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# Stem diameter growth rates in a fire-prone savanna correlate with photosynthetic rate and branch-scale biomass allocation, but not specific leaf area

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#### 1 ABSTRACT

2 Plant growth rates strongly determine ecosystem productivity and are a central element of plant ecological strategies. For laboratory and glasshouse-grown seedlings, specific leaf area 3 4 (SLA; ratio of leaf area to mass) is a key driver of interspecific variation in growth rate (GR). 5 Consequently, SLA is often assumed to drive GR variation in field-grown adult plants. 6 However, there is increasing evidence that this is not the general case. This suggests that GR 7 - SLA relationships (and perhaps those for other traits) may vary depending on the age or 8 size of the plants being studied. Here we investigated GR – trait relationships and their size 9 dependence among 17 woody species from an open-canopy, fire-prone savanna in northern 10 Australia. We tested the predictions that SLA and stem diameter growth rate would be 11 positively correlated in saplings but unrelated in adults while, in both age classes, faster-GR 12 species would have higher light-saturated photosynthetic rate (A<sub>sat</sub>), higher leaf nutrient 13 concentrations, higher branch-scale biomass allocation to leaf versus stem tissues, and lower 14 wood density (WD). SLA showed no relationship to stem diameter GR, even in saplings, and 15 the same was true of leaf N and P concentrations, and WD. However, branch-scale leaf:stem 16 allocation was strongly related to GR in both age groups, as was Asat. Together, these two 17 traits accounted for up to 80% of interspecific variation in adult GR, and 41% of sapling GR. 18 A<sub>sat</sub> is rarely measured in field-based GR studies, and this is the first report of branch-scale 19 leaf:stem allocation (analogous to a benefit:cost ratio) in relation to plant growth rate. Our 20 results suggest that we may yet find general trait-drivers of field growth rates, but SLA will not be one. 21

22

23 **Keywords**: relative growth rate, specific leaf area, plant functional traits, leaf:stem

24 allocation, leaf economic spectrum

#### 25 INTRODUCTION

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32 variation. Although other formulations are possible (Cernusak *et al.* 2008; Enquist *et al.* 

Growth rate differences between species are critical in determining the outcome of

competition, while forest, shrubland and grassland yields - and carbon sequestration - are

driven by growth rates. Since the 1970s, seedling relative growth rate (RGR; the dry mass

increase per unit dry mass per unit time) has been treated as a key element of plant ecological

strategies (Grime and Hunt 1975; Grime et al. 1997; Lambers and Poorter 1992; Lambers et

al. 1998). A chief focus in that literature has been on identifying the key drivers of RGR

33 2007; Lambers and Poorter 1992), RGR is most commonly mathematically decomposed as

34 follows:

35 
$$RGR = NAR \times SLA \times LMF$$
; (Eqn 1)

36 where NAR is net assimilation rate (dry mass growth rate per unit leaf area), LMF is leaf 37 mass fraction (leaf dry mass/plant dry mass), and SLA is specific leaf area (leaf area/leaf dry 38 mass). A meta-analysis of 111 studies concerning herbaceous species found SLA to account 39 for 64 % of RGR variation, whereas NAR and LMF accounted for just 26 % and 11 %, 40 respectively (Poorter and van der Werf 1998). For woody species, meta-analyses have shown 41 LMF has only a weak influence on RGR, with NAR and SLA being the dominant terms 42 (Shipley 2006; Veneklaas and Poorter 1998). For seedlings, NAR variation is thought to 43 largely reflect photosynthetic rate; for older plants, other factors presumably become 44 increasingly influential on NAR, most especially carbon losses from tissue turnover, tissue respiration and root exudates (Konings 1989; Li et al. 2016; Poorter and Garnier 2007). 45

But do high SLA species achieve faster growth rates also as adults? Intuitively one
might think so. After all, high SLA species typically have high leaf N and P concentrations
and various other traits indicative of a "fast" lifestyle (Reich 2014). However, it seems

49 increasingly apparent that SLA and field-measured growth rates are most commonly 50 unrelated (Aiba and Nakashizuka 2009; Coomes and Grubb 1998; Easdale and Healey 2009; 51 Gower et al. 1993; Hérault et al. 2011; Martínez-Vilalta et al. 2010; Paine et al. 2015; 52 Poorter et al. 2008; Rüger et al. 2012). In only a minority of cases – mostly for saplings or 53 small trees – has a positive SLA-growth relationship been reported and even then, only 54 weakly (Poorter and Bongers 2006; Prior et al. 2004; Reich et al. 1992; Rossatto et al. 2009; 55 Wright *et al.* 2010). This raises the question, how is it that, through ontogeny – or with 56 increasing plant size – variation in SLA becomes less strongly correlated with plant growth 57 rate? This is an important question, with species-dimensions running from slow to fast growth rates, and low to high SLA, being widely accepted as pivotal in plant ecological 58 59 strategies (Grime et al. 1997; Reich 2014; Westoby et al. 2002).

60 One potential explanation is that low SLA species accumulate more massive canopies 61 over time (because of their longer leaf lifespans), and this counteracts their slower per-gram 62 metabolic rates such that annual productivity can be as high as that of a high SLA species 63 (Chabot and Hicks 1982; Matyssek 1986; Reich et al. 1992). Another potential explanation 64 was outlined by Gibert et al. (2016). In their first-principles plant growth model, SLA always 65 had a positive influence on growth rate but, as plants grow, this effect becomes increasingly 66 masked by leaf turnover costs (which are higher in high SLA species, because of short leaf 67 lifespans) and by increasing sapwood respiration costs – because larger plants have relatively 68 more sapwood (King 1999; Poorter et al. 2012). Other predictions from this model were that, 69 irrespective of plant size, light-saturated photosynthetic rate (Asat) should correlate positively 70 with growth rate, and wood density negatively. Meta-analysis across a range of trait – growth 71 studies showed broad support for these predictions, though low sample size (number of 72 studies) was a recurring problem for the analysis (Gibert et al. 2016).

73 Considering plant growth as an outcome of the balance between sapwood respiration 74 and leaf photosynthesis is not new. For example, slower growth rates in larger (older) trees is 75 likely a result of whole-plant sapwood volume increasing more rapidly over time than canopy 76 leaf area (Ryan 1989). Of particular interest for the present study, one can make a related 77 argument at branch-scale. That is, species (or indeed, individuals) deploying more leaf 78 relative to stem wood and bark on terminal branches (i.e., with higher branch-scale LMF) 79 should – all else equal – achieve faster whole-plant growth rates (Pickup et al. 2005). This 80 intriguing proposition remains untested.

81 In this study we investigated the size-dependence of relationships between growth 82 rate and several key plant functional traits (including branch-scale leaf:stem allocation), for a 83 range of woody species from a fire-prone savanna in northern Australia. We chose this 84 vegetation type because we expected the predicted relationships to be most clearly expressed 85 in a situation where competition for light is minimal, and potential photosynthetic benefits are 86 more likely to be realised. For each species we sampled traits both on small-DBH individuals 87 and on larger-DBH individuals (henceforth called "saplings" and "adults"), and matched the 88 trait data to previously reported long-term, stem-diameter growth rate data, re-calculated 89 separately for small and large individuals. We tested the following predictions: (1) The GR – 90 SLA relationship would be positive in saplings but absent among adult plants; (2) Asat 91 (expressed per unit leaf area) would explain substantial variation in GR, both in saplings and 92 adults; (3) Trends in leaf N and P would mirror those in Asat (e.g., because N-rich proteins 93 and P-rich energetic molecules play key roles in plant metabolism); (4) Sapwood tissue 94 density would be negatively related to GR both in saplings and adults; (5) Branch-scale 95 leaf:stem biomass allocation would correlate positively with GR, both in saplings and adults.

96

#### 97 MATERIALS AND METHODS

## 98 Trait dataset: site and species selection

99 We sampled species at Howard Springs Nature Reserve, 30 km east from Darwin. The 100 savanna vegetation there is typical of the region, with an overstory dominated by eucalypts 101 and an understory dominated by C4 grasses. Overstory leaf area index ranges between about 102 0.6 in the dry season to 1 in the wet season (Hutley et al. 2001). At Darwin airport (20 km to 103 the west), long-term annual rainfall is 1736 mm, typically with > 95% of rain falling during 104 the wet season (October – April). Mean annual temperature is 27.6 °C (data from 105 www.bom.gov.au). Fires occur regularly in the dry season; typical fire return intervals in the 106 region are 1 - 3 years (Russell-Smith *et al.* 2003). Soils at the site are sandy and low in 107 nutrients: mean (and standard deviation) nutrient concentrations in eight soil samples (0-20 108 cm depth) collected in September 2010 were as follows: total C = 2.7 % (1.8), total N = 0.093% (0.074), total P = 72.7 mg kg<sup>-1</sup> (22.6). 109

110 Deciduous, semi-deciduous and evergreen species are all present in this savanna. 111 Hence, we sampled plant traits at two time points: at the end of one dry season (September 112 2010), when deciduous species are leafless and the canopies of semi-deciduous species are at 113 their thinnest; and right at the end of the subsequent wet season (early May 2011), when most 114 species still tend to have full or nearly-full canopies (Williams et al. 1997). We selected 17 115 species to study, based on available growth rate data and their availability across two nearby 116 sub-sites (12°27'10" S, 131°6'30" E; 12°27'57" S, 131°6'51" E). This species-set represents a range of functional types (trees and shrubs; nitrogen-fixers and non-fixers; deciduous, semi-117 118 deciduous and evergreen species; Table 1). We used diameter at breast height (DBH) to identify "adults" versus "saplings" for each species, but we use these terms informally (i.e., 119 120 not necessarily reflecting degree of reproductive maturity), and synonymously with "small

121 individuals" and "large individuals" of each species (Prior et al. 2006). For species that are 122 large trees at maturity we used a DBH cut-off of 6 cm to distinguish saplings from adults; for 123 smaller trees and shrub species we used a 3 cm cut-off, with the exception of the small, short-124 lived and fast growing species Acacia difficilis, for which we used a cut-off of 1.5 cm DBH 125 (Table 1). When choosing plants to sample we prioritised individuals of each species which 126 clearly fell into these size-defined 'sapling' and 'adult' categories, so as to minimise ambiguity in this regard. For each age/size class of each species we aimed to make trait 127 128 measurements on each of five individuals.

129

# 130 Leaf traits

Five recently-matured, fully-expanded and undamaged leaves (including petioles) were collected from each individual for determination of one-sided projected leaf area (flatbed scanner), oven-dried mass (70°C for at least 48 hours), and thus SLA (area per dry mass; cm<sup>2</sup> g<sup>-1</sup>). Total N concentration of leaf and soil samples was measured with a LECO TruSpec CHN analyser; total P by ICP-OES, on nitric acid digests (analyses run at Appleton Lab, University of Queensland). Area-based leaf N and P concentrations were calculated from mass-based concentrations and the mean SLA value for each species/age class.

Light-saturated rates of photosynthesis ( $A_{sat}$ ) and stomatal conductance to water vapour ( $g_s$ ) were measured during the wet season only, using a Li-Cor 6400XT portable infra-red gas analyser (5-8 replicate plants per species/age class). These measurements were made on leaves held on detached branches, > 1 m in length; branch-ends were re-cut and kept in water post-harvest until photosynthesis was measured (within 5 minutes of harvesting). Cuvette block temperature was kept at 26-27°C, reference CO<sub>2</sub> was set to 400 ppm and cuvette photosynthetic photon flux density was maintained at 2000 µmol m<sup>-2</sup> s<sup>-1</sup>. Cuvette

145	vapour pressure deficit was only loosely controlled, averaging 1.22 kPa across all
146	measurements (standard deviation 0.28). Leaf lamina material used in photosynthetic
147	measurements was oven-dried and pooled per species/age class, then analysed for ${}^{13}C/{}^{12}C$
148	stable isotope composition (hereafter " $\delta^{13}$ C") at the Stable Isotopes Laboratory, Australian
149	National University, Canberra. The $\delta^{13}$ C provides an integrated measure of the extent of CO <sub>2</sub>
150	drawdown during photosynthesis (Farquhar <i>et al.</i> 1982). We also measured $\delta^{13}$ C on
151	representative whole-leaf (including petiole) samples collected during September 2010 (dry
152	season).

153

#### 154 Branch traits

155 A single terminal branch, 80 cm in length, was sampled from each of several individuals of 156 each species/age class (average, 4.8 branches per species/age). These were divided into 157 segments cut at 5, 10, 20, 40 and 80 cm from the terminal end, and oven-dried at 70°C for at 158 least 5 days. For each 80 cm branch (and including material on any side-branches) we 159 calculated (1) total leaf dry mass, (2) total stem dry mass, (3) the ratio of leaf:stem dry mass 160 for this 80cm segment (hereafter LM:SM), (4) the ratio of leaf area: stem dry mass (by 161 multiplying leaf mass by the appropriate SLA value; hereafter LA:SM), and (5) the allometric 162 slope describing leaf mass accumulation versus stem mass accumulation (hereafter 163 a LMSM). This slope was estimated for accumulations from the tip going back down the 164 stem, expressed at the 5, 10, 20, 40 and 80 cm sampling points. These allometries were 165 calculated as standardised major axis slopes (Warton et al. 2006) fitted to log-transformed data. The "static ratio" (LM:SM, LA:SM) and allometric (a LMSM) descriptors were of 166 course correlated, but sufficiently weakly (mean  $r^2$  across age/season datasets = 0.37) that 167 they contained substantial independent information about branch-scale leaf:stem allocation. 168

169 Stem tissue density (dry mass per fresh volume) was measured on 2 cm long branch 170 segments of approximately 1 cm over-bark diameter. For dry season samples, density was 171 measured on whole stems, including bark. For wet season samples, the bark was first 172 removed, allowing us to measure relative dry mass allocation to bark versus sapwood. For 173 these samples, stem density refers to that of the sapwood only (plus any pith, if present). 174 Hereafter we refer to these quantities as "WD" (wood density). At each date, five samples 175 were taken per species/age class. Sample volumes were measured using standard procedure 176 (via displacement), and dry masses after oven-drying at 70°C for seven days.

177

178 *Growth rates* 

179 The stem increment (growth rate) data were not measured as part of this study. Data for 180 northern Australian savanna species came from two sources: (1) The "Kapalga" dataset; and 181 (2) the "Three Parks" dataset (Murphy et al. 2010). Kapalga (12°50'S, 132°50'E) is located 182 200 km east of Darwin in Kakadu National Park. It was run as a CSIRO research station from 183 the mid-1970s to the mid-1990s. Various combinations of fire frequency and severity were 184 generated between 1990 and 1995 (Cook and Corbett 2003; Williams et al. 2003). Stem 185 diameters at 1.3 m height (DBH) were measured every 12 months. The "Three Parks" dataset 186 contains repeat-measured DBH data for savanna vegetation in Kakadu, Litchfield and 187 Nitmiluk National Parks (Northern Territory, Australia). At each of 163 locations, all 188 individuals with DBH  $\geq$  5 cm were initially measured between 1994 and 1997, then re-189 measured twice, each time approximately five years apart. DBH measurements were made 190 during the wet season only (to avoid the stem shrinkage that may occur during dry 191 conditions). Fire severity and frequency was determined for each location/census period post 192 hoc, using survey data and aerial photographs (Murphy et al. 2010). Annual rainfall ranges 193 from 900 mm at Nitmiluk to 1470 mm at Kakadu (including Kapalga), with sites showing

similar seasonality (Murphy *et al.* 2010). Murphy et al (2010) – one of our two sources of
growth rate data – showed that DBH growth rate was unrelated to rainfall across this region,
suggesting that the GR data should be broadly representative of our study species, despite
Howard Springs receiving higher rainfall.

198 In savanna, fire damage to stems may result in very low or even negative stem 199 diameter increments between censuses. We took three steps to counter this issue. First, we 200 discarded all DBH data from sites designated as having experienced "severe" (Kapalga) or 201 "frequent, severe" (Three Parks) fire. Second, all negative increment data were discarded. 202 This resulted in a dataset with 7897 rows, each row describing a stem diameter increment for 203 some individual plant, for some census period. (Data from Kapalga contributed 5155 rows; 204 from Three Parks, 1340 rows from Kakadu NP, 765 from Litchfield NP, 637 from Nitmiluk 205 NP). Finally, we summarised the set of stem increment data for each species/age class using 206 80<sup>th</sup> percentiles, rather than a measure of central tendency (Clark and Clark 1999; Rozendaal 207 et al. 2010; Wright et al. 2010). Doing so should minimise the influence of fire-damaged 208 plants and better facilitate quantifying trait-growth relationships of "successful" individuals 209 of each species and age class (by analogy, we measured traits such as photosynthetic rates on 210 recently-matured leaves which are near their peak physiological condition). There were too 211 few data to confidently estimate GR for saplings of *Acacia difficilis* and *Grevillea decurrens*, 212 meaning that we had GR data for 17 species as adults, and 15 species as saplings. Trait data 213 were averaged to give a single value for saplings and for adults, for each species, for each 214 sampling period (Table S1; Figs S1, S2). Almost all individuals of the two deciduous species 215 were leafless during dry season sampling, limiting analyses of GR-trait relationships for that 216 period to 16 species as adults and 13 species as saplings (Table 2).

In this study we were working across species with very different sizes at maturity.Consequently we chose to use absolute rather than relative growth rate as our preferred

growth index, relative growth rate being strongly size-dependent (Iida *et al.* 2014; Rees *et al.*2010), systematically decreasing with increasing plant size. By contrast, absolute growth
rates are generally relatively stable across broad ranges in plant size (Prior *et al.* 2006; Prior *et al.* 2004).

223

224 Analyses

Ordinary least squares regression (including multiple regressions, with interaction terms) was used to quantify relationships between GR and the various plant traits. For key bivariate relationships we tested for heterogeneity among regression slopes fitted to each age-season group. Where deemed non-heterogeneous (slope test, P > 0.05) a common slope can be fitted, and whole-model explanatory power ( $r^2$ ) and statistical significance can be reported. All analyses were run in IBM SPSS Statistics v. 22, with the General Linear Model module used for testing slope heterogeneity and calculating descriptors of common slopes.

An additional analysis specified by a reviewer and editor is reported in Supplementary Appendix 1. There we report results from an analysis of phylogenetically independent contrasts (PICs) calculated for GR in relation to key traits, for the wet season - adult data subset only (this subset including all 17 species, and all traits including photosynthetic rates).

236

## 237 **RESULTS**

Sapling GR varied *ca*. 3-fold, from 0.37 cm yr<sup>-1</sup> (*Corymbia bleeseri*) to 1.0 cm yr<sup>-1</sup> (*Grevillea pteridifolia*), median = 0.50 cm yr<sup>-1</sup> (Table S1). Adult GR varied *ca*. 6-fold, from 0.25 cm yr<sup>-1</sup> (*Persoonia falcata*) to 1.56 cm yr<sup>-1</sup> (*Acacia difficilis*, for which we lacked data for saplings), median 0.42 cm yr<sup>-1</sup>. Leaf-level traits (Table S1) varied 9-fold or less among species. By

contrast, leaf mass on terminal branches varied 16-fold among species, wood mass 34-fold,and their ratio, 17-fold.

244

### 245 Growth Rate – leaf trait relationships

SLA was unrelated to GR in any of the four comparisons (two age classes, two sampling 246 periods; all  $r^2 \le 0.03$ , P > 0.4; Fig. 1a, Table 2). Leaf N and P, expressed either per mass or 247 per area, were also unrelated to GR in every case (all  $r^2 \le 0.15$ , with most  $r^2 \le 0.07$ , and all P 248 > 0.14; Table 2). By contrast, A<sub>sat</sub> explained a substantial proportion of growth rate variation 249 in adults ( $r^2 = 0.32$ ; P = 0.017) as well as saplings ( $r^2 = 0.23$ ; P = 0.071; Fig. 1b). The 250 common fitted slope of  $\log_{10}$ GR on  $\log_{10}A_{sat}$  was 1.0 (whole-model  $r^2 = 0.31$ ), indicating 251 252 direct proportionality: on average, a two-fold increase in A<sub>sat</sub> corresponded to a two-fold 253 increase in stem diameter growth rate.

Variation in A<sub>sat</sub> can be underpinned by variation in stomatal conductance to water (g<sub>s</sub>) and by the extent of CO<sub>2</sub> drawdown during photosynthesis, indexed here via leaf  $\delta^{13}$ C. The observed GR – A<sub>sat</sub> relationships were seemingly underpinned more by variation in g<sub>s</sub> than by CO<sub>2</sub> drawdown, g<sub>s</sub> being more consistently related to GR than was  $\delta^{13}$ C (Table 2), and relationships between A<sub>sat</sub> and g<sub>s</sub> being tighter than those between A<sub>sat</sub> and  $\delta^{13}$ C ( $r^2 = 0.63 - 0.70 vs. 0.25 - 0.35$ , respectively).

260

### 261 Growth Rate – stem trait relationships

Growth rate was unrelated to either sapwood or whole-stem tissue density (Fig. 2a; Table 2).
Fractional bark allocation varied from *ca*. 20 % to 60 % by mass and was unrelated to GR in

saplings, and negatively correlated with GR among adults ( $r^2 = 0.30$ , P = 0.024; Fig 2b). That

said, the sapling and adult slopes were deemed not significantly different from one another (slope test, P = 0.223), with the common fitted slope significantly negative (P = 0.016; model  $r^2 = 0.21$ ). This suggests some tendency for species with higher relative investment in bark to have slower stem diameter growth rates.

269

# 270 Relationships between GR and branch-scale biomass allocation

Total leaf mass was unrelated to GR in each of the four comparisons (Fig 3a; all P > 0.1). For adult plants, stem mass was negatively related to growth rate ( $r^2 = 0.27 - 0.32$ ; Fig. 3b); for saplings, no relationship was observed (Table 2). Nonetheless, the four stem mass – GR slopes did not differ significantly from one another (P = 0.731), with the common slope

explaining 25% of GR variation (significantly negative, P < 0.001).

276 Leaf and stem mass tended to explain more variation in growth rate when considered 277 together, rather than one at a time. LM:SM explained between 16 % and 65 % variation in 278 GR, depending on plant age and season (Fig. 3c). The four GR – LM:SM regression slopes 279 were deemed non-heterogeneous (P = 0.414), with the common fitted slope explaining 33% 280 variation in GR (significantly positive slope, P < 0.001). The allometric descriptor a LMSM 281 explained between 32 % and 51% variation in GR (Fig. 3d; Table 2). The four GR a LMSM regressions were deemed non-heterogeneous (P = 0.300), with a common fitted 282 283 slope explaining 36% variation (significantly positive slope, P < 0.001).

284

## 285 Combined explanatory power of leaf:stem allocation and photosynthesis

286 To be consistent with treating photosynthetic rate (Asat) on a per area basis, for this analysis

287 we used the ratio of leaf *area* to stem mass (LA:SM), rather than LM:SM. Regression models

288	including an interaction term between Asat and LA:SM were run first. With these interactions
289	deemed non-significant for both saplings and adults (both $P > 0.2$ ), we next ran models
290	including main effects only. For adults, Asat and LA:SM together explained 64 % of GR
291	variation, with each variable contributing similar explanatory power (as judged by their
292	respective F-statistics; Table 3). For saplings, the two traits explained 28 % of GR variation,
293	just a modest improvement over A <sub>sat</sub> on its own ( $r^2 = 0.23$ ; Table 2), and in this case the
294	coefficient for LA:SM was non-significant and that for Asat just marginally so (Table 3).
295	For the allometric descriptor we continued to use a_LMSM (the mass-basis
296	allometry). For adults, Asat and a_LMSM explained 80% of GR variation, in a regression
297	model that included a highly significant, positive interaction between the traits (Table 3).
298	This interaction term indicated that the effect of higher Asat on GR was stronger in species
299	with higher a_LMSM, and vice versa. For saplings, the two traits explained 41% of GR
300	variation (Table 3), just a modest improvement of a_LMSM on its own ( $r^2 = 0.32$ ; Table 2),
301	and in this case the coefficient for $A_{sat}$ was non-significant (Table 3).
302	For the adult dataset these relationships were strongly influenced by Acacia difficilis.
303	Re-running the analyses with this species removed, Asat and a_LMSM together still explained
304	57% of GR variation (multiple regression with interaction term; not shown), and $A_{sat}$ and
305	LA:SM explained 34 % (main effects only; not shown).
306	A supplementary analysis, incorporating phylogenetic information as
307	Phylogenetically Independent Contrasts (Supplementary Appendix 1), gave additional
308	strength to our results. In those analyses also, variation in growth rate was clearly
309	(significantly) connected to variation in leaf:stem allometry, Asat and fractional bark
310	allocation, but not to variation in either SLA or wood density.
311	

#### 312 **DISCUSSION**

## 313 Photosynthetic rate as a driver of GR variation

314 Light-saturated photosynthetic rate, Asat, explained roughly 30% of interspecific variation in 315 stem diameter growth rates. This mirrors the generally positive A<sub>sat</sub> – growth rate relationship 316 known from controlled-environment seedling studies (Kruger and Volin 2006) and older 317 seedlings grown for two years in a shade house (Li et al. 2016). By contrast, there have been 318 rather few field-based tests of GR – Asat relationships, especially for sapling or adult plants. A 319 positive relationship has been reported across several vegetation types in northern Australia 320 (Prior et al. 2004), from open forest in Hawaii (Stratton and Goldstein 2001), and from 321 saplings of 53 species growing in high-light gaps within a Bolivian rainforest (Poorter and 322 Bongers 2006). No relationship was observed among 24 savanna and forest species in Brazil 323 (Rossatto et al. 2009).

A generally positive  $GR - A_{sat}$  relationship makes intuitive sense, especially in high light situations. That said, species with higher  $A_{sat}$  could in principle deploy less total leaf area, or have higher leaf replacement costs (shorter leaf lifespan), or have higher belowground or sapwood respiration costs, and these costs could potentially cancel out the growth benefit of higher  $A_{sat}$ . However, for both adults and saplings of the 17 species examined here this was seemingly not the case, in support of the prediction that the  $GR - A_{sat}$ relationship does not vary with plant stature (Gibert *et al.* 2016).

331

#### 332 *Leaf:stem allocation as a driver of GR variation*

This study represents the first test of the proposition that higher relative allocation to leafversus stems, considered at branch level, should drive faster growth at the whole-plant level

335 (Pickup et al. 2005). Those authors argued that higher allocation to leaf represented greater 336 potential for photosynthetic benefits, while higher allocation to stem would incur higher costs 337 for maintenance respiration. All else equal, a more positive carbon balance at branch-level 338 should lead to more carbon exported to other parts of the plant, and thus faster growth 339 (Pickup et al. 2005). Our results are consistent with this interpretation. However, as it turned 340 out, the positive influence of leaf:stem allocation on growth rate was more strongly driven by 341 stem allocation (which was negatively correlated with GR) than by leaf mass (which was 342 unrelated to GR).

Higher investment in stem tissues must also have benefits, for example it may
represent more biomechanical support and hydraulic supply to leaves, greater potential for
water storage in sapwood and the living inner bark layers (Rosell 2016), and greater
protection against fire, especially from the outer bark layers (Pausas 2014). Presumably, all of
these features can be important in seasonally-dry, frequently-burnt savannas, such as that
studied here.

349

# 350 No general relationship between GR and SLA for field-grown plants

351 The other key finding from this study is the null result, that SLA failed to explain variation in 352 GR for either saplings or adult plants. The size-dependence of GR – SLA relationships has 353 been investigated in several recent studies, but conclusions have varied. SLA and stem 354 diameter growth rate were: positively correlated among Puerto Rico rainforest species, with 355 no size-related trend in relationship strength (Lasky et al. 2015); unrelated across all size 356 classes in a very detailed study of Panamanian rainforest species (Visser et al. 2016); 357 unrelated across all size classes except mid-size trees (DBH = 16-18 cm) in a seasonally dry 358 forest, Brazil (Prado-Junior et al. 2016); and positively correlated among adult rainforest

trees in Taiwan, but *negatively* correlated among juveniles (Iida *et al.* 2014). Mostly these
studies have focused on forests with a marked vertical light gradient and (at least partially as
a result) marked shifts in leaf traits between young and old plants. Our study represents an
important contrast, coming from open vegetation with little vertical light gradient, and where
leaf and wood traits of saplings were indistinguishable from adults (Fig. S1).

364

# 365 *GR* unrelated to stem tissue density and leaf nutrient concentrations

366 Other hypotheses detailed in the Introduction receiving no support were the predictions that 367 GR would be negatively related to WD, and positively related to leaf N and P concentrations. 368 A negative relationship between stem diameter growth rates and trunk wood density has been 369 reported many times (Iida et al. 2014; King et al. 2006; Lasky et al. 2015; Martínez-Vilalta et 370 al. 2010; Poorter et al. 2008; Wright et al. 2010), although null relationships have also been 371 reported (Russo et al. 2010). Even when statistically significant the explanatory power is 372 generally rather low in these studies, typically < 10% (but see Martínez-Vilalta et al. 2010), 373 and sample size (number of species) is usually far higher than used here. Perhaps we lacked 374 sufficient power to detect a relationship between growth rate and wood density, if indeed 375 there is a general tendency for this to be true among Australian savanna species. Here we 376 measured tissue density on terminal branches rather than main trunks; however, these 377 properties would likely be related (Swenson and Enquist 2008), and their relationships to GR therefore similar. 378

Growth rates were also unrelated to leaf N and P concentrations (considered per unit leaf area or mass). Perhaps this should not be surprising since  $A_{sat}$  was at best marginally correlated with  $N_{area}$  (in saplings, r = 0.46, P = 0.062) and unrelated to  $P_{area}$  in both age groups (P > 0.140; not shown), and the causal pathway between leaf N (and P) and GR

arguably flows via their respective links to photosynthesis. Considered on a leaf mass basis,
leaf N, leaf P and photosynthetic rate were all correlated with SLA (not shown). Covariation
in this suite of traits can be thought of representing variation along a "leaf economic
spectrum" (Wright *et al.* 2004). Hence, the finding that all of SLA, N<sub>mass</sub> and P<sub>mass</sub> were
unrelated to GR (Table 2) can also be taken to illustrate how strategy variation along the leaf
economic spectrum does not necessarily map on to variation in growth rate of field-grown
plants.

390

# 391 Successful growth – trait strategies

392 Our results help us understand the biology underpinning differences among species in their 393 growth rates. For example, the short-lived, fire-sensitive species A. difficilis was the fastest 394 growing species, with this high GR seemingly driven by the combination of high leaf:stem mass ratio (1.6 - 2.9 g.g<sup>-1</sup> depending on season; Table S1), low fractional allocation to bark 395 (23 %) and fast Asat (26 µmol m<sup>-2</sup> s<sup>-1</sup>), itself associated with profligate photosynthetic water 396 use (gs; 1087 mmol m<sup>-2</sup> s<sup>-1</sup>). Grevillea pteridifolia seemingly achieves its fast GR by teaming 397 high LM:SM (ca. 3.6 g g<sup>-1</sup> in saplings, 2.4 g g<sup>-1</sup> in adults) with medium-high A<sub>sat</sub> (18.2 – 19.7 398  $\mu$ mol m<sup>-2</sup> s<sup>-1</sup>), while *Eucalyptus miniata* does so by teaming very rapid A<sub>sat</sub> (*ca.* 25  $\mu$ mol m<sup>-2</sup> 399  $s^{-1}$ ) with mid-range LM:SM (1.2 – 1.7 g.g<sup>-1</sup>). Conversely, the slow GR of adult *Petalostigma* 400 pubescens was associated with the lowest mean  $A_{sat}$  value (12.4 µmol m<sup>-2</sup> s<sup>-1</sup>), slow  $g_s$  (392) 401 mmol m<sup>-2</sup> s<sup>-1</sup>) and low LM:SM (ca. 1.0 g g<sup>-1</sup>). The slow GR of sapling Buchanania obovata 402 was associated with the lowest  $A_{sat}$  values (11.6  $\mu$ mol m<sup>-2</sup> s<sup>-1</sup>) seen for saplings, the lowest  $g_s$ 403 (301 mmol m<sup>-2</sup> s<sup>-1</sup>), low LM:SM (0.6 - 0.9 g g<sup>-1</sup>, depending on season), and the highest 404

405 fractional allocation to bark (60%).

406 That these species vary widely in stem diameter growth rate yet still co-occur 407 indicates that there are many equally successful growth and survival strategies in this system. 408 Of particular importance, not addressed in this study, is the need for sufficient individuals of 409 each species to overcome the demographic bottleneck that frequent, intense fires create – the 410 so-called sapling "fire trap" (Bond 2008; Prior et al. 2010). For investigating that aspect of 411 the growth strategy, additional data for height growth rate would be especially valuable. 412 413 **Svnthesis** 414 Our results suggest that branch-level biomass allocation is a property deserving serious 415 attention in future studies of plant growth. Here we presented two types of indices: static 416 ratios (LM:SM, LA:SM), and an allometric descriptor (a LMSM). Not surprisingly a LMSM 417 and LM:SM were correlated, but weakly enough that each variable contains considerable 418 independent information. Besides explaining considerable variation in GR, branch-level 419 biomass allocation also has the virtue of being easy to measure. 420 Together, Asat and leaf:stem deployment explained up to 41% GR variation in 421 saplings and up to 80% in adults (57% with Acacia difficilis excluded). These results are 422 remarkable given that we did not consider interspecific variation in, for example, canopy

423 architecture, below-ground allocation, or tissue turnover rates. Of course, the extent to which

424 these results generalise to other savannas or other vegetation types will require further study.

425 On the face of it, one might predict lower explanatory power for these traits in vegetation

426 types where many individuals are shaded, or where many individuals are very large (i.e.,

427 situations where the benefits of higher  $A_{sat}$  are likely muted, and where sapwood respiration

428 costs begin to dominate GR variation). Testing this hypothesis would be a useful next step.

429 Currently we lack clear demonstration (i.e., within a single empirical study) that SLA 430 - growth rate relationships flip as predicted, from positive in small plants to null in large 431 plants. Nonetheless we suggest that this is likely the case, given the wealth of seedling 432 studies showing strong positive relationships and the increasing number of sapling-adult 433 studies showing little if any relationship. Even so, it is as yet unclear at what plant size these 434 shifts occur and whether they occur systematically earlier or later in particular light 435 environments or vegetation types, or in plants with particular architectures or growth 436 strategies.

437 That SLA is often unrelated to growth rates of field-grown plants has been 438 disappointing for authors expecting plant functional traits to neatly explain demographic 439 variation (Paine et al. 2015; Wright et al. 2010). Nonetheless, SLA remains a key trait for understanding leaf economic variation (Westoby et al. 2002); a key descriptor of how canopy 440 441 mass interconverts with canopy light-capturing area (and thus an important property in plant 442 growth models); and it is related to important ecological variation at other scales - for 443 example, herbivory rates, flammability, and litter decomposition rate (Poorter et al. 2009). 444 But, it seems increasingly clear that SLA and plant growth rate are generally correlated only 445 in very small plants, and this fact should be more widely appreciated.

446

#### 447 Author contributions

IJW and CEL designed the study. All authors contributed to fieldwork and discussion of
results. IJW analysed the data and drafted the manuscript. All authors contributed to
subsequent versions.

#### 452 **REFERENCES**

- 453 Aiba M. & Nakashizuka T. (2009) Growth properties of 16 non-pioneer rain forest tree
- 454 species differing in sapling architecture. *J Ecol* **97**, 992-9.
- 455 Bond W. J. (2008) What Limits Trees in C-4 Grasslands and Savannas? Annu Rev Ecol Evol
- 456 *Syst* **39**, 641-59.
- 457 Cernusak L. A., Winter K., Aranda J. & Turner B. L. (2008) Conifers, angiosperm trees, and
- 458 lianas: Growth, whole-plant water and nitrogen use efficiency, and stable isotope
- 459 composition (delta C-13 and delta O-18) of seedlings grown in a tropical environment. *Plant*
- 460 *Physiology* **148**, 642-59.
- 461 Chabot B. F. & Hicks D. J. (1982) The ecology of leaf life spans. *Annu Rev Ecol Syst* **13**,
- 462 229-59.
- 463 Clark D. A. & Clark D. B. (1999) Assessing the growth of tropical rain forest trees: Issues for
  464 forest modeling and management. *Ecol Appl* 9, 981-97.
- 465 Cook G. D. & Corbett L. K. (2003) Kapalga and the Fire Experiment. In: Fire in Tropical
- 466 Savannas: The Kapalga Fire Experiment (eds A. N. Andersen, G. D. Cook and R. J.
- 467 Williams) pp. 15-32. Springer-Verlag, New York.
- 468 Coomes D. A. & Grubb P. J. (1998) A comparison of 12 tree species of Amazonian caatinga
- using growth rates in gaps and understorey, and allometric relationships. *Funct Ecol* 12, 42635.
- 471 Easdale T. A. & Healey J. R. (2009) Resource-use-related traits correlate with population
- 472 turnover rates, but not stem diameter growth rates, in 29 subtropical montane tree species.
- 473 *Perspect Plant Ecol Evol Syst* **11**, 203-18.
- 474 Enquist B. J., Kerkhoff A. J., Stark S. C., Swenson N. G., McCarthy M. C. & Price C. A.
- 475 (2007) A general integrative model for scaling plant growth, carbon flux, and functional trait
- 476 spectra. *Nature* **449**, 218-22.

- 477 Farquhar G. D., O'Leary M. H. & Berry J. A. (1982) On the Relationship between Carbon
- 478 Isotope Discrimination and the Inter Cellular Carbon Dioxide Concentration in Leaves. *Aust*479 *J Plant Physiol* 9, 121-38.
- 480 Gibert A., Gray E. F., Westoby M., Wright I. J. & Falster D. S. (2016) On the link between
- 481 functional traits and growth rate: meta-analysis shows effects change with plant size, as
- 482 predicted. *J Ecol* **104**, 1488-503.
- 483 Gower S. T., Reich P. B. & Son Y. (1993) Canopy dynamics and aboveground production of
- 484 five tree species with different leaf longevities. *Tree Physiol* **12**, 327-45.
- 485 Grime J. P. & Hunt R. (1975) Relative growth-rate: its range and adaptive significance in a
- 486 local flora. *J Ecol* **63**, 393-422.
- 487 Grime J. P., Thompson K., Hunt R., Hodgson J. G., Cornelissen J. H. C., Rorison I. H.,
- 488 Hendry G. A. F., Ashenden T. W., Askew A. P., Band S. R., Booth R. E., Bossard C. C.,
- 489 Campbell B. D., Cooper J. E. L., Davison A. W., Gupta P. L., Hall W., Hand D. W., Hannah
- 490 M. A., Hillier S. H., Hodkinson D. J., Jalili A., Liu Z., Mackey J. M. L., Matthews N.,
- 491 Mowforth M. A., Meal A. M., Reader R. J., Reiling K., Ross-Fraser W., Sutton F., Tasker D.
- 492 E., Thorpe P. C. & Whitehouse J. (1997) Integrated screening validates primary axes of
- 493 specialisation in plants. *Oikos* **79**, 259-81.
- 494 Hérault B., Bachelot B., Poorter L., Rossi V., Bongers F., Chave J., Paine C. E. T., Wagner F.
- 495 & Baraloto C. (2011) Functional traits shape ontogenetic growth trajectories of rain forest
- 496 tree species. *J Ecol* **99**, 1431-40.
- 497 Hutley L. B., O'Grady A. P. & Eamus D. (2001) Monsoonal influences on evapotranspiration
- 498 of savanna vegetation of northern Australia. *Oecologia* **126**, 434-43.
- 499 Iida Y., Kohyama T. S., Swenson N. G., Su S.-H., Chen C.-T., Chiang J.-M. & Sun I. F.
- 500 (2014) Linking functional traits and demographic rates in a subtropical tree community: the
- 501 importance of size dependency. *J Ecol* **102**, 641-50.

- 502 King D. A. (1999) Juvenile foliage and the scaling of tree proportions, with emphasis on
- 503 Eucalyptus. *Ecology* **80**, 1944-54.
- 504 King D. A., Davies S. J., Tan S. & Noor N. S. M. (2006) The role of wood density and stem
- support costs in the growth and mortality of tropical trees. *J Ecol* **94**, 670-80.
- 506 Konings H. (1989) Physiological and morphological differences between plants with a high
- 507 NAR or a high LAR as related to environmental conditions. In: *Causes and consequences of*
- 508 variation in growth rate and productivity of higher plants (eds H. Lambers, M. L.
- 509 Cambridge, H. Konings and T. L. Pons) pp. 101-23. SPB Academic Publishing, The Hague,
- 510 The Netherlands.
- 511 Kruger E. L. & Volin J. C. (2006) Reexamining the empirical relation between plant growth
- and leaf photosynthesis. *Funct Plant Biol* **33**, 421-9.
- 513 Lambers H. & Poorter H. (1992) Inherent variation in growth rate between higher plants: a
- search for ecological causes and consequences. *Adv Ecol Res* 23, 187-261.
- 515 Lambers H., Poorter H. & Van Vuren M. M. I. (1998) Inherent variation in plant growth :
- 516 physiological mechanisms and ecological consequences. Backhuys, Leiden.
- 517 Lasky J. R., Bachelot B., Muscarella R., Schwartz N., Forero-Montana J., Nytch C. J.,
- 518 Swenson N. G., Thompson J., Zimmerman J. K. & Uriarte M. (2015) Ontogenetic shifts in
- trait-mediated mechanisms of plant community assembly. *Ecology* **96**, 2157-69.
- 520 Li X., Schmid B., Wang F. & Paine C. E. T. (2016) Net Assimilation Rate Determines the
- 521 Growth Rates of 14 Species of Subtropical Forest Trees. *PLOS ONE* **11**, e0150644.
- 522 Martínez-Vilalta J., Mencuccini M., Vayreda J. & Retana J. (2010) Interspecific variation in
- 523 functional traits, not climatic differences among species ranges, determines demographic
- 524 rates across 44 temperate and Mediterranean tree species. *J Ecol* **98**, 1462-75.
- 525 Matyssek R. (1986) Carbon, water and nitrogen relations in evergreen and deciduous
- 526 conifers. *Tree Physiol* **2**, 177-87.

- 527 Murphy B. P., Russell-Smith J. & Prior L. D. (2010) Frequent fires reduce tree growth in
- 528 northern Australian savannas: implications for tree demography and carbon sequestration.
- 529 *Glob Change Biol* **16**, 331-43.
- 530 Paine C. E. T., Amissah L., Auge H., Baraloto C., Baruffol M., Bourland N., Bruelheide H.,
- 531 Daïnou K., de Gouvenain R. C., Doucet J.-L., Doust S., Fine P. V. A., Fortunel C., Haase J.,
- 532 Holl K. D., Jactel H., Li X., Kitajima K., Koricheva J., Martínez-Garza C., Messier C.,
- 533 Paquette A., Philipson C., Piotto D., Poorter L., Posada J. M., Potvin C., Rainio K., Russo S.
- 534 E., Ruiz-Jaen M., Scherer-Lorenzen M., Webb C. O., Wright S. J., Zahawi R. A. & Hector A.
- 535 (2015) Globally, functional traits are weak predictors of juvenile tree growth, and we do not
- 536 know why. *J Ecol* **103**, 978-89.
- 537 Pausas J. G. (2014) Bark thickness and fire regime. *Funct Ecol*, n/a-n/a.
- 538 Pickup M., Westoby M. & Basden A. (2005) Dry mass costs of deploying leaf area in relation
  539 to leaf size. *Funct Ecol* 19, 88-97.
- 540 Poorter H. & Garnier E. (2007) Ecological significance of inherent variation in relative
- 541 growth rate and its components. In: *Functional Plant Ecology* (eds F. I. Pugnaire and F.
- 542 Valladares) pp. 67-100. CRC Press, Boca Raton, FL USA.
- 543 Poorter H., Niinemets U., Poorter L., Wright I. J. & Villar R. (2009) Causes and
- 544 consequences of variation in leaf mass per area (LMA): a meta-analysis. *New Phytol* **182**,
- 545 565-88.
- 546 Poorter H., Niklas K. J., Reich P. B., Oleksyn J., Poot P. & Mommer L. (2012) Biomass
- 547 allocation to leaves, stems and roots: meta-analyses of interspecific variation and
- 548 environmental control. *New Phytol* **193**, 30-50.
- 549 Poorter H. & van der Werf A. (1998) Is inherent variation in RGR determined by LAR at low
- 550 irradiance and by NAR at high irradiance? A review of herbaceous species. In: Inherent

- 551 variation in plant growth. Physiological mechanisms and ecological consequences (eds H.
- Lambers, H. Poorter and M. M. I. Van Vuuren) pp. 309-36. Backhuys Publishers, Leiden.
- 553 Poorter L. & Bongers F. (2006) Leaf traits are good predictors of plant performance across 53
- rain forest species. *Ecology* **87**, 1733-43.
- 555 Poorter L., Paz H., Wright S. J., Ackerly D. D., Condit R., Ibarra-Manriquez G., Harms K. E.,
- 556 Licona J. C., Martinez-Ramos M., Mazer S. J., Muller-Landau H. C., Peña-Claros M., Webb
- 557 C. O. & Wright I. J. (2008) Are functional traits good predictors of demographic rates?
- 558 Evidence from five Neotropical forests. *Ecology* **89**, 1908-20.
- 559 Prado-Junior J. A., Schiavini I., Vale V. S., Raymundo D., Lopes S. F. & Poorter L. (2016)
- 560 Functional traits shape size-dependent growth and mortality rates of dry forest tree species.
- 561 Journal of Plant Ecology.
- 562 Prior L. D., Brook B. W., Williams R. J., Werner P. A., Bradshaw C. J. A. & Bowman D. M.
- 563 J. S. (2006) Environmental and allometric drivers of tree growth rates in a north Australian
- 564 savanna. *For Ecol Manag* **234**, 164-80.
- 565 Prior L. D., Eamus D. & Bowman D. M. J. S. (2004) Tree growth rates in north Australian
- savanna habitats: seasonal patterns and correlations with leaf attributes. Aust J Bot 52, 303-
- 567 14.
- 568 Prior L. D., Williams R. J. & Bowman D. M. J. S. (2010) Experimental evidence that fire
- 569 causes a tree recruitment bottleneck in an Australian tropical savanna. *J Trop Ecol* 26, 595570 603.
- 571 Rees M., Osborne Colin P., Woodward F. I., Hulme Stephen P., Turnbull Lindsay A. &
- 572 Taylor Samuel H. (2010) Partitioning the Components of Relative Growth Rate: How
- 573 Important Is Plant Size Variation? *The American Naturalist* **176**, E152-E61.
- 574 Reich P. B. (2014) The world-wide 'fast–slow' plant economics spectrum: a traits manifesto.
- 575 *J Ecol* **102**, 275-301.

- 576 Reich P. B., Walters M. B. & Ellsworth D. S. (1992) Leaf life-span in relation to leaf, plant,
- and stand characteristics among diverse ecosystems. *Ecol Monogr* **62**, 365-92.
- 578 Rosell J. A. (2016) Bark thickness across the angiosperms: more than just fire. *New Phytol*579 **211**, 90-102.
- 580 Rossatto D. R., Hoffmann W. A. & Franco A. C. (2009) Differences in growth patterns
- between co-occurring forest and savanna trees affect the forest-savanna boundary. *Funct Ecol*23, 689-98.
- 583 Rozendaal D. M. A., Soliz-Gamboa C. C. & Zuidema P. A. (2010) Timber yield projections
- for tropical tree species: The influence of fast juvenile growth on timber volume recovery.
- 585 *For Ecol Manag* **259**, 2292-300.
- Rüger N., Wirth C., Wright S. J. & Condit R. (2012) Functional traits explain light and size
  response of growth rates in tropical tree species. *Ecology* 93, 2626-36.
- 588 Russell-Smith J., Yates C., Edwards A., Allan G. E., Cook G. D., Cooke P., Craig R., Heath
- 589 B. & Smith R. (2003) Contemporary fire regimes of northern Australia, 1997–2001:
- 590 change since Aboriginal occupancy, challenges for sustainable management. Int J Wildland
- 591 *Fire* **12**, 283-97.
- 592 Russo S. E., Jenkins K. L., Wiser S. K., Uriarte M., Duncan R. P. & Coomes D. A. (2010)
- 593 Interspecific relationships among growth, mortality and xylem traits of woody species from
- 594 New Zealand. *Funct Ecol* **24**, 253-62.
- 595 Ryan M. G. (1989) Sapwood Volume for Three Subalpine Conifers Predictive Equations and
- 596 Ecological Implications. *Can J For Res* **19**, 1397-401.
- 597 Shipley B. (2006) Net assimilation rate, specific leaf area and leaf mass ratio: which is most
- 598 closely correlated with relative growth rate? A meta-analysis. *Funct Ecol* **20**, 565-74.
- 599 Stratton L. C. & Goldstein G. (2001) Carbon uptake, growth and resource-use efficiency in
- one invasive and six native Hawaiian dry forest tree species. *Tree Physiol* **21**, 1327-34.

- 601 Swenson N. G. & Enquist B. J. (2008) The relationship between stem and branch wood
- 602 specific gravity and the ability of each measure to predict leaf area. *Am J Bot* **95**, 516-9.
- 603 Veneklaas E. & Poorter L. (1998) Growth and carbon partitioning of tropical tree seedlings in
- 604 contrasting light environments. In: Inherent variation in plant growth. Physiological
- 605 mechanisms and ecological consequences (eds H. Lambers, H. Poorter and M. M. I. Van
- 606 Vuuren) pp. 337-61. Backhuys Publishers, Leiden.
- 607 Visser M. D., Bruijning M., Wright S. J., Muller-Landau H. C., Jongejans E., Comita L. S. &
- de Kroon H. (2016) Functional traits as predictors of vital rates across the life cycle of
- tropical trees. *Funct Ecol* **30**, 168-80.
- 610 Warton D. I., Wright I. J., Falster D. S. & Westoby M. (2006) Bivariate line-fitting methods
- 611 for allometry. *Biological Reviews* **81**, 259-91.
- 612 Westoby M., Falster D. S., Moles A. T., Vesk P. A. & Wright I. J. (2002) Plant ecological
- 613 strategies: some leading dimensions of variation between species. Annu Rev Ecol Syst 33,
- 614 125-59.
- 615 Williams R. J., Muller W. J., Wahren C. H., Setterfield S. A. & Cusack J. (2003) Fire impacts
- 616 on savanna vegetation. In: Fire in Tropical Savannas: The Kapalga Fire Experiment (eds A.
- 617 N. Andersen, G. D. Cook and R. J. Williams) pp. 79-106. Springer-Verlag, New York.
- 618 Williams R. J., Myers B. A., Muller W. J., Duff G. A. & Eamus D. (1997) Leaf phenology of
- 619 woody species in a north Australian tropical savanna. *Ecology* **78**, 2542-58.
- 620 Wright I. J., Reich P. B., Westoby M., Ackerly D. D., Baruch Z., Bongers F., Cavender-Bares
- 621 J., Chapin F. S., Cornelissen J. H. C., Diemer M., Flexas J., Garnier E., Groom P. K., Gulias
- J., Hikosaka K., Lamont B. B., Lee T., Lee W., Lusk C., Midgley J. J., Navas M.-L.,
- 623 Niinemets Ü., Oleksyn J., Osada N., Poorter H., Poot P., Prior L., P'yankov V. I., Roumet C.,
- 624 Thomas S. C., Tjoelker M. G., Veneklaas E. J. & Villar R. (2004) The world-wide leaf
- 625 economics spectrum. *Nature* **428**, 821-7.

- 626 Wright S. J., Kitajima K., Kraft N. J. B., Reich P. B., Wright I. J., Bunker D. E., Condit R.,
- 627 Dalling J. W., Davies S. J., Díaz S., Engelbrecht B. M. J., Harms K. E., Hubbell S. P., Marks
- 628 C. O., Ruiz-Jaen M. C., Salvador C. M. & Zanne A. E. (2010) Functional traits and the
- 629 growth–mortality trade-off in tropical trees. *Ecology* **91**, 3664-74.

**Table 1.** List of study species, some key life history characteristics, the stem diameter cut-offs (at 1.3 m height) used to distinguish small plants ("saplings") from large plants ("adults"), and the sample size (number of data points) used for estimating stem-diameter growth rates. Semideciduous species are evergreen, but exhibit a noticeable thinning of the canopy during the dry season.

Species	Family	Leaf phenology	Habit	N <sub>2</sub>	DBH cut-off	GR sample size
				fixer	(cm)	(saplings, adults)
Acacia difficilis	Fabaceae	Evergreen	Shrub	Yes	1.5	-,19
Acacia latescens	Fabaceae	Evergreen	Shrub	Yes	3	77, 67
Acacia mimula	Fabaceae	Evergreen	Shrub	Yes	3	15, 31
Buchanania obovata	Anacardiaceae	Semi-deciduous	Tree	No	6	31, 95
Cochlospermum fraseri	Bixaceae	Deciduous	Tree	No	3	8, 21
Corymbia bleeseri	Myrtaceae	Semi-deciduous	Tree	No	6	44, 220
Corymbia porrecta	Myrtaceae	Semi-deciduous	Tree	No	6	207, 752
Erythrophleum chlorostachys	Fabaceae	Semi-deciduous	Tree	Yes	6	273, 741
Eucalyptus miniata	Myrtaceae	Evergreen	Tree	No	6	635, 1559
Eucalyptus tetrodonta	Myrtaceae	Evergreen	Tree	No	6	346, 1810
Grevillea decurrens	Proteaceae	Semi-deciduous	Shrub	No	3	-,9
Grevillea pteridifolia	Proteaceae	Semi-deciduous	Shrub	No	3	12, 94
Lophostemon lactifluus	Myrtaceae	Semi-deciduous	Tree	No	6	34, 159
Persoonia falcata	Proteaceae	Semi-deciduous	Shrub	No	3	4, 36
Petalostigma pubescens	Picrodendraceae	Evergreen	Tree	No	6	24, 48
Planchonia careya	Lecythidaceae	Deciduous	Tree	No	3	4, 147
Xanthostemon paradoxus	Myrtaceae	Semi-deciduous	Tree	No	6	207, 168

**Table 2.** Tests for correlation between stem-diameter growth rates and various plant functional traits for the four age-season datasets.Correlations at least marginally significant (P < 0.1) are shown in bold, and the sign of these relationship is also indicated.Trait abbreviations. SLA: specific leaf area;  $N_{mass}$ ,  $N_{area}$ : leaf N per mass and area;  $P_{mass}$ ,  $P_{area}$ : leaf P per mass and area;  $A_{sat}$ : light-saturatedphotosynthetic rate;  $g_s$ : stomatal diffusional conductance to water; LM: leaf mass; SM: stem mass; LA: leaf area; a\_LM/SM: allometric slopedescribing leaf mass allocation relative to stem mass allocation. Note, "wood density" refers to whole-stem tissue density for dry seasonsampling, and density of sapwood only for the wet season.

Trait	Saplings (dry season)	Saplings (wet season)	Adults (dry season)	Adults (wet season)
	$r^2$ , P, n, sign	$r^2$ , P, n, sign	$r^2$ , P, n, sign	$r^2$ , P, n, sign
$\log SLA (cm^2 g^{-1})$	0.02, 0.664, 13	0.002, 0.870, 15	0.02, 0.633, 16	0.03, 0.492, 17
log N <sub>mass</sub> (%)	0.06, 0.425, 13	0.03, 0.536, 15	0.04, 0.499, 15	0.13, 0.149, 17
log N <sub>area</sub> (g m <sup>-2</sup> )	0.05, 0.468, 13	0.05, 0.409, 15	0.08, 0.315, 15	0.07, 0.302, 17
log P <sub>mass</sub> (%)	0.15, 0.193, 13	0.11, 0.226, 15	0.02, 0.606, 15	0.05, 0.397, 17
log P <sub>area</sub> (g m <sup>-2</sup> )	0.04, 0.526, 13	0.11, 0.223, 15	0.04, 0.487, 15	<0.001, 0.997, 17
log A <sub>sat</sub> (µmol m <sup>-2</sup> s <sup>-1</sup> )		0.23, 0.071, 15, +		0.32, 0.017, 17, +
$\log g_{\rm s} (\rm mmol \ m^{-2} \ s^{-1})$		0.20, 0.095, 15, +		0.17, 0.095, 17, +
Leaf d <sup>13</sup> C (‰)	0.03, 0.59, 12	0.25, 0.056, 15, -	0.01, 0.736, 16	0.13, 0.157, 17
log LM (g)	0.22, 0.101, 13	0.03, 0.520, 15	0.02, 0.613, 16	0.09, 0.252, 17
log SM (g)	0.13, 0.225, 13	0.09, 0.290, 15	0.27, 0.039, 16, -	0.32, 0.019, 17, -
log LM:SM (g g <sup>-1</sup> )	0.65, 0.001, 13, +	0.16, 0.134, 15	0.18, 0.097, 16, +	0.44, 0.004, 17, +
a_LM/SM	0.51, 0.006, 13, +	0.32, 0.027, 15, +	0.41, 0.008, 16, +	0.32, 0.017, 17, +
$\log LA:SM (cm^2 g^{-1})$	0.66, 0.001, 13, +	0.08, 0.313, 15	0.15, 0.132, 16, +	0.35, 0.013, 17, +
Bark % mass		0.05, 0.443, 15		0.30, 0.024, 17, -
Wood density (g cm <sup>-3</sup> )	0.002, 0.896, 13	<0.001, 0.947, 15	0.001, 0.923, 15	0.03, 0.473, 17

**Table 3.** Multiple regressions exploring the interactive effects of photosynthetic rate and leaf:stem allocation on plant growth rates

GR ~ Asat, LA:SM (interac	tion was never	significant)	
ADULTS ( $r^2 = 0.64, P < 0.0$	01, df = 16)		
Variable	F Coefficients		Р
Intercept	29.83	-2.53 (-3.52, -1.54)	< 0.001
log A <sub>sat</sub>	11.29	1.12 (0.41, 1.84)	0.005
log LA:SM	12.25	0.41 (0.16, 0.66)	0.004
SAPLINGS $(r^2 = 0.28, P = 0)$	0.022, df = 14)		
Variable	F	Coefficients	Р
Intercept	6.945	-1.43 (-2.61, -0.25)	0.022
log Asat	3.261	0.75 (-0.16, 1.66)	0.096
log LA:SM	0.766	0.10 (-0.15, 0.35)	0.399
<b>GR</b> ~ A <sub>sat</sub> , <b>a_LMSM</b> , A <sub>sat</sub> × ADULTS (model $r^2 = 0.80$ , ) Variable			Р
	<b>r</b> 13.36		_
Intercept	15.30	6.27 (2.56, 9.98)	0.003
log A <sub>sat</sub> a LMSM	20.22	-5.76 (-8.84, -2.69)	0.001
_	20.22	-9.64 (-14.27, -5.01)	<0.001
$\log A_{sat} \times a\_LMSM$	22.44	8.34 (4.54, 12.14)	<0.001
SAPLINGS (model $r^2 = 0.41$	P = 0.042, df =	= 14)	
Variable	F	Coefficients	Р
Intercept	5.86	-1.15 (-2.18, -0.12)	0.032
log A <sub>sat</sub>	1.78	0.53 (-0.34, 1.4)	0.207
a LM:SM	3.69	0.26 (-0.03, 0.55)	0.079

#### **FIGURE CAPTIONS**

**Figure 1.** Relationships between stem diameter growth rate and (a) specific leaf area, SLA; and (b) Light-saturated photosynthetic assimilation rate,  $A_{sat}$ , for each of the age-season datasets (note, photosynthesis measurements were made during the wet season only). Each data point represents a different species. Significant regression slopes (P < 0.05) shown as solid lines; marginally significant slopes (0.05 < P < 0.10) with dashed lines (full details given in Table 2). Symbols. Pink circles: adults, dry season; blue circles: adults, wet season; grey triangles: saplings, dry season; black triangles: saplings, wet season.

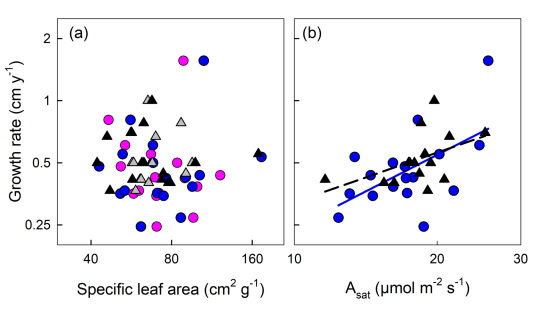
**Figure 2.** Relationship between stem diameter growth rate and properties of 1 cm diameter terminal stems. (a) Whole-stem (dry season) or sapwood (wet season) tissue density. (b) Percentage of stem mass allocated to bark (wet season only). Solid regression line indicates the relationship was significant (P < 0.05); see Table 2. Symbols. Pink circles: adults, dry season; blue circles: adults, wet season; grey triangles: saplings, dry season; black triangles: saplings, wet season.

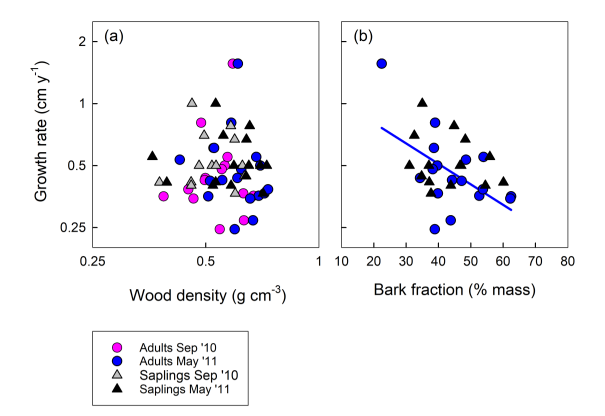
**Figure 3.** Relationship between stem diameter growth rate and leaf versus stem deployment on terminal branches. (a) Total leaf mass; (b) Total stem mass (including bark); (c) Ratio of leaf:stem mass. (d) Allometric coefficient describing the rate of leaf mass accumulation versus stem mass accumulation along the branch. Significant regression slopes (P < 0.05) shown as solid lines; marginally significant slopes (0.05 < P < 0.10) with dashed lines (full details given in Table 2). Symbols. Pink circles: adults, dry season; blue circles: adults, wet season; grey triangles: saplings, dry season; black triangles: saplings, wet season.

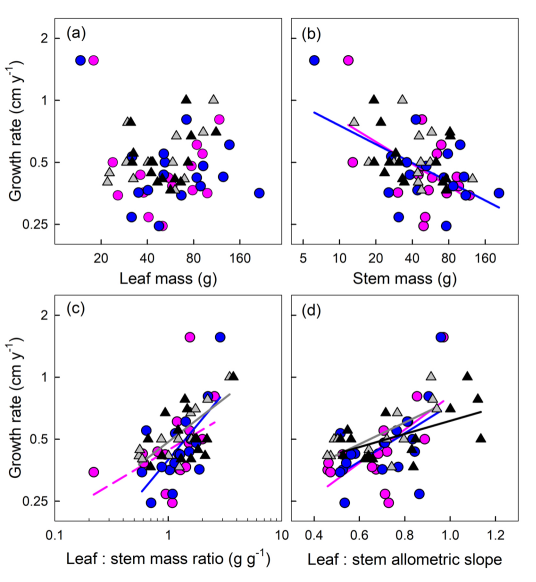
**Figure S1.** Comparison of growth rate and trait values of adults and saplings for each of the study species (Table 1). For trait data each data point represents the mean of *ca*. five replicates (see Methods). In each panel the 1:1 line is shown in black. There were insufficient

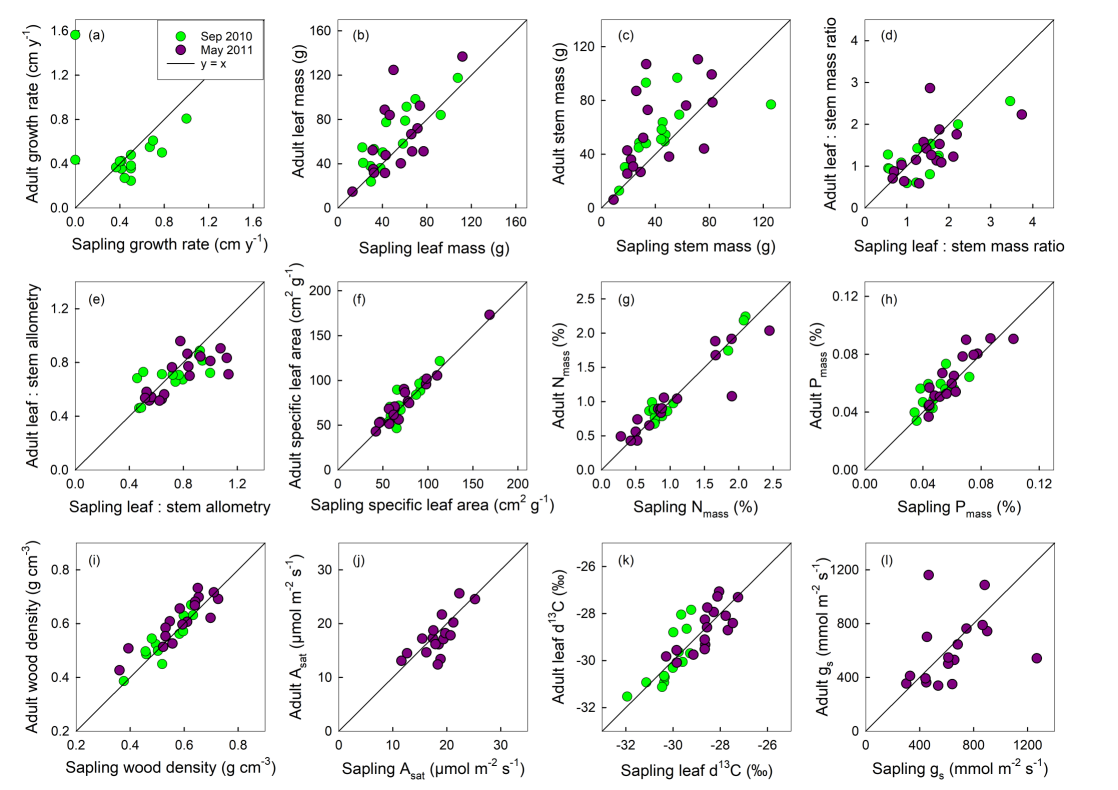
data to calculate growth rates for saplings of *Acacia difficilis* and *Grevillea decurrens*, so in panel (a) these are shown with GR = 0. September 2010 = dry season; May 2011 = wet season.

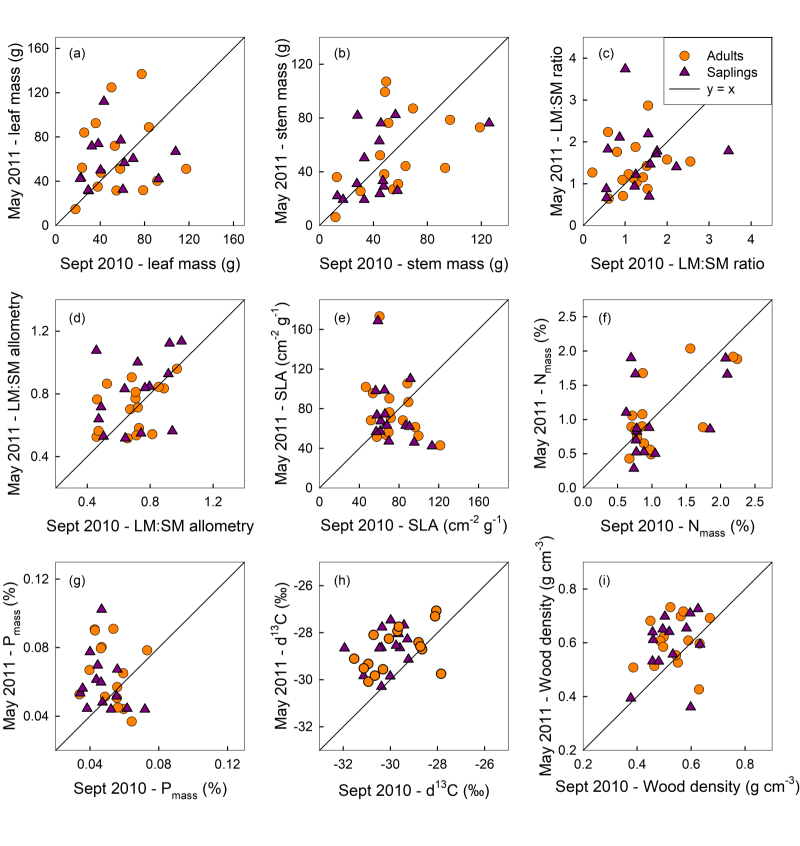
**Figure S2:** Comparison of trait values between the two seasons sampled (wet season, May 2011; dry season September 2010) for both adults (orange circles) and saplings (purple triangles) of 17 species. Points represent species-mean values calculated from *ca*. five individual plants (see Methods). In each panel the 1:1 line is shown in black. 'Wood' density measurements in 2010 (dry season) included bark and sapwood. Repeating this in 2011 (wet season) we removed the bark, giving the density of sapwood only (and in most cases higher values).



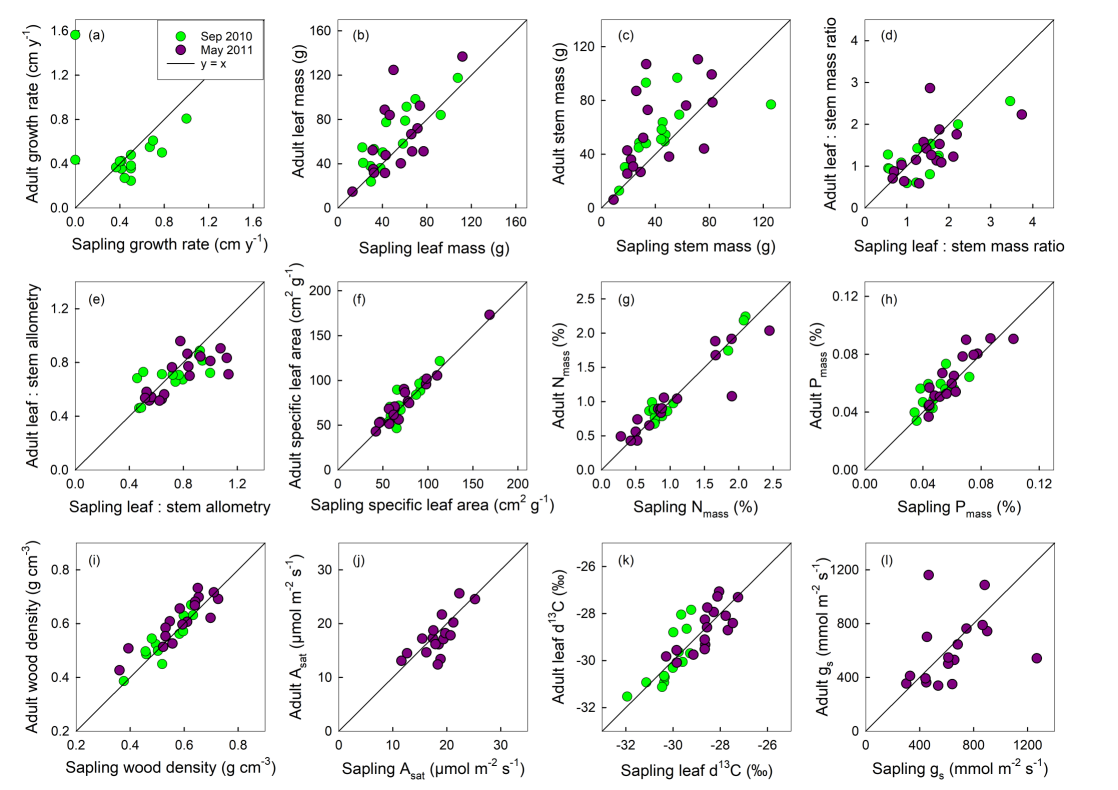




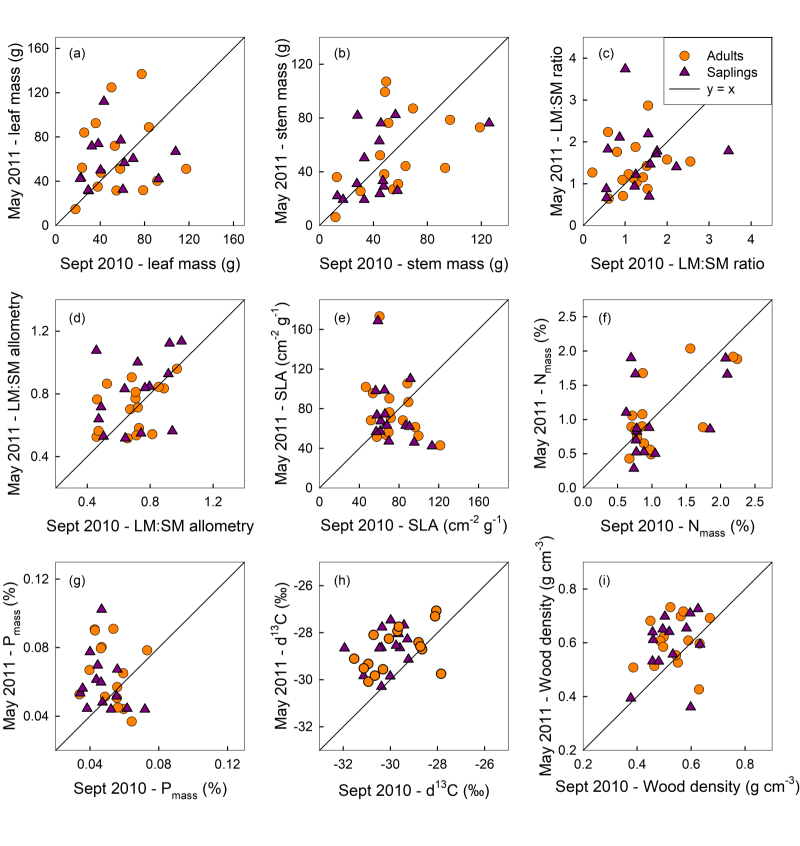




**Appendix S1.** Comparison of growth rate and trait values of adults and saplings for each of the study species (Table 1). For trait data each data point represents the mean of *ca*. five replicates (see Methods). In each panel the 1:1 line is shown in black. There were insufficient data to calculate growth rates for saplings of *Acacia difficilis* and *Grevillea decurrens*, so in panel (a) these are shown with GR = 0. September 2010 = dry season; May 2011 = wet season.



**Appendix S2.** Comparison of trait values between the two seasons sampled (wet season, May 2011; dry season September 2010) for both adults (orange circles) and saplings (purple triangles) of 17 species. Points represent species-mean values calculated from *ca*. five individual plants (see Methods). In each panel the 1:1 line is shown in black. 'Wood' density measurements in 2010 (dry season) included bark and sapwood. Repeating this in 2011 (wet season) we removed the bark, giving the density of sapwood only (and in most cases higher values).



Species	age	date	GR	SLA	leaf N	leaf N	leaf P	leaf P	A <sub>sat</sub>	gs	leaf d <sup>13</sup> C	Leaf mass	Stem mass (g)	LM:SM	LA:SM	a_LMSM	Wood density	Bark fraction
			(cm yr <sup>-1</sup> )	(cm <sup>2</sup> g <sup>-1</sup> )	(%)	(g m <sup>-2</sup> )	(%)	(g m <sup>-2</sup> )	(mmol m <sup>-2</sup> s <sup>-1</sup> )	(mmol m <sup>-2</sup> s <sup>-1</sup> )		(g)		(g g <sup>-1</sup> )	(cm <sup>2</sup> g <sup>-1</sup> )		(g cm <sup>-3</sup> )	(%)
Acacia difficilis	ad	May'11	1.56	105.5	2.04	1.93	0.091	0.086	25.61	1087	-29.56	14.7	6.2	2.87	302.2	0.959	0.608	22.5
Acacia difficilis	ad	Sept'10	1.56	88.5	1.56	1.76	0.054	0.061			-30.31	17.8	11.8	1.55	137.1	0.970	0.590	
Acacia difficilis	sap	May'11		110.2	2.45	2.22	0.086	0.078	22.34	882	-29.85	13.1	9.1	1.55	171.2	0.778	0.547	24.4
Acacia latescens	ad	May'11	0.50	68.2	1.88	2.76	0.057	0.083	16.13	530	-30.09	52.1	36.0	1.58	107.5	0.834	0.698	39.6
Acacia latescens	ad	Sept'10	0.50	83.9	2.24	2.67	0.056	0.067			-30.93	23.8	12.9	2.00	167.6	0.888	0.562	
Acacia latescens	sap	May'11	0.78	62.9	1.66	2.63	0.044	0.071	18.50	656	-29.84	31.2	22.0	1.40	88.1	1.122	0.654	44.8
Acacia latescens	sap	Sept'10	0.78	86.8	2.10	2.42	0.062	0.071			-31.13	29.7	13.2	2.22	192.7	0.923	0.583	
Acacia mimula	ad	May'11	0.36	70.7	1.92	2.71	0.067	0.095	17.29	645	-28.26	35.0	25.5	1.88	132.6	0.701	0.691	52.6
Acacia mimula	ad	Sept'10	0.36	71.7	2.19	3.05	0.040	0.055			-30.06	37.8	30.4	1.25	89.6	0.673	0.669	
Acacia mimula	sap	May'11	0.50	63.0	1.90	3.01	0.053	0.085	17.42	682	-28.65	31.8	19.3	1.78	112.0	0.847	0.726	47.0
Acacia mimula	sap	Sept'10	0.50	67.8	2.07	3.06	0.034	0.051			-29.59	29.3	17.5	1.76	119.3	0.796	0.626	
Buchanania obovata	ad	May'11	0.35	51.5	0.49	0.95	0.078	0.152	13.08	354	-27.07	214.1	207.8	1.03	53.0	0.526	0.508	62.5
Buchanania obovata	ad	Sept'10	0.35	57.7	0.99	1.71	0.073	0.127			-28.05	98.3	77.0	1.28	73.7	0.460	0.386	
Buchanania obovata	sap	May'11	0.41	57.1	0.28	0.50	0.067	0.118	11.61	301	-28.05	60.4	76.2	0.88	50.1	0.640	0.394	60.1
Buchanania obovata	sap	Sept'10	0.41	61.0	0.74	1.21	0.056	0.092			-29.65	69.7	125.7	0.55	33.6	0.473	0.377	
Cochlospermum fraseri	ad	May'11	0.53	173.1	1.08	0.62	0.091	0.052	13.39	350	-27.94	31.8	26.8	1.15	199.2	0.516	0.426	48.5
Cochlospermum fraseri	sap	May'11	0.55	168.6	1.90	1.13	0.102	0.061	18.85	642	-28.28	32.5	29.1	1.22	205.5	0.548	0.361	55.9

## APPENDIX S3. Growth rate and trait data used in this study.

Corymbia bleeseri	ad	May'11	0.37	53.5	0.89	1.67	0.051	0.096	21.67	1161	-28.71	40.3	44.1	0.87	46.8	0.771	0.716	39.8
Corymbia bleeseri	ad	Sept'10	0.37	60.4	0.86	1.43	0.043	0.071			-29.69	78.8	54.5	1.43	86.2	0.656	0.629	
Corymbia bleeseri	sap	May'11	0.37	47.1	0.83	1.76	0.048	0.102	19.10	467	-27.68	56.6	76.1	0.70	32.8	0.838	0.710	37.7
Corymbia bleeseri	sap	Sept'10	0.37	58.7	0.70	1.19	0.047	0.080			-29.28	60.5	47.5	1.25	73.5	0.742	0.598	
Corymbia porrecta	ad	May'11	0.55	52.5	0.89	1.69	0.044	0.084	20.21	501	-29.33	51.2	78.5	0.64	33.5	0.764	0.681	53.9
Corymbia porrecta	ad	Sept'10	0.55	67.0	0.70	1.05	0.049	0.073			-28.65	91.4	63.8	1.54	102.9	0.707	0.571	
Corymbia porrecta	sap	May'11	0.67	45.9	0.86	1.87	0.044	0.096	21.22	612	-28.64	77.1	82.4	0.94	43.1	0.716	0.641	48.3
Corymbia porrecta	sap	Sept'10	0.67	69.7	0.78	1.12	0.047	0.068			-29.42	61.7	45.4	1.57	109.8	0.767	0.596	
Erythrophleum chlorostachys	ad	May'11	0.38	95.6	1.68	1.75	0.080	0.084	16.16	362	-28.10	88.7	87.0	1.13	108.4	0.542	0.732	53.8
Erythrophleum chlorostachys	ad	Sept'10	0.38	99.7	1.74	1.75	0.060	0.060			-30.93	58.1	96.8	0.61	60.6	0.463	0.449	
Erythrophleum chlorostachys	sap	May'11	0.50	98.0	1.66	1.70	0.078	0.079	17.91	449	-27.77	42.1	25.9	1.71	167.6	0.563	0.651	46.7
Erythrophleum chlorostachys	sap	Sept'10	0.50	95.4	1.85	1.94	0.052	0.055			-30.38	58.5	56.3	1.23	117.0	0.488	0.519	
Eucalyptus miniata	ad	May'11	0.61	68.1	0.74	1.08	0.090	0.132	24.54	744	-29.75	136.8	99.3	1.42	96.8	0.811	0.526	38.6
Eucalyptus miniata	ad	Sept'10	0.61	53.7	0.87	1.62	0.047	0.087			-30.71	83.9	69.3	1.19	63.9	0.814	0.523	
Eucalyptus miniata	sap	May'11	0.70	56.7	0.53	0.93	0.070	0.123	25.24	900	-29.14	112.0	81.8	1.47	83.2	1.001	0.557	32.5
Eucalyptus miniata	sap	Sept'10	0.70	56.4	0.76	1.35	0.040	0.071			-30.35	92.5	57.8	1.73	97.8	0.941	0.495	
Eucalyptus tetrodonta	ad	May'11	0.48	42.9	0.56	1.30	0.037	0.086	17.10	547	-27.75	92.3	52.2	1.76	75.5	0.712	0.621	38.2

Eucalyptus tetrodonta	ad	Sept'10	0.48	51.7	0.78	1.52	0.043	0.083			-27.85	77.4	48.5	1.52	78.4	0.708	0.551	
Eucalyptus tetrodonta	sap	May'11	0.50	42.2	0.50	1.19	0.044	0.104	19.38	613	-28.55	73.8	31.0	2.19	92.3	1.135	0.698	37.1
Eucalyptus tetrodonta	sap	Sept'10	0.50	57.0	0.89	1.55	0.044	0.078			-29.23	43.4	28.1	1.60	91.0	0.721	0.532	
Grevillea decurrens	ad	May'11	0.43	101.8	0.83	0.82	0.051	0.050	14.49	411	-28.41	51.1	38.1	1.53	155.5	0.844	0.607	34.3
Grevillea decurrens	ad	Sept'10	0.43	121.5	0.97	0.80	0.064	0.053			-29.64	36.0	44.9	0.80	97.8	0.722	0.499	
Grevillea decurrens	sap	May'11		98.6	0.87	0.88	0.052	0.052	12.65	329	-27.47	66.8	50.2	1.78	175.8	0.927	0.611	34.2
Grevillea decurrens	sap	Sept'10		113.2	1.05	0.93	0.072	0.064			-29.76	38.3	27.7	1.55	175.5	0.999	0.503	
Grevillea pteridifolia	ad	May'11	0.81	56.2	0.65	1.15	0.045	0.080	18.21	789	-29.52	71.8	42.8	2.23	125.5	0.905	0.584	39.0
Grevillea pteridifolia	ad	Sept'10	0.81	46.5	0.76	1.64	0.056	0.120			-28.80	117.5	48.0	2.55	118.8	0.854	0.486	
Grevillea pteridifolia	sap	May'11	1.00	67.5	0.70	1.03	0.044	0.066	19.72	866	-28.66	71.6	19.2	3.74	252.7	1.076	0.531	35.0
Grevillea pteridifolia	sap	Sept'10	1.00	65.3	0.78	1.19	0.055	0.085			-29.99	107.9	33.1	3.46	226.1	0.916	0.459	
Lophostemon lactifluus	ad	May'11	0.42	90.3	0.90	1.00	0.065	0.072	17.81	541	-29.11	124.7	106.9	1.23	110.8	0.581	0.553	44.2
Lophostemon lactifluus	ad	Sept'10	0.42	69.5	0.89	1.28	0.056	0.081			-31.12	53.3	93.2	0.59	41.2	0.683	0.497	
Lophostemon lactifluus	sap	May'11	0.42	73.5	0.88	1.20	0.061	0.084	20.70	1268	-28.66	50.0	33.2	2.11	154.9	0.528	0.531	37.2
Lophostemon lactifluus	sap	Sept'10	0.42	61.6	0.77	1.25	0.038	0.062			-30.46	32.4	33.1	1.00	61.7	0.458	0.457	
Persoonia falcata	ad	May'11	0.25	61.4	0.43	0.70	0.053	0.086	18.73	762	-29.83	47.6	76.2	0.70	43.3	0.535	0.597	38.9
Persoonia falcata	ad	Sept'10	0.25	70.3	0.86	1.22	0.059	0.084			-31.53	50.2	49.5	1.08	76.2	0.729	0.544	
Persoonia falcata	sap	May'11	0.50	62.0	0.52	0.84	0.056	0.091	17.51	747	-30.29	42.9	62.8	0.66	41.1	0.517	0.594	31.0

Persoonia falcata	sap	Sept'10	0.50	57.7	0.96	1.66	0.044	0.076			-31.94	40.4	47.0	0.87	50.1	0.504	0.480	
Petalostigma pubescens	ad	May'11	0.27	86.6	1.04	1.20	0.060	0.069	12.38	392	-28.59	31.6	30.8	1.09	94.3	0.864	0.668	43.8
Petalostigma pubescens	ad	Sept'10	0.27	96.4	0.67	0.70	0.034	0.035			-30.66	40.7	51.1	0.95	91.6	0.713	0.632	
Petalostigma pubescens	sap	May'11	0.44	74.5	1.10	1.48	0.060	0.080	18.34	443	-28.56	42.4	23.6	1.82	135.8	0.831	0.640	34.8
Petalostigma pubescens	sap	Sept'10	0.44	90.5	0.77	0.86	0.036	0.039			-30.38	22.7	44.3	0.56	50.6	0.642	0.633	
Planchonia careya	ad	May'11	0.42	76.3	1.06	1.38	0.079	0.104	17.19	700	-27.31	83.9	72.9	1.26	96.5	0.563	0.513	47.2
Planchonia careya	ad	Sept'10	0.42	89.5							-28.72	54.7	58.4	0.94	83.9	0.526		
Planchonia careya	sap	May'11	0.40	77.8	0.91	1.17	0.075	0.096	15.45	454	-27.25	46.6	34.4	1.58	123.3	0.659	0.522	43.8
Xanthostemon paradoxus	ad	May'11	0.35	74.6	0.43	0.57	0.054	0.072	14.64	338	-27.28	66.6	110.5	0.59	43.7	0.515	0.656	62.2
Xanthostemon paradoxus	ad	Sept'10	0.35	70.2	0.72	1.02	0.046	0.066			-28.10	25.7	118.9	0.22	15.4	0.472	0.464	
Xanthostemon paradoxus	sap	May'11	0.40	79.2	0.43	0.54	0.063	0.079	16.23	537	-28.12	66.0	71.6	1.30	102.5	0.624	0.584	54.5
Xanthostemon paradoxus	sap	Sept'10	0.40	65.7	0.63	0.95	0.046	0.071				22.0	44.7	0.59	38.7	0.638	0.457	

## **Appendix 4. Phylogenetic analysis**

Editors and reviewers sometimes request authors apply "phylogenetic" analyses to their trait datasets in order to "correct" for the degree of relatedness among species. The idea here is that phylogenetic relatedness is a nuisance that needs to be, and can be, statistically controlled (Felsenstein 1985; Harvey and Pagel 1991). Others take a different view (Uyeda *et al.* 2018; Westoby *et al.* 1998; Westoby *et al.* 1995), viewing phylogenetic and "cross-species" analyses as complementary approaches that ask subtly different questions. A cross-species analysis asks (in this case) whether variation in growth rates of species that occur today in savanna vegetation can be understood as being driven more or less by particular plant traits. For that question, every species is an independent statistical replicate, irrespective or what phylogenetic structure connects them: they are each an independent item of evidence for what types of traits and growth rates are successful in that situation. By contrast, a phylogenetic analysis can be used to ask the question "*Have evolutionary divergences in growth rate and in trait X been associated with one another throughout evolutionary history, more often than <i>expected by chance alone?*". For this reason, phylogenetic methods are sometimes referred to as "correlated divergence" analyses (Moles *et al.* 2005; Westoby *et al.* 1998).

Here we ran a simple correlated divergence analysis based on phylogenetically independent contrasts or PICs (Harvey and Pagel 1991). First, a phylogeny was constructed that described the hypothesised evolutionary relationships among the 17 species (Fig. S3). Tree macro-topology was based on information from Angiosperm Phylogeny Website (www.mobot.org/MOBOT/research/APweb/) version 14 (July 2017). Within-Myrtaceae relationships were derived from (Wilson *et al.* 2001), and within-*Acacia* relationships were resolved with tribe-level information from www.worldwidewattle.com. Second, trait and growth rate (GR) values were calculated for each internal node as the arithmetic average of the trait values for the two daughter species or nodes originating from that node. Third, the set of PICs (divergences) was calculated, each contrast being the difference between the trait (or GR) values for the two nodes or species descending from the contrast-node. (The direction of subtraction in calculating contrasts is unimportant, providing all traits are treated in the same manner). A correlation coefficient was then calculated between the set of GR contrasts and those for each trait (Table S1), assuming N-1 degrees of freedom (Harvey and Pagel 1991) where N is the number of internal nodes providing contrasts (in this case 16). Figure S3. Hypothesised phylogenetic relationships among the 17 study species

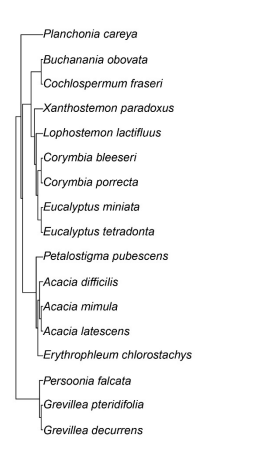


Table S1. Phylogenetically independent contrast analysis of stem diameter growth rates in relation to six plant functional traits.

Trait	r	Р
LMSM80	0.793	0.0001
a_LMSM	0.568	0.017
SLA	0.227	0.380
WD	-0.369	0.145
Asat	0.703	0.002
bark%	-0.613	0.009

From these results one can see that when considering the matter as evolutionary divergences the conclusion remains that variation in stem diameter growth rate is clearly connected to variation in leaf:stem allometry (LMSM80; a\_LMSM) and in photosynthetic rate (A<sub>sat</sub>), but not to variation in SLA or wood density (WD). In this analysis, divergences in growth rate

were negatively correlated with those in fractional bark allocation – a result we reported with some caution in the cross-species results, since it seemed potentially heavily influenced by *Acacia difficilis* (fast growth rate; very thin bark). The PIC analysis suggests that result was in fact more robust than we suspected.

## REFERENCES

Felsenstein J. (1985) Phylogenies and the comparative method. Am Nat 125, 1-15.

Harvey P. H. & Pagel M. D. (1991) *The comparative method in evolutionary biology*. Oxford University Press, Oxford.

Moles A. T., Ackerly D. D., Webb C. O., Tweddle J. C., Dickie J. B., Pitman A. J. & Westoby M. (2005) Factors that shape seed mass evolution. *P Natl Acad Sci USA* **102**, 10540-4.

Uyeda J. C., Zenil-Ferguson R. & Pennell M. W. (2018) Rethinking phylogenetic comparative methods. *Syst Biol*, in press. DOI 10.1093/sysbio/syy031.

Westoby M., Cunningham S. A., Fonseca C., Overton J. & Wright I. J. (1998) Phylogeny and variation in light capture area deployed per unit investment in leaves: designs for selecting study species with a view to generalizing. In: *Variation in Growth Rate and Productivity of Higher Plants* (eds H. Lambers, H. Poorter and M. M. I. van Vuuren) pp. 539-66. Backhuys Publishers, Leiden, The Netherlands.

Westoby M., Leishman M. R. & Lord J. M. (1995) On misinterpreting the 'phylogenetic correction'. *J Ecol* **83**, 531-4.

Wilson P. G., O'Brien M. M., Gadek P. A. & Quinn C. J. (2001) Myrtaceae Revisited: A Reassessment of Infrafamilial Groups. *Am J Bot* **88**, 2013-25.