneth, A.,
 Arroyo, L., Chao, K. J., Dezzeo, N., Erwin, T., di Fiore, A., Higuchi, N., Honorio
- 988 Coronado, E., Jimenez, E. M., Killeen, T., Lezama, A. T., Lloyd, G., López-González,
- 989 G., Luizão, F. J., Malhi, Y., Monteagudo, A., Neill, D. A., Núñez Vargas, P., Paiva, R.,
- 990 Peacock, J., Peñuela, M. C., Peña Cruz, A., Pitman, N., Priante Filho, N., Prieto, A.,
- Ramírez, H., Rudas, A., Salomão, R., Santos, A. J. B., Schmerler, J., Silva, N., Silveira,
 M., Vásquez, R., Vieira, I., Terborgh, J., and Lloyd, J., 2012. Basin-wide variations
- in Amazon forest structure and function are mediated by both soils and climate.Biogeosciences 9, 2203-2246.
- R Core Team, 2012. R: A language and environment for statistical computing. R
 Foundation for Statistical Computing, Vienna, Austria. <u>http://www.R-</u>
 project.org/.
- Ranson, S.L., Thomas, M., 1960. Crassulacean Acid Metabolism. Annual Review ofPlant Physiology 11, 81-110.
- 1000 Raunkiær, C., 1934. The Life Forms of Plants and Statistical Plant Geography.1001 Clarendon Press.
- 1002 Reich, P.B., 2014. The world-wide 'fast–slow' plant economics spectrum: a traits
- 1003 manifesto. Journal of Ecology 102, 275-301.

- 1004 Reich, P.B., Walters, M.B., Ellsworth, D.S., 1997. From tropics to tundra: global
 1005 convergence in plant functioning. Proceedings of the National Academy of
 1006 Sciences 94, 13730-13734.
- Reid, D.E.B., Silins, U., Mendoza, C., Lieffers, V.J., 2005. A unified nomenclature for
 quantification and description of water conducting properties of sapwood xylem
 based on Darcy's law. Tree Physiology 25, 993-1000.
- 1010 Sakschewski, B., Bloh, W., Boit, A., Rammig, A., Kattge, J., Poorter, L., Peñuelas, J.,
- 1011 Thonicke, K., 2015. Leaf and stem economics spectra drive diversity of functional 1012 plant traits in a dynamic global vegetation model. Global Change Biology 21,
- 1013 2711-2725.
- Scheiter, S., Langan, L., Higgins, S.I., 2013. Next generation dynamic global
 vegetation models: learning from community ecology. New Phytologist 198, 957969.
- 1017 Sharkey, T.D., Bernacchi, C.J., Farquhar, G.D., Singsaas, E.L., 2007. Fitting
 1018 photosynthetic carbon dioxide response curves for C3 leaves. Plant, Cell &
 1019 Environment 30, 1035-1040.
- 1020 Shugart, H.H., 1984. A Theory of Forest Dynamics. Springer-Verlag, New York.
- 1021 Sitch, S., Smith, B., Prentice, I.C., Arneth, A., Bondeau, A., Cramer, W., Kaplan, J.O.,
- Levis, S., Lucht, W., Sykes, M.T., Thonicke, K., Venevsky, S., 2003. Evaluation of
 ecosystem dynamics, plant geography and terrestrial carbon cycling in the LPJ
 dynamic global vegetation model. Global Change Biology 9, 161-185.
- Smith, B., Prentice, I.C., Sykes, M.T., 2001. Representation of vegetation dynamics
 in the modelling of terrestrial ecosystems: comparing two contrasting
 approaches within European climate space. Global Ecology and Biogeography 10,
 621-637.
- Sperry, John S., 2003. Evolution of water transport and xylem structure.International Journal of Plant Sciences 164, S115-S127.
- Swaine, M., Whitmore, T., 1988. On the definition of ecological species groups intropical rain forests. Vegetatio 75, 81-86.
- 1033 Ter Braak, C.J., Prentice, I.C., 1988. A theory of gradient analysis. Advances in1034 Ecological Research 18, 271-317.
- 1035 Togashi, H.F., Prentice, I.C., Atkin, O.K., Macfarlane, C., Prober, S., Bloomfield, K.J.,
- Evans, B., 2017. Thermal acclimation of leaf photosynthetic traits in an evergreen
 woodland, consistent with the co-ordination hypothesis. Biogeosciences Discuss.,
- 1038 https://doi.org/10.5194/bg-2017-449, in review, 2017.
- Togashi, H.F., Prentice, I.C., Evans, B.J., Forrester, D.I., Drake, P., Feikema, P.,
 Brooksbank, K., Eamus, D., Taylor, D., 2015. Morphological and moisture
 availability controls of the leaf area-to-sapwood area ratio: analysis of
 measurements on Australian trees. Ecology and Evolution 5, 1263-1270.
- 1043 Turner, I.M., 2001. The Ecology of Trees in the Tropical Rain Forest. Cambridge1044 University Press.
- 1045 Tyree, M.T., Ewers, F.W., 1991. The hydraulic architecture of trees and other 1046 woody plants. New Phytologist 119, 345-360.
- 1047 van Bodegom, P.M., Douma, J.C., Verheijen, L.M., 2014. A fully traits-based
- approach to modeling global vegetation distribution. Proceedings of the NationalAcademy of Sciences 111, 13733.
- 1050 Verheijen, L., Brovkin, V., Aerts, R., Bönish, G., Cornelissen, J., Kattge, J., Reich, P.,
- 1051 Wright, I., Van Bodegom, P., 2013. Impacts of trait variation through observed

- 1052 trait-climate relationships o performance of a representative Earth System1053 Model: a conceptual analysis. Biogeosciences 10, 5497-5515.
- Wang, H., Harrison, S.P., Prentice, I.C., Yang, Y., Bai, F., Togashi, H.F., Wang, M.,
 Zhou, S., Ni, J., 2018. The China Plant Trait Database: toward a comprehensive
 regional compilation of functional traits for land plants. Ecology 99, 500-500.
- 1057 Wang, H., Prentice, I.C., Keenan, T.F., Davis, T.W., Wright, I.J., Cornwell, W.K.,
- 1057 Wang, H., Pfentice, I.C., Keenan, T.F., Davis, T.W., Wright, I.J., Cornwell, W.K.,
 1058 Evans, B.J., Peng, C., 2017. Towards a universal model for carbon dioxide uptake
 1059 by plants. Nature Plants 3, 734-741.
- 1060 Warton, D.I., Wright, I.J., Falster, D.S., Westoby, M., 2006. Bivariate line fitting
 1061 methods for allometry. Biological Reviews 81, 259-291.
- Weerasinghe, L.K., Creek, D., Crous, K.Y., Xiang, S., Liddell, M.J., Turnbull, M.H.,
 Atkin, O.K., 2014. Canopy position affects the relationships between leaf
 respiration and associated traits in a tropical rainforest in Far North Queensland.
 Tree Physiology 34, 564-584.
- 1066 Whitehead, D., Edwards, W.R.N., Jarvis, P.G., 1984. Conducting sapwood area, 1067 foliage area, and permeability in mature trees of *Picea sitchensis* and *Pinus* 1068 *contorta*. Canadian Journal of Forest Research 14, 940-947.
- 1069 Whitmore, T., 1982. On pattern and process in forests. In: Newman, E. I. (ed.), 1070 The plant community as a working mechanism. pp. 45–59. Blackwell, Oxford.
- 1071 Wickham, H., 2010. ggplot2: Elegant Graphics for Data Analysis. Springer.
- 1072 Woodward, F.I., 1987. Climate and Plant Distribution. Cambridge University1073 Press.
- 1074 Wright, I.J., Reich, P.B., Westoby, M., Ackerly, D.D., Baruch, Z., Bongers, F.,
- 1075 Cavender-Bares, J., Chapin, T., Cornelissen, J.H.C., Diemer, M., Flexas, J., Garnier,
- 1076 E., Groom, P.K., Gulias, J., Hikosaka, K., Lamont, B.B., Lee, T., Lee, W., Lusk, C.,
- 1077 Midgley, J.J., Navas, M.-L., Niinemets, U., Oleksyn, J., Osada, N., Poorter, H., Poot, P.,
- Prior, L., Pyankov, V.I., Roumet, C., Thomas, S.C., Tjoelker, M.G., Veneklaas, E.J.,
 Villar, R., 2004. The worldwide leaf economics spectrum. Nature 428, 821-827.
- 1080 Yang, Y. , Wang, H. , Harrison, S.P., Prentice, I.C., Wright, I.J., Peng, C., Lin, G., 2018.
- 1081 Quantifying leaf trait covariation and its controls across climates and biomes.
 1082 New Phytologist (online). doi:10.1111/nph.15422
- 1083 Zhou, S., Duursma, R.A., Medlyn, B.E., Kelly, J.W. and Prentice, I.C., 2013. How
- 1084 should we model plant responses to drought? An analysis of stomatal and non-
- 1085 stomatal responses to water stress. Agricultural and Forest Meteorology 182, 1086 204-214.