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- 1 Nitrogen and phosphorus availability interact to modulate leaf trait scaling
- 2 relationships across six plant functional types in a controlled-environment study
- 3
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34 Summary

35 Nitrogen (N) and phosphorus (P) have key roles in plant leaf metabolism resulting in a strong 36 coupling of leaf-chemical composition traits to metabolic rates in field-based studies. 37 However, in such studies, it is difficult to disentangle the effects of nutrient supply per se on 38 trait-trait scaling relationships. 39 Our study assessed how high and low N (5 mM and 0.4 mM respectively) and P (1 mM and 2 40 μ M P respectively) supply on 37 species from six plant functional types (PFT) affected photosynthesis (A) and respiration (R) ({in darkness (R_{Dark}) and light (R_{Light})) in a controlled-41 42 environment. Low P supply only reduced increased scaling-exponents (slopes) of area-based the log-log -A-N 43 44 or R-N relationships when N supply was not limiting (high N), whereas there was no P-effect inunder low N supply-grown plants. By contrast, scaling-exponents slopes-of A-P and R-P 45 relationships were altered by both P and N supply, with lower N resulting in lower metabolic 46 rates for a given P. While NnNeither the R:A ratio nor light inhibition of leaf R the R Light: R Dark 47 ratio-wereas affected by nutrient supply. Light inhibition of respiration was 26% across 48 nutrient treatments; but heherbaceous species had exhibited , though some PFT differences 49 occurred.a lower degree of light inhibition than woody species. Slopes of A N and R N 50 51 relationships were lower in plants grown on low P and high N, whereas there was no P-effect 52 in low N-grown plants. As a function of leaf P, slopes of bivariate relationships were altered by 53 both P and N supply, with lower N resulting in lower metabolic rates for a given P. Neither the R:A ratio nor the RLight Reark ratio was affected by nutrient supply, though some PFT 54

55 differences-occurred.

- Because variations in N and P supply modulates leaf trait-trait relationships, it is vital to
 characterise the mechanistic impact of nutrient supply per se on these relationships given
 their application in the next generation of terrestrial biosphere models may need to consider
 how limitations in N and P availability affect trait-trait relationships when predicting carbon
 exchange-using leaf chemical composition need to incorporate nutrient variations across
 biomes.
- 62

- **Keywords:** day respiration, glasshouse, leaf economic spectrum, nitrogen, nutrient supply,
- **phosphorus,** photosynthesis, plant functional groups.

66 Introduction

67 Nitrogen and phosphorus are both essential plant nutrients involved in many important plant metabolic functions, including rates of light-saturated photosynthesis (A) and leaf respiration (R). 68 Strong positive relationships are found between both A, R and foliar N (Field & Mooney, 1986; 69 70 Evans, 1989; Ryan, 1995; Reich et al., 2008), reflecting the presence of N in key proteins. As a result, 71 N deficiencies can have negative effects on reduce capacity of photosynthetic and respiratory 72 enzymes, leading to lower leaf CO₂ exchange (Evans & Terashima, 1988; Noguchi & Terashima, 73 2006). Phosphorus (P) is a major component of nucleic acids, ATP, sugar phosphates and phospholipids, all of which are intrinsically related to photosynthesis and respiration (Rao & Terry, 74 75 1995). Consequently, P-deficient plants often exhibit: reduced stomatal conductance (Thomas et al., 2006); reduced ribulose bisphosphate (RuBP) regeneration, carboxylation activity and A 76 77 (Bloomfield et al., 2014; Ellsworth et al., 2015); and, lower allocation of N to RuBP 78 carboxylase/oxygenase (Rubisco) (Warren & Adams, 2002). Increased N allocation to non-79 photosynthetic components and/or inactivation of Rubisco are additional factors that might 80 contribute to lower A under P-deficiency (Stitt & Schulze, 1994). Phosphorus also plays a crucial role 81 in regulating glycolysis and mitochondrial electron transport (Plaxton & Podesta, 2006), with low P 82 supply having the potential to result in reduced R (Gonzalez-Meler et al., 2001). Limitations in P supply can lead to accumulation of starch (Hammond & White, 2008), -and-increases in dry mass 83 per unit leaf area (M_a) (Fyllas et al., 2009; Turnbull et al., 2016). Importantly, the effect of P-84 85 deficiency on such plant traits is likely to differ depending on whether P is more or less limiting than 86 N, reflecting Liebig's law of the minimum (e.g. Aerts *et al.*, 1992; Meir *et al.*, 2001)(Ågren *et al.*, 2012).-87

The coupling of metabolism to leaf chemical composition is important for predictions of 88 plant carbon exchange at a global scale (Sitch et al., 2008) (Kattge et al., 2009), with 'scaling 89 relationships' having been developed linking variation in metabolic rates being linked to variation 90 into M_a and mass-based leaf N (N_m) (Wright et al., 2004; Kattge et al., 2011). Plants growing on 91 92 nutrient-poor soils often have longer lived leaves, higher Ma values, lower Nm, and lower mass-93 based A (A_m) and leaf R in darkness (R_{Dark,m}) than plants growing on nutrient-rich soils (Wright & 94 Westoby, 2003) (Wright et al., 2003; Wright & Westoby, 2003). When trait-trait relationships are 95 compared for sites differing in nutrient availability, distinct patterns are often observed. For 96 example, the slope of bivariate linear relationships regressions linking A to leaf N_m are often lower

97 in plants growing on low P-soils (Reich & Walters, 1994; Denton et al., 2007; Kattge et al., 2009). Comparing tropical leaf traits at sites differing in P availability, Kattge et al. (2009) reported lower 98 rates of area-based maximum carboxylation rates (V_{cmax,a}) at given area-based N (N_a) in plants 99 100 growing on low P soils. Similarly, the slope (or scaling-exponent) of the log-log A-N relationships is 101 lower in tropical trees growing on low P soils than temperate trees growing on high P soils (Reich et 102 al., 2009). Moreover, Bahar et al. (2016) reported lower N-based rates of A and V_{cmax} in tropical 103 rainforest species growing on low P soils in Peru (compared to high P soils). Collectively, these field-104 based studies suggest that coupling of photosynthetic metabolism with N may depend on P-105 availability and hence a single 'universal' scaling relationships may not be realistic.

106

107 Relationships between leaf N and dark respiration are also affected by P-supply. Meir et al. 108 (2001) found that <u>area-based</u> leaf $R_{\text{Dark},a}$ at a given N_a being higher at P-limited sites. By 109 contrast, Rowland et al. (2016) reported higher area-based R_{Dark}-(R_{Dark,a}) at a given N_a in tropical 110 trees growing on low nutrient soils; this study also found that R_{Dark,a} was less affected by nutrient 111 limitations than A, resulting in higher R:A ratios in plants growing on nutrient-deficient soils. 112 Further, Reich et al. (2006) found that the log-log relationship between whole-plant R_{Dark,m} and N_m 113 was unaffected by nutrient supply, a finding also observed at the leaf-level (Atkin et al., 2013). In comparisons of log-log plots, Wright et al. (2004) showed intercept elevation (proportionality-114 <u>coefficient</u>) differences for among sites differing in nutrient availability but with constant -R_{dark,m}-N_m 115 slopes-scaling-exponents across sites. Notwithstanding the contrasting observations of these 116 117 studies, what is clear is that there is no single R_{Dark}-leaf N relationship. What is less clear, however, is what role soil N and P play in determining rates of metabolism and relationships linking A and R 118 119 to other leaf traits.

HAs noted above, some studies assessing impact of nutrient availability on trait-trait 120 relationships have used linear relationships fitted to untransformed data, while others have log 121 transformed data on both axes. (e.g. Atkin et al., 2013) [KC1] ow might deficiencies in N and/or P 122 123 affect log-log relationships fitted to multi-species data setsmade up of contrastingacross? When grown on limiting N supply, leaf N concentration can decrease to a greater extent than does the 124 125 rate of metabolism, underpinned by an increase in the proportion of N allocated to metabolic processes (Makino et al., 1994; Pons et al., 1994; Cheng & Fuchigami, 2000); in such cases, low N 126 127 treatment would result in higher rates of A per unit leaf N (i.e. higher PNUEAN). If the proportional

128 increase in $A_{\rm N}$ -is similar among species, then only the proportionality-coefficient of log-log A-N 129 plots would change (see Supporting Information Notes S1 for further details). However, if 130 proportional changes in $A_{\rm N}$ are greater in some species than others - although may $A_{\rm H}$ (e.g. Warren 131 et al., 2003; Pons & Westbeek, 2004; Hikosaka, 2010), then limitations in N supply might alter the scaling-exponent of log-log A-N relationships. (Warren et al. 2003, (Pons & Westbeek, 2004; 132 Hikosaka, 2010)For example, Warren et al. (2003) found that the maximal rate of carboxylation by 133 134 Rubisco (V_{emax}) per unit leaf N of Pinus sylvestris remained constant irrespective of N supply; others 135 have reported similar observations (Pons & Westbeek, 2004; Hikosaka, 2010). Thus, it is uncertain how limitations in N supply affect the slope and/or elevation of log log trait trait relationships for 136 data sets that include a wide range of species differing in leaf structure/chemistry and life history 137 138 characteristics Similar issues are at play for how low P supply impacts on log-log A-N relationships, 139 depending on: (1) the extent to which leaf phosphorus can be re-allocated from non-metabolic to 140 metabolic pools (Woodrow et al., 1984; Foyer & Spencer, 1986; Raghothama, 1999); (2) whether low P supply impacts on N allocation within leaves (Whitehead et al., 2005; Reich et al., 2009); and 141 142 (3) the extent to which these responses differ among species representative of on the leaf economic 143 spectrum (Wright et al., 2004). 144 Leaf respiration takes place both in the light (*R*_{Light}) and the dark, with *R*_{Light} typically lower than R_{Dark} (Way et al., 2015), even when accounting for re-fixation of respiratory CO₂ (Pärnik et al., 145 146 2007). Failure to account for light-induced inhibition of leaf R can lead to overestimation of 147 ecosystem respiration (Wehr et al., 2016). The extent of inhibition is highly variable, ranging from 148 80% inhibition through to cases where R_{Light} is 30% higher than R_{Dark} (Heskel et al., 2014; Way & 149 Yamori, 2014). While the mechanisms underpinning light inhibition are not fully understood, 150 photorespiratory metabolism is likely involved (Randall et al., 1990; Tcherkez et al., 2008). 151 Although the impact of nutrient gradients on light inhibition has been examined in a few field 152 studies (Atkin et al., 2013; Heskel et al., 2013), little if any attention has been given to the main and 153 interactive effects of N and P deficiency on the degree of light inhibition, or how N and P supply 154 affect scaling relationships linking leaf R_{Light} to related traits. In addition, no study has yet addressed the question of whether there are systematic differences in light inhibition among species or plant 155 156 functional types (PFTs). PFTs are used in terrestrial biosphere models (TBMs) as the starting point

157 for predicting metabolic rates across the globe, with TBM-dependent foliar N often being the

starting point for predicting rates of *A* and *R* (Clark *et al.*, 2011; Oleson *et al.*, 2013).

| 159 | | Given the importance of leaf trait relationships for the functioning of individual plants and | | | | | | | |
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| 160 | whole | ecosystems , with <u>strong tendencies for leaf N and P to co-vary in nature (</u> Sterner & Elser, | | | | | | | |
| 161 | 2002; | Reich & Oleksyn, 2004), it is difficult to isolate the effects of low P from low N in explaining | | | | | | | |
| 162 | <u>broad</u> | patterns of variation in relationships between metabolism and leaf nutrients. This is | | | | | | | |
| 163 | <u>especi</u> | ally so in field studies comparing species from a mong biomes, and among sites within | | | | | | | |
| 164 | individ | l <u>ual biomes (</u> Kattge <i>et al.,</i> 2009; Reich <i>et al.,</i> 2009; Domingues <i>et al.,</i> 2015) <u>. Given these</u> | | | | | | | |
| 165 | <u>challer</u> | nges, we used a factorial design using 37 species from six PFTs to isolate the individual and | | | | | | | |
| 166 | combined effects of N and P availability supply on a range of leaf traits associated with | | | | | | | | |
| 167 | photos | synthesis and respiration. Our study provides the first multi-PFT comparison assessing | | | | | | | |
| 168 | nutrie | nt impacts on leaf trait relationships under controlled environment conditions. and the fact | | | | | | | |
| 169 | that n | utrient availability varies markedly in nature, it is vital that the impacts of nutrient supply per | | | | | | | |
| 170 | se on l | eaf trait trait relationships be characterised. Past studies assessing impacts of N and P | | | | | | | |
| 171 | availat | pility on trait-trait relationships have been field-based and, as such, not able to definitively | | | | | | | |
| 172 | assess | the effects of nutrient supply on leaves. It was with this in mind that we conducted a | | | | | | | |
| 173 | contre | lled environment study using 37 species from six PFTs, with each species grown on high and | | | | | | | |
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| 174 | low N | and P supply. We tested the following hypotheses: | | | | | | | |
| 174 175 | low N 1. | and P supplyWe tested the following hypotheses: -Growth under low P supply will reduce <u>average rates of</u> leaf A and R <u>, with the inhibitory</u> | | | | | | | |
| 174 175 176 | low N 1. | and P supply. We tested the following hypotheses: -Growth under low P supply will reduce <u>average rates of</u> leaf A and R <u>, with the inhibitory</u> <u>effect being greater</u> in high N <u>than low N</u> -grown plants , but have limited <u>minimal</u> effect in | | | | | | | |
| 174 175 176 177 | low N 1. | and P supply. We tested the following hypotheses: -Growth under low P supply will reduce <u>average rates of</u> leaf A and R, <u>with the inhibitory</u> <u>effect being greater</u> in high N <u>than low N</u> -grown plants , but have limited <u>minimal</u> effect in plants grown on<u>when N is limiting limitingN, reflecting Liebig's law of the minimum.</u> | | | | | | | |
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| 174 175 176 177 178 179 | low N 1. | and P supply. We tested the following hypotheses: -Growth under low P supply will reduce <u>average rates of leaf A and R, with the inhibitory</u> <u>effect being greater</u> in high N <u>than low N</u> -grown plants , but have limited <u>minimal</u> effect in <u>plants grown onwhen N is limiting limitingN, reflecting Liebig's law of the minimum.</u> <u>Similarly, the deleterious effect of low N supply on A and R will be less pronounced when P</u> <u>is limiting.</u> Such effects are expected to be similar among PFTs. <u>Similarly, the deleterious</u> | | | | | | | |
| 174 175 176 177 178 179 180 | low N 1. | and P supply. We tested the following hypotheses: -Growth under low P supply will reduce <u>average rates of leaf A and R, with the inhibitory</u> <u>effect being greater</u> in high N <u>than low N</u> -grown plants , but have limited <u>minimal effect in</u> <u>plants grown onwhen N is limiting limitingN, reflecting Liebig's law of the minimum.</u> <u>Similarly, the deleterious effect of low N supply on A and R will be less pronounced when P</u> <u>is limiting.</u> Such effects are expected to be similar among PFTs. <u>Similarly, the deleterious</u> <u>effect of low N supply on A and R will be less pronounced when P is limiting.</u> | | | | | | | |
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| 189 | 4. | <i>R</i> : <i>A</i> ratios will be greater for nutrient-limited plants for <u>compared to</u> plants grown on high N |
|-----|----|--|
| 190 | | and P supply, reflecting the greater inhibitory effect of limiting nutrient supply on |
| 191 | | photosynthesis compared to that of respiration (Evans & Terashima, 1988). |
| 192 | 5. | (Shapiro <i>et al.,</i> 2004; Tcherkez <i>et al.,</i> 2008; Ayub <i>et al.,</i> 2011; Crous <i>et al.,</i> 2012; Griffin & |
| 193 | | Turnbull, 2013) Nutrient treatments that reduce rates of photorespiration (i.e. limitations in |
| 194 | | <u>N and/or P supply) willmay will result in lower There is no systematic difference in the</u> |
| 195 | | degrees of light inhibition of leaf <i>R</i> -among species or nutrient treatments (see SI Notes S2 |
| 196 | | <u>for further details)</u> |
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- 198

199 Materials and Methods

200

201 Plant material and experimental design

The study comprised 20 woody and 17 herbaceous species from 17 families, representing six PFTs: 202 203 broadleaved trees (BITBLT), broadleaved shrubs (S), coniferous needle-leaved trees (NITNLT), C₃ 204 grasses (C3G), C₃ herbs (C3H) and C₄ grasses (C4G) (Table S1). Four Protaceae species which can access P more efficiently under low P (Lambers et al., 2010) were included in the woody species 205 group (Table S1). The woody species were planted in November 2008, and measured starting in 206 207 January 2009, whereas the herbaceous species were planted in November 2009 and measured in 208 January 2010 onwards. Woody plants were sourced mainly from the Wagga Wagga Forestry 209 Nursery (Wagga Wagga, NSW, Australia) whereas the herbaceous species were sourced from the Woodbridge nursery (Woodbridge, Tasmania) or grown from seeds (Nindethana Australian Seeds, 210 211 Albany, WA, Australia). Twenty-four plants of each species were potted in sterilised sand in 50 cm 212 tall PVC, 9 cm diameter, 3.18 L pipes_to achieve six replicates per nutrient treatment. Two adjacent 213 glasshouses were used with three replicate blocks in each glasshouse. Species and nutrient 214 treatments were randomly allocated within each block. Species were grown in the glasshouse 215 under natural light conditions at 25°C/18°C (day and night).

Nutrient treatments represented four combinations of different N and P concentrations: High
N-High P (H_NH_P), High N-Low P (H_NL_P), Low N-High P (L_NH_P) and Low N-Low P (L_NL_P). The H_NH_P
solution was based on Hoagland No. 1 solution (Hoagland & Arnon, 1950) with the following
macronutrient composition: 1 mM KH₂PO₄, 5 mM KNO₃, 0.07 mM CaCl₂, 0.45 mM MgSO₄.7H₂O. The

220 Whereas the final concentrations of 'high N' solutions were 5 mM KNO₂, 'low N' solutions had a 221 final-concentration of 0.4 mM KNO₃ (modified after Atkinson *et al.*, 2007). The 'high P' solutions 222 contained 1 mM KH₂PO₄ (Edwards et al., 2006) whereas 'low P' had 2.0 µM KH₂PO₄ to limit storage 223 of P in the vacuole as a buffer (after Campbell & Sage, 2006). Thus, N:P supply ratios varied from 5:1 224 for H_NH_P to 2500:1 for H_NL_P, 0.4:1 for L_NH_P and finally 200:1 for L_NL_P. Here, our aim was to achieve 225 changes in foliar chemistry and function, that was also reflected in rates of leaf plant metabolism, 226 rather than mimic naturally occurring N:P ratios as such rather than mimic natural soil N:P ratios. 227 Each nutrient solution was balanced for cations and contained the same amount of micronutrients 228 and FeEDTA. Micronutrients were constant in all solutions: 4.2 μ M B, 1.2 μ M Mn, 0.8 μ M Zn, 0.03 229 μ M Cu, 0.04 μ M Mo and 0.01 μ M Co. Iron was added as ferric EDTA to a level of about 8 μ M Fe. 230 These micronutrient concentrations were one-tenth of the recommended Hoagland solution 231 because full strength has resulted in toxic symptoms (Leggett, 1971). Nutrient solutions were made up in 200 L black storage tubs, which were refilled regularly. Once per day, nutrient solutions were 232 pumped out of 200 L tubs and delivered to the pots using irrigation tubing and drippers at 20 ml per 233 234 minute. Plants received on average about 120-150 mL solution each day and were watered at the 235 end of each day.

236

237 Gas exchange measurements

238 Gas exchange was measured between 9 am and 2:30 pm with portable infrared gas analysers (LiCor 239 6400, LiCor Inc., NE, USA). 873 light-response curves were conducted on the most recently fully 240 developed leaves in each treatment-at a single timepoint, starting with light-saturating conditions 241 (irradiance: 1800 μmol m⁻² s⁻¹; flow: 500 μmol s⁻¹; leaf temperature: 25°C; reference CO₂: 400 μmol 242 mol⁻¹). After these initial conditions, light was decreased to 1500 µmol m⁻² s⁻¹ and then 100 µmol m⁻ 2 s⁻¹, and finally in steps of 5-10 μ mol m⁻² s⁻¹ to achieve estimates of respiration during the day 243 (RLight) using the Kok method (Kok, 1948). RDark was obtained 10 minutes after the light was turned 244 245 off (flow 300 µmol s⁻¹) to avoid post-illumination transients (Atkin et al., 1998). For each leaf, 246 measurements were made at a single time point.

The Kok effect refers to the break in the slope of plots of net CO_2 uptake vs irradiance near the light-compensation point (Ayub *et al.*, 2011). Using this approach, a regression line is extrapolated from the linear part of the light-response curve (over the irradiance range 20-60 µmol m⁻² s⁻¹), with the *y*-axis intercept representing R_{Light} . R_{Light} was corrected for changes in C_i (intercellular CO_2 concentration) associated with changes in-irradiance (Kirschbaum & Farquhar, 1987). All curves were
analysed using R (R Core Development Team, 2015).

253

254 Structural and chemical leaf composition

Leaves were collected after each light response curve and the leaf area inside the leaf cuvette was measured before drying at 70°C for at least 24 h to determine leaf mass per area (*M*_a). Total N and P were determined via Kjeldahl digestion (Allen, 1974), with digests analysed using a flow injection analyser (Lachat instruments, Loveland, Colorado, USA) for N and P using the indophenol blue (Scheiner, 1976) and ammonium molybdate methods (John, 1970) respectively. Leaves from 16 woody species were analysed for soluble sugars and starch according to Loveys *et al.* (2003).

261

262 Statistical <u>a</u>Analyses

Initial data exploration revealed cases of outlying (but not erroneous) values, unequal variation 263 among treatment classes and non-normal distributions. Many of our variables, notably leaf N and 264 P, showed distributions that were heavily right-skewed and therefore . For consistency of 265 presentation and interpretation all variables have been log-transformed in the analyses that follow. 266 Log transformation was the best option chosen from a number of alternative methods (e.g. square 267 root and inverse) and that with improvement applied both to the underlying variables and to model 268 residuals KC2].__Because most variables followed a rightly skewed distribution, values were log-269 270 transformed before screening for outliers and subsequent analysis, as outlined in SI SM1. Data were filtered for outliers identified as those falling beyond twice the interguartile range with for a given 271 variable. Ffilters for separate variables were applied to the dataset in the following order: Pm, Nm, 272 273 Ma, R_{Dark,m}, and A_m. Application of these filters resulted in a total of 15 outliers/rows being removed 274 from the dataset. Whenever an outlier was identified, the entire observational row was removed. Application of these filters resulted in a total of 15 outliers being removed from the dataset. After 275 filtering, 37 species, six PFTs, and four nutrient treatments remained (Table S1) totalling 873 276 277 measurements (with 148 PFT species by treatment combinations). Woody species represented 472 278 observations. Further filtering was necessary for RLight, removing three species (Banksia spinulosa, 279 Pinus halepensis and Banksia integrifolia) due to inadequate replication or measurements with 280 negative R_{Light} values (i.e. positive y-axis interceptselevation from Kok estimates – see 281 abovemethod). Data falling beyond twice the interquartile range resulted in further removal of

282 twelve rows. Hence the R_{Light} dataset consisted of 710 values. Calculations of means and ANOVA 283 tests were conducted in JMP (SAS Institute, NC). We used individual plant data when comparing bivariate relationships across the six PFTs and four nutrient treatments, consistent with the 284 285 rationale used in past studies (Reich et al., 2008; Xiang et al., 2013). Standardized major axis (SMA) 286 analysis was used to determine the best-fitting lines (α = 0.05) for the key log-log trait-trait 287 relationships (Falster et al., 2006; Warton et al., 2006; Warton et al., 2012) and leaf traits. We 288 tested for differences among nutrient supply treatments and PFT classes. SMA, descriptive statistics 289 and filtering Data exploration, screening for outliers and SMA analysis was performed using the R language and environment were done in R (R Development Core Team, 2015) and the smatr 290 291 package (Warton et al. 2012). Multiple comparison tests employed in the smatr package are explained in Appendix D of Warton et al. (2006) - see SI Notes S1 for further details. - Data are to 292 be-freely available and published on the Australian Research Data portal (-doi to be inserted; here). 293 294

295 Results

296

297 Leaf trait variation

298 Across the entire dataset, there was large variation in each of the selected structural, chemical 299 composition and physiology traits, reflecting the diverse range of species (Fig.s 2 and 3). For all 300 traits, there were significant differences among PFTs (P < 0.0001, Table 1). When averaged across 301 all treatments, mean M_a for each across PFTs varied five-fold, with needle-leaf coniferous (NLT) trees <u>having</u> exhibited ing the highest values (225 \pm 10 g m⁻²) while C₄-grasses (C4G) showed the 302 lowest M_a (39 ± 2 g m⁻²) (Fig. 2ac, Table S2). This pattern was also reflected in N_a (3.6 ± 0.3 g m⁻² in 303 NIT and 0.7 ± 0.02 g m⁻² in C4G) and P₂ (0.93 ± 0.13 g m⁻² in NLT) with the average of broadleaved 304 shrubs and trees (BLT) being 0.37 ± 0.05 g m⁻² while grasses exhibited mean P_a values of 0.18 g m⁻² 305 (C3G) and 0.13 g m⁻² (C4G) (Fig. 2b). Overall, woody species exhibited higher N_a than their 306 307 herbaceous counterparts, largely reflecting differences in Ma among PFTs and PFT-dependent differences in N_m (Table S2). Highest P_a were again found in NIT (0.93 \pm 0.13 g m⁻²) with the average 308 of broadleaved shrubs and trees being 0.37 ± 0.05 g m⁻², while grasses exhibited mean P_a values of 309 0.18 g m⁻² (C3G) and 0.13 g m⁻² (C4G)(Fig. 2c). N:P ratios showed significant differences between 310 311 each treatment (Table 2) with N:P above 15 in broad-leaved trees (BLT), shrubs (S) and C₃ grasses 312 (C3G) ; PFT differences are shown in (Table S2). Mass-based values of both nitrogen (N_m) and

- phosphorus (P_m) were lowest in S and NLT and highest in C4G and C3H (Table S2, Fig. 2). Within
- 314 each PFT, high and low nutrient treatment concentrations were reflected in the leaf traits
- Summarised in Table S2 and Figure S1. Mean A_a values were similar across PFTs, ranging from 10.1 to 15.3 µmol m⁻² s⁻¹ for C3G and S, respectively (Fig. 3a2d); A_m was highest in C4G and C3H and lowest for NITNLT (data not shown). Highest rates of $R_{\text{Dark},a}$ were found for S and BITBLT (mean 1.13 µmol m⁻² s⁻¹) with the lowest means in C3G (0.49 µmol m⁻² s⁻¹) (Fig. 3b2e). By contrast, $R_{\text{Light},a}$ was highest in NITNLT and C4G (mean 0.95 µmol m⁻² s⁻¹) and lowest in C3G (0.44 µmol m⁻² s⁻¹) (Fig. 3c2f, Table S2).
- 321

322 Impacts of nutrient supply on mean values

Table S2 shows mean values for each PFT both among and across treatments, while Table 1 shows 323 324 the two-way ANOVA assessing main and interactive effects of PFT and treatment. Growth of plants 325 on contrasting N and P supply resulted in significant changes in foliar N and P, both on a mass and 326 area basis (Table 1). Although there were differences in leaf chemistry among PFTs (see above and 327 Table S2), overall treatments receiving high N supply exhibited the highest N_m (24.5 mg g⁻¹, 328 averaged across both H_NH_P and H_NL_P; Table 2) compared to the low N treatments with an average N_m of 15.9 mg g⁻¹. Low P treatment reduced N_m and N_a with high N supply, whereas under low N 329 supply, P supply did not impact on leaf N (Table 2, Fig. S1). The highest P_m were found in high P 330 supply both for high N and low N conditions, with the lowest P_m found in high N-low P supply 331 (H_NL_P). Low P supply resulted in a decline in P_a on both high and low N supply. Interestingly, low N-332 333 grown plants exhibited higher P_m compared to high N-grown plants (Table 2, Fig. S1). For woody 334 PFTs (i.e. **BIT**<u>BLT</u>, **NIT**<u>NLT</u> and S), there was no effect of low N on area-based soluble sugars (Table</u> 335 3). Under high N conditions, low P supply led to a 60% increase in starch and a 15% increase in total 336 non-structural carbohydrates (TNC) per unit leaf area in the woody PFTs (Table 3 and SI Table S3). 337 Whereas N and P supply had large effects on foliar chemistry, nutrient supply had no effect on M_{a} , averaged across all PFTs (Tables 1 and 2, Fig. S1a). Moreover, correcting M_a values for 338 339 accumulation of TNC did not alter this conclusion, when considering woody species alone. The 340 stability of *M*^a suggests we can equally assess the effect of nutrient supply on metabolic traits on 341 either an area- or mass-basis.

Nutrient supply had a marked effect on fluxes of area- and mass-based rates of
 photosynthesis and respiration (Tables 1 and 2). The highest rates of both A and R occurred in the

344 H_NH_P treatment and all other treatments exhibited similar, lower rates of A_a and A_m. - A_a and A_m 345 were reduced to a similar extent when either N or P were in low supply (reduced by 20% for A_a) and 346 with no further reduction when both N and P were in low supply (Table 2). Similarly, for R_{Dark} (both 347 on area and mass bases) low N treatment resulted in significant declines (-27%), as did low P supply 348 (-16%) compared to the H_NH_P treatment. However, unlike photosynthesis, L_NL_P supply resulted in 349 further significant declines in R_{Dark} compared to when either N or P were in low supply (Table 2). 350 R_{Light} showed similar patterns as R_{Dark} , with lowest rates in L_NL_P. Compared to the H_NH_P treatment, 351 R_{Light} exhibited reduced rates in low N regardless of P supply (Table 2). Hence, there was no effect of 352 P deficiency on both photosynthesis and respiration (in either the dark or light) in low N-grown 353 plants (both L_NH_P and L_NL_P), with P deficiency only affecting fluxes in high N-grown plants (Table 2). 354 When averaged across all PFTs, Pphotosynthetic N use efficiency (A_N) was 11% enhanced 355 when plants experienced low N supply (compared to H_NH_P). By contrast, A_N was reduced by low P 356 supply only reduced average A_N in plants grown on high N (-20%) but not when N was limiting (Table 2). For respiration, N and P deficiencies had little effect on N-based rates of R_{Dark} and R_{Light} 357 358 (Table 2). For both A and R, rates expressed on a foliar P basis were significantly higher in low P-359 grown plants, irrespective of N supply (Table 2), suggesting greater P-use efficiency under low P conditions. 360

N-deficiency had no effect on respiration:photosynthesis ratios (i.e. the inhibitory effect of low N on A and R was similar) when P was non-limiting (Table 2). By contrast, under P-deficient conditions, low N supply resulted in a slight decrease in the R_{Dark} :A and R_{Light} :A ratios, reflecting the significant decline in respiration alone (i.e. no change in A) under L_NL_P compared to H_NL_P conditions (Table 2). Finally, nutrient availability had no effect on the R_{Light} : R_{Dark} ratio (Table 2), with the average the R_{Light} : R_{Dark} ratio among treatments being 0.74 µmol m⁻² s⁻¹ (n = 710). Thus, light inhibited leaf respiration by 26% across the entire dataset.

Table 1 also provided insights into whether the effect of N supply on each trait differed among PFTs. The absence of a significant interaction term indicated that all PFTs were similarly affected by availability of N and P for M_a , A_a and $R_{\text{Light},a}$ (Table 1). For $R_{\text{Dark},a}$, the effect of treatment did differ among PFTs (Table S2); in general, low N supply reduced *R* in <u>BITBLT</u> and S but not in NITNLT or grasses. Growth on H_NL_P supply led to significantly lower $R_{\text{Dark},a}$ in <u>BITBLT</u> alone, with no effect of P supply on other PFTs (Table S2). At low N, low P supply only reduced rates of $R_{\text{Dark},a}$ in C3G. Taken together, these results point to a general pattern of low N supply having its greatest impact on metabolic traits when P is non-limiting (<u>thus-supporting Hypothesis 1</u>) and that the
 impact of nutrient supply is more commonly seen in broadleaved trees and shrubs than the other
 PFTs.

378

379 *Leaf trait relationships*

The above analyses point to increased rates of A_N under low N supply (irrespective of P supply), and reduced A_N when P is limiting (for high N plants only), suggesting nutrient-mediated shifts in relationships between photosynthesis and N. We explored whether such changes were associated with shifts in proportionality-coefficient (i.e. elevation of the log-log relationships) elevation and/or scaling-exponents (i.e. slope) slopes of the A_a -N_a relationship (as hypothesized in Fig. 1) and other area-based trait relationships via standardized major axis (SMA) analyses.

386 Figure 4-3 shows bivariate relationships among N_a , P_a and M_a , with the corresponding SMA 387 results shown in Table 4. While substantial scatter occurred within each bivariate relationship, significant positive relationships were apparent for each separate nutrient combination. For the Na-388 *M*^a relationship, there were no significant differences in <u>scaling-exponents</u> of the four 389 390 treatments, but significant differences were found in the proportionality-coefficient elevation of 391 the fitted lines (Fig. 4a3a, Table 4). Here, N_a at a given M_a was significantly higher in the H_NH_P 392 treatment, followed by H_NL_P , which in turn was higher than L_NH_P and L_NL_P relationships (i.e. H_NH_P > 393 $H_NL_P > L_NH_P = L_NL_P$). Thus, leaves consistently accumulated more leaf N for a given M_a under high N 394 conditions, with P deficiency slightly decreasing how much N accumulated at given M_a . For the P_{a^-} 395 *M*_a relationship (Fig. 3b), there was a significant difference in slopes among the treatments because 396 $L_{N}H_{P}$ had a steeper greater scaling-exponent slope-compared to the other treatments (Table 4); 397 Figure 4b-3b also shows that P_a values at a given M_a were lower in the two low-P treatments than 398 the two high-P treatments. When considering the Na-Pa relationship, SMA analyses revealed 399 different ces among the scaling-exponents slopes, with $L_N H_P$ exhibiting a flatter N_a - P_a response than 400 the other treatments. N_a at a given P_a was highest in H_NL_P plants (Fig. $4c_{3c}$), followed by L_NL_P and 401 H_NH_P grown plants, which in turn were markedly higher than L_NH_P plants. Thus, the treatments 402 differed in a manner consistent with N:P values (Table 2).

We now explore how the treatments affected bivariate relationships linking A_a to N_a , P_a and M_a (Fig. 54, Table 4). For A_a - N_a relationships (Fig. 5a4a), showed different slopes. Ssimilar growth on low N supply resulted in greater scaling-exponents slopes-compared to high N grown plants,

406 irrespective of P supply (Table 4). This finding reflected two interesting outcomes. Firstly, for For 407 plants grown on high P supply, growth on low N (L_NH_P) only increased A at a given N (i.e. increased A_N) in species with inherently high N_a values, with little effect of N supply on A_N in low N_a species. 408 409 By contrast, when P supply was limiting, low N supply had little effect on A_N in high N_a species, but increased A_N in low N_a species. Thus, the effect of N supply on A_N differs among the contrasting 410 411 species used in our study, and depends on whether P supply was limiting. were exhibited by the 412 two low N treatments (i.e. L_NH_P & L_NL_P) but plants grown on H_NL_P exhibited a much steeper slope 413 than the other treatments. HenceUnderpinning this observation was that, with high N availability, 414 growth on-low P supply reduced A_N in low N_a species, but had little effect on A_N in high N_a species. 415 had a marked inhibitory effect on photosynthesis, but when N was limiting, P availability had 416 no effect.—For A_a - P_a relationships (Fig. <u>5b4b</u>), H_NL_P grown plants exhibited a significantly steeper 417 greater scaling-exponent slope-than the other treatments (Table 4), with A_a at a given P_a differing 418 between high and low N treatments. For the low P-grown plants, those provided with high N 419 exhibited generally greater A_a at a given P_a than low N-grown plants (i.e. comparing green with 420 yelloworange, and blue with red black lines symbols in Fig. 5b4b; note: Aa-Pa relationship was not 421 significant for $L_N L_P$ plants). Finally, when considering the relationship between A_a and M_a , no 422 relationship was found for high N-grown plants (Table 4), with a common relationship being shared 423 by low N grown (L_NH_P and L_NL_P) plants (grey line in Fig. 4c). Taken together, photosynthesis was 424 reduced with low P supply in high N-grown plants with respect to the A_a-N_a relationship, whereas 425 but with low N supply leading to reduced rates of A_a for a given P_a were reduced, both in high and 426 low P availability.

427 For relationships linking leaf respiration to other traits Na, Pa and mass:area Ma, we found that 428 treatment had no effect on the <u>scaling-exponents</u> slopes of *R*_{Dark,a}-N_a, but did alter the <u>scaling</u>-429 exponents slopes of the R_{Light,a}-N_a relationship (Fig. 654, Table 4). For R_{Dark,a}-N_a, plants grown on 430 high N, P-deficiency resulted in lower $R_{\text{Dark},a}$ at a given N_a (i.e. lower proportionality-coefficient)₇ 431 particularly in plants where foliar N concentrations were low. By contrast, at low N supply, 432 availability of P had no effect on $R_{\text{Dark},a}$ at a given N_a (Fig. $\frac{6a_{5}4a_{1}}{6a_{5}4a_{1}}$); this was also true for the $R_{\text{Light},a}$ -433 N_a relationship (Fig. 6d54dg). Thus, as was the case for photosynthesis, P availability only modifies 434 the relationship between leaf respiration and N_a when N supply is not limiting. Growth on low P resulted sexhibited in higher R_{Dark,a} and R_{Light,a} at a given P_a, irrespective of N 435

436 supply (Fig. 6b54eb, eh). Moreover, limitations in the availability of N resulted in lower R_{Dark,a} and

R_{Light.a} at a given P_a; this was true for plants grown on low and high P. Thus, relationships linking leaf 437 438 respiration to P_a are-were influenced not only by the availability of P, but also by N supply. Finally, 439 whereas no significant relationships were found when considering $R_{\text{Light,a}}$ as a function of M_a (Fig. 440 6f54fi, Table 4), R_{Dark,a}-M_a relationships were significant for all four treatments (albeit with a low R²; 441 Fig. 6c54fe, Table 4). Interestingly, expressing rates on a mass basis resulted in highly significant 442 RLight, and RDark, and Ma (Fig. 4f), nutrient (SI Fig. S2, Table S4). For RDark, a-Ma (Fig. 4f), nutrient 443 treatment had no effect on relationship slopes, with elevations being ranked from highest to 444 lowest: $H_NH_P > H_NL_P = L_NH_P > L_NL_P$. Thus, the general pattern was one of $R_{Dark,a}$ at a given M_a 445 decreasing as nutrient availability declined (Table 2).

446 Nutrient availability did not significantly affect rates of R_{Dark,a} at a given A_a indicated via a common R_{Dark,a}-A_a relationship across all four treatments (shown by the common grey regression 447 line in Fig. 7a65a). -While there were significant differences in scaling-exponents slopes-when 448 449 considering $R_{\text{Light,a}}$ - A_a (Fig. $\frac{7b_65b}{7b_65b}$, Table 4), the overall pattern was similar to that seen in $R_{\text{Dark,a}}$ - A_a 450 Figure 7a. On first inspection, the finding that $R_{\text{Dark},a}$ - A_a was unaffected by nutrient supply contrasts 451 with the comparison of means shown in Table 2, where limitations in N supply reduced R_{Dark,a}:A_a 452 ratios in low P-grown plants. However, given the scattered nature of the bivariate plots and the 453 slight differences in R_{Dark,a} at a given A_a among the treatments (Fig. 765a), caution is needed when 454 considering small changes in mean-alone data shown in Table 2. Thus, the most parsimonious 455 conclusion is that nutrient supply had little or no effect on R_{Dark,a}:A_a ratios. This was also true for the 456 <u>*R*_{Light,a}-*R*_{Dark,a} relationships among four treatments</u>, with nutrient supply having no effect (Fig. 265ca; Table 44); consequently, the degree of light inhibition was unaffected by nutrient supply (~ 457

458 <u>26% inhibition; Table 2).</u>

459 While nutrient supply had no significant effect on the slope in Figure 7a6a, there remained considerable scatter in the overall R_{Darka}-A_a relationship. Thus, other factors distinct from nutrient 460 supply must be responsible for the scatter. Differences between PFTs woody and herbaceous 461 species showed is an obvious candidate via no significant difference in slopes scaling-exponents (P = 462 0.37) for R-A relationships, w-but significant differences in elevation (Table S5). Wwoody species 463 exhibiteditinged a higher proportionality-coefficient elevation (i.e. less negative y axis intercept) 464 compared to herbaceous PFTs (Fig. S3a65d, Table S54). Hence, for a given Aa, woody species overall 465 had higher dark respiration compared to herbaceous species, with R_{Dark,a} at a given A_a being highest 466 467 in shrubs and broadleaf trees followed by C₃ herbs, while needle leaf NLT trees, C₃ and C₄ grasses

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| 468 | exhibited the lowest $R_{\text{Dark},a}$ at a given A_a (Fig. S3, Table S5). For $R_{\text{Light},a}$ - A_a (Fig. 5e), there was no |
| 469 | difference between herbs and woody species in the scaling-exponent slope or proportionality- |
| 470 | <u>coefficient elevation of the <i>R</i>_{Light,a}-A_a-relationship-(but there was a shift moving along the same</u> |
| 471 | relationship); this is -in-contrast to R _{Dark,a} -A _a (Fig. 65ed) where woody species exhibited a higher |
| 472 | proportionality-coefficient than herbs. For the R_{Light} - R_{Dark} relationship, hHerbaceous species |
| 473 | showed exhibited a higher greater scaling-exponent slope in the R _{Light} -R _{Dark} relationship-compared to |
| 474 | woody species (Fig. 56f, Table S5) with tThe lowest degree of light inhibition of leaf respiration |
| 475 | was found in C4G (as shown by the highest <i>R</i> Light: <i>R</i> Dark_slope, Table S5) and the highest degree of |
| 476 | light inhibition found in NLT (lowest R _{Light} :R _{Dark} scaling-exponentslope). |
| 477 | |
| 478 | with R _{Dark,a} at a given A _a being highest in shrubs and broadleaf trees followed by C ₃ herbs, |
| 479 | with needle-leaf trees, C ₃ -and C ₄ grasses exhibiting the lowest R _{Dark,a} at a given A _a (Fig. 8, Table 5). |
| 480 | SMA analyses on woody and herbaceous PFTs indicated no significant difference in slopes (P = 0.37) |
| 481 | but different elevations, with woody species exhibiting a higher elevation (i.e. less negative y-axis |
| 482 | intercept) compared to herbaceous PFTs (Fig. S3a, Table 5). Hence, for a given A _a , woody species |
| 483 | had higher dark respiration compared to herbaceous species. |
| 484 | For R _{Light,a} -A _a , there was less consistency in the bivariate relationships among PFTs (Table 5). |
| 485 | When combining PFTs into woody and herbaceous categories, then there was no difference in slope |
| 486 | or intercept of the R _{Light,a} -A _a -relationship (but there was a shift moving along the same relationship) |
| 487 | in contrast to R _{Dark,a} -A _e (Fig. S3b). Lastly, Figure 9 shows A _{Light,a} -A _{Dark,a} -relationships among four |
| 488 | treatments, with nutrient supply having no effect (Fig. 9a; Table 4); consequently, the degree of |
| 489 | light inhibition was unaffected by nutrient supply (~ 26% inhibition; Table 2)However, the lowest |
| 490 | degree of light inhibition found in C4G (highest R _{Light} :R _{Dark} slope, Table 5) and the highest degree of |
| 491 | light inhibition found in NIT <u>NLT</u> (lowest R _{Light} :R _{Dark} slope). Combining PFT into woody and |
| 492 | herbaceous species indicated a higher R _{Light} :R _{Dark} slope in herbaceous compared to woody species |
| 493 | (Fig. 9b, Table 5). |
| 494 | |
| • | |

- 496 Discussion
- 497

498 **Study rationale and Eextent of nutrient limitations**

499 Nitrogen and phosphorus exert strong and fundamental controls on metabolic performance of leaves, as seen across both natural and fertilized soil gradients (Reich & Schoettle, 1988; Cordell et 500 501 al., 2001; Whitehead et al., 2005). However, given strong tendencies for leaf N and P to co-vary in 502 nature (Sterner & Elser, 2002; Reich & Oleksyn, 2004; Ågren et al., 2012), it is difficult to isolate the 503 effects of low P from low N in explaining broad patterns of variation in relationships between metabolism and leaf nutrients. This is especially so in field studies comparing species from among 504 biomes and among sites within individual biomes (Kattge et al., 2009; Reich et al., 2009; Domingues 505 et al., 2015; Bahar et al., 2016). Given these challenges, we used a factorial design to isolate the 506 individual and combined effects of N and P availability on a range of leaf traits associated with 507 photosynthesis and respiration. Our findings show that both N-and P-supply can mediate leaf trait-508 trait relationships, suggesting that the next generation of Earth System Models should consider 509 510 how limitations in N and P supply in individual biomes may affect trait relationships (Coll et al., 511 2012).

512 In setting up our experiment, there was a risk that the low P and N treatments would not result in P- and N-deficient phenotypes (in terms of lower metabolic rates) needed to test our working 513 514 hypotheses. -Past studies have shown that total P concentrations are strongly influenced by storage 515 of P in vacuoles (Foyer & Spencer, 1986; Raghothama, 1999) and this storage pool can buffer 516 cytosolic P. - hHence measured differences in foliarleaf P may not showing a metabolically different phenotypes despite measured differences in foliar P. Similarly, the fact that investment of leaf N in 517 518 metabolism is a plastic trait – for example, increasing as a proportion of total N when N supply is 519 limiting (Cheng & Fuchigami, 2000) - means that low N treatments will not on't necessarily result in 520 lower metabolic rates. However, we found rates of A_a and R_a that were lower in low P-grown plants 521 (compared to H_NH_P), demonstrating that the supply of P was sufficiently low to create P-deficient 522 phenotypes (Table 2). Moreover, as reported previously for a range of plant species (Hammond & 523 White, 2008), starch accumulation occurred in low P-grown plants of the woody PFTs (Table 3), 524 consistent with a P₋-deficient phenotype. The low N treatment also resulted in reductions in A_a and 525 R_{a} , and reduced mass-based concentrations of soluble sugars in woody PFTs (Table 3), 526 demonstrating that N-deficient phenotypes were achieved. Importantly, the impact of the low P-

and N treatments on metabolic rates and area-basedlog-log trait-trait relationships depended on
 the extent to which <u>supply of</u> each nutrient was limiting. Our study provides the first multi PFT
 comparison assessing nutrient impacts on leaf trait relationships under controlled environment
 conditions.

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532 How does P mediate A-R-N relationships, and is the impact of P independent of N supply? 533 Past work has shown that rates of A at a given N are often lower at sites with low P availability 534 (Meir et al., 2002; Kattge et al., 2009; Reich et al., 2009; Domingues et al., 2010; Bahar et al., 2016). Based on knowledge of how P deficiencies affect leaf metabolism (Warren & Adams, 2002; Plaxton 535 536 & Tran, 2011; Bloomfield et al., 2014; Ellsworth et al., 2015) and studies comparing trait 537 relationships of plants growing on soils with contrasting P levels (Kattge et al., 2009; Reich et al., 2009), we predicted that P deficiency would reduce the proportionality-coefficient of log-log A-N, 538 539 <u>R_{Dark}-N and R_{Light}-N relationships, but with no change in the scaling-exponent of the log-log</u> relationships shallower A-N slopes in P-limited plants (Fig. 1). However, contrary to the 540 541 hypothesized relationships outlined in Fig. 1, inspection of Figure 4a shows that growth on limiting 542 P supply did not result in a *decreased* slope in the A_a-N_a relationship; rather, when plants were 543 grown on high N, low P resulted in a significant increase in scaling-exponent slope-of A_a-N_a 544 relationship, reflecting a decrease in A_a in low N_a species with low, with P deficiency having little or <u>no effect on A_a in species with high N_a . The same was true for the $R_{\text{Dark},a}$ - N_a relationship (Fig. 4d)₇</u> 545 consistent with Rowland et al. (2016) reporting higher rates of R_{Dark} for a given N in low nutrient 546 547 soils. While the reasons for this increase in scaling-exponent under low P-high N conditions remain uncertain, the results highlight how contrasting species differing in leaf structure/chemistry differ in 548 549 their responses to P deficiency Stimulation [KC3] of A_a (or R_{Dark, a}) under low P conditions (when N is 550 not limiting but not under N-limiting conditions) might reflect the fact that N supply can play a role in stimulating P uptake in low P soils (Treseder & Vitousek, 2001). 551 One of our working hypotheses was that the effect of We expected that low P supply on 552

⁵⁵²One of our working hypotheses was that the effect of <u>We expected that</u> low P supply on ⁵⁵³metabolic rates would be greatest in plants supplied with high N, based on past observations on the ⁵⁵⁴stimulatory effect of P addition being greatest when N is non-limiting (e.g. Aerts *et al.*, 1992; Meir ⁵⁵⁵*et al.*, 2001). We found<u>Our results support that L</u>ow P supply had minimal effects on metabolic ⁵⁵⁶rates when N was limiting (Table 2) - both for photosynthesis and respiration - while the inhibitory ⁵⁵⁷effect of low P was greatest when N was not limiting. These results are consistent with Liebig's law where the most limiting element has the greatest impact. A phosphorusP-supply-mediating effect
on metabolic relationships with N has been reported in several tropical studies (Reich & Walters,
1994; Denton *et al.*, 2007; Meir *et al.*, 2007; Mercado *et al.*, 2011; Quesada *et al.*, 2012), which
typically have lower P availability compared to N supply reflected in high N:P ratios.

562 Interestingly, P had no effect at all on the $\log \log A_m - N_m$ relationship irrespective of N supply 563 (Fig. S2). -The latter result contrasts with Reich *et al.* (2009), where $\log - \log A_m - N_m$ relationships 564 differed among plants growing at P-rich and P-deficient sites from field studies representing a range 565 of nutrient availability, soil properties and mycorrhizal associations, thermal environment, water 566 availability and species composition. In <u>controlled--environment conditions</u>our study, we compared 567 the effect of low P supply on a common set of species grown under common, controlled 568 environment conditions, suggesting that contrasting relationships (via different slopes-scaling-569 exponents or intercepts proportionality-coefficientselevation) from different sites and biomes may include variation other than just soil nutrient supply. Given this, care is needed when interpreting 570 the direct role of nutrient availability on A_m-N_m relationship slopes across field sites. 571

572

573 Mechanisms underpinning changes in photosynthetic N use efficiency.

Consistent with our working hypotheses (Fig. 1), we found that overall mean values of leaf-level N-574 575 use efficiency of photosynthesis (A_N) was were enhanced by low N supply, both under low and high 576 P supply (Table 2, Fig. 5a4a). Cheng & Fuchigami (2000) found that A_N of apple leaves also 577 increased with decreasing N supply. However, not all studies have reported this response, with 578 Warren et al. (2003), Hikosaka (2010) and Pons & Westbeek (2004) all reporting little effect of N supply on photosynthetic N use efficiency. Our data suggest that while low N treatment increased 579 580 $A_{\rm N}$ of some species – particularly species that exhibit inherently low N_m values (Fig. S2) – low N 581 treatment had little or no effect on other species (e.g. high N_m species). Given that species with inherently low N_m tend to also be slow-growing and/or found in less productive environments 582 583 (Wright *et al.*, 2004), this suggests that plasticity of A_N in response to variation in N supply may 584 systematically vary among species adapted to contrasting environments. The increase in A_N in low 585 N supply can occur via several possible ways (Hikosaka, 2004), such as: (1) greater fraction of leaf N 586 allocated to photosynthetic processes, particularly the relative investment of leaf N in Rubisco (Evans & Seeman, 1989; Poorter & Evans, 1998; Warren et al., 2003); (2) increase in the activation 587 588 state of Rubisco (Stitt & Schulze, 1994; Warren et al., 2000); (3) reduction of N investment in

589defence compounds, cell walls and/or nucleic acids (Chapin & Kedrowski, 1983; Onada *et al.*, 2004;590Harrison *et al.*, 2009); or (4) a decrease in inorganic N accumulation (e.g. NO_3^- in vacuoles) to591maintain organic N investment in metabolism. Similar factors are also likely to play a role in592decreased A_N exhibited by high N-grown plants with limiting P supply (Table 2) (Warren & Adams,5932002; Bloomfield *et al.*, 2014). Indeed, there was a negative relationship in all PFTs between N-use594efficiency and N:P ratio with more P-limited plants (high N:P ratio) exhibiting lower nitrogen-use595efficiencies for both A_N and $R_{Dark,N}$ (Fig. S4).

596

597 Plant functional types and bivariate relationships

598 In our study, 37 species from six PFTs were used to create a spectrum of leaf traits varying in structure, chemistry and function, which enabled us to assess whether the effect of nutrient supply 599 on leaf traits differed among the PFTs. Table 1 showed that for all leaf chemistry traits, tThe 600 601 magnitude of nutrient-mediated changes in leaf traits differed among the PFTs (Table 1). - Proteoid species exhibited steeper A_a-N_a relationships when P was low compared to broadleaved trees but 602 603 not when P was high (Fig. S5). However, the distinct PFTs exhibited relatively similar responses to 604 treatments when considering structural (i.e. M_a) and metabolic traits (particularly A_a and $R_{\text{Light},a}$). 605 Thus, despite enormous differences in leaf life span, leaf nutrients and structural components among the PFTs, generally speaking all PFTs exhibited similar metabolic responses to nutrient 606 607 availability, consistent with previous field studies (Reich et al., 2008).

608 While nNutrient supply had no effect on the ratio between leaf dark respiration and 609 photosynthesis (i.e. no significant difference in slope a common relationship across all four treatments in Fig. 5a). However, our finding of there was PFT-dependent variation in R_{Dark,a}-A_a 610 611 scaling (Fig. 85d6, Fig. S3), with PFTs differing in the proportional-coefficients of relationships that 612 shared common scaling-exponents. This contrasts with recent field-based comparisons of PFTs in 613 respiration-photosynthesis relationships (Atkin et al., 2015), where there was little evidence of C₃ herbs/grasses, broad leaved trees, shrubs and needle-leaved trees differing in rates of R_{Dark,a} at a 614 615 given A_a. However, such comparisons may be strongly influenced by a range of environmental factors that may mask underlying PFT-dependent patterns. Our study, conducted under controlled 616 617 environment conditions provided strong evidence of PFT-mediated variations in the respiration: photosynthesis ratio in plants driven by differences between woody and herbaceous species. 618 619 Future work will be needed to determine what underlying factors (e.g. respiratory costs associated

with protein turnover, phloem loading and maintenance of ion gradients in leaf cells; Lambers,
 1985; Noguchi & Yoshida, 2008) are responsible for this PFT-dependent variation in *R*_{Dark,a}-*A*_a
 scaling.

623

624 *R*Light versus *R*Dark

625 One of the objectives of our study was to assess whether nutrient supply affected the degree of 626 light inhibition of non-photorespiratory mitochondrial CO₂ release (see SI Notes S2 for further 627 details). Past work has reported that *R*_{Light} is often lower than *R*_{Dark}, with the degree of inhibition being highly variable (Griffin & Heskel, 2013) but typically around 30% (Crous et al., 2012; 628 629 Weerasinghe et al., 2014). In leaves, respiratory energy is needed for assimilation of N and protein 630 turnover, as well as for supporting the ATP demands from photosynthesis (Krömer, 1995). Given 631 that N deficiency may limit such demands for respiratory products, one possibility was that the degree of inhibition could be linked to nutrient supply, with low N potentially affecting factors such 632 as demand for TCA cycle intermediates and photorespiratory capacity. In our study, we found that 633 634 (Ayub et al., 2014) However, as shown in Fig. 96c, across a broad range of species, nutrient supply 635 had no effect on the relationship between R_{Dark} and R_{Light} (Fig. 5c), with the overall degree of light inhibition being 26% across all treatments and species. Interestingly, the degree of light inhibition 636 637 was generally less in herbaceous species compared to woody species (Fig. 9b6f), a result that merits testing in a broader set of species. 638

639

640 Conclusions

641 Our study assessed the impact of P limitations both under high and low N conditions, with the

642 results leading to the following insights: <u>Our findings show that both N- and P-supply can mediate</u>

643 leaf trait-trait relationships, suggesting that the next generation of Earth System Models should

644 <u>consider how limitations in N and P supplyavailability in individual biomes may affect trait</u>

645 <u>relationships (Goll *et al.*, 2012).</u>

- 646 <u>Linking plant metabolism to leaf nutrients, both A-N and R-N relationships were only affected by</u>
- 647 Iow P when N availability was high. Low P supply did not have an effect on the slope of A-N and R-N
- 648 relationships when N was limiting. Thus, P availability only modified the relationship between leaf R
- 649 and N when N supply was not limiting (high N). Our findings do not support the hypothesized
- 650 responses shown in Figure 1; rather, rates of leaf A and R at a given leaf N were dependent on P

651 supply in species with inherently low N_a values but not in those with inherently high N_a values. Further work assessing patterns of N and P allocation will be needed to address what factors are 652 653 responsible for these observations. Interestingly, our results also did not support Hypothesis 4, 654 with This finding supports our H.2 and is consistent with results from field-based studies (especially in tropical ecosystems with high N:P ratios). Whereas less P is needed to meet metabolic demands 655 in N-deficient plants, in high N grown plants, the effect of low P supply may be underpinned by 656 insufficient P allocated to metabolic processes or the extent to which P supply affect N allocation in 657 the plant. [KC4] 658 In A P and R P relationships, low P reduced A and R for a given P_a compared to high P, supporting 659 the first part of H.1 and H.3, but low N supply reduced A and R rates both in high and low P 660 availability. In contrast to H.4, Photosynthesis rates were reduced under low P supply in high N-661 grown plants in A N and R N relationships. By contrast, P supply had no effect at a given N_a when N 662 supply was limiting. Low N supply led to reduced A and R, both in high and low P supply. Thus, P 663 availability only modified the relationship between leaf *R* and N when N supply was not limiting 664 665 (high N). 666 Nnutrient supply had having no effect on the ratio between R_{Dark} and A_a, although woody species had a higher respiration rates for a given A compared to herbaceous species. This The fact that 667 coupling between respiration and photosynthesis remained constant suggests that deficiencies in 668 nutrient availability have similar inhibitory effects on the capacity of processes. suggests that 669 670 respiration proportionally declined with photosynthesis across species and nutrient treatments. LastlyFurther, light inhibition of respiration was constant across nutrient treatments (26% on 671 average) – again not supporting our working hypothesis - but with lower degrees of light inhibition 672 673 in herbaceous than in woody species. Both findings have relevance to ecosystem models seeking to predict future rates of carbon exchange, as the degree of light inhibition can strongly influence 674 ecosystem respiration estimates (Wohlfahrt et al., 2005; Wehr et al., 2016). Finally, the fact that 675 676 both N- and P-supply can mediate leaf trait-trait relationships suggests that the next generation of 677 Earth System Models may need to consider how limitations in N and P availability affect trait-trait 678 relationships used to predict large scale carbon fluxes. Nutrient supply had no effect on the ratio between R_{Light} and R_{Dark}, with a light inhibition across 679 680 treatments of 26%. Interestingly, our results point to a lower degree of light inhibition in 681 herbaceous than in woody species.

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691 Author contributions

- 0.K.A., K.L.G., M.H.T., P.M. and K.Y.C. designed the study. K.Y.C., O.S.S., J.Z. and A.C.A.N. collected
- 693 the gas exchange data while O.S.S. also-conducted leaf nutrient analyses. K.Y.C. led data processing,
- analyses and writing with substantial contributions from O.K.A. and K.J.B.. All authors made
- 695 conceptual contributions throughout the manuscript development.
- 696

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- 917 The following Supporting Information is available for this article:
- 918 Notes S1 Analysing trait-trait relationships
- 919 Notes S2 Light inhibition of leaf respiration to photorespiratory metabolism: Hypothesis 5

920 Supplementary Figures and Tables:

- 921 **Table S1** Species averages ± standard error of the mean per nutrient treatment for leaf mass per
- 922 area ratio (M_{a}), area based leaf nitrogen (N_{a}) and leaf phosphorus (P_{a}), light saturated
- 923 photosynthesis (A_{a}), leaf respiration in darkness ($R_{\text{Dark},a}$) and in the light ($R_{\text{Light},a}$).
- 924 **Table S2** Means ± standard errors for several leaf (nitrogen (N), phosphorus (P) and leaf mass per
- 925 area ratio, M_a) and metabolic traits (light-saturated photosynthesis (A_a), leaf respiration in darkness
- 926 (R_{Dark, a}) and in the light (R_{Light,a}) on either an area- or mass basis (indicated with a or m subscripts
- 927 respectively) for each functional group and nutrient treatment.
- Table S3 Means ± standard error of mass-based total soluble sugars, starch and TNC (total non structural carbohydrates) on 16 woody species for each nutrient treatment.
- 930 **Table S4** Standardized Major Axis regression slopes and their confidence intervals for mass-based
 931 log transformed bivariate relationships in each nutrient treatment.
- 932 Table S5 Standardized Major Axis regression slopes and their confidence intervals for area-based
 933 log transformed bivariate relationships in each PFT or between herbaceous and woody PFT groups.
- 934

Fig. S1 Box plots showing the variation of leaf structural and chemical traits for four different
nutrient treatment combination of nitrogen (N) and phosphorus (P) supply: H_NH_P (high N – high P),
H_NL_P (High N – Low P), L_NH_P (Low N – High P) and L_NL_P (Low N – Low P). Traits shown are: (a) leaf
mass per unit leaf area (M_a); (b) area-based leaf nitrogen concentration (N_a); and (c) area-based leaf
phosphorus concentration (P_a).

- 940
- Fig. S2 <u>RLog-log r</u>elationships between (a, d) mass-based light-saturated photosynthesis (A_a), (b, e)
 mass-based dark respiration (R_{Dark,a}) and (c, f) mass-based mitochondrial respiration in the light

| 1 | |
|------|--|
| 943 | (R _{Light,a}) as a function mass-based-leaf nitrogen concentrations-(N _m) (upper panels) and as a function |
| 944 | of leaf mass per unit leaf area (M_a) (lower panels) . SMA analyses were done on individual |
| 945 | observations within each nutrient combination treatment: $H_{M}H_{P}$ (high N – high P, red), $H_{M}L_{P}$ (High N |
| 946 | - Low P, green), L_NH_P (Low N - High P, blue) and L_NL_P (Low N - Low P, yellow). |
| 947 | |
| 948 | Fig. S3 Log-log relationships between area-based dark respiration (Report a) and light-saturated |
| 949 | nhotosynthesis showing individual observations within each plant functional type across all nutrient |
| 950 | treatments: C46_grass with C4 photosynthetic pathway: C26_grass species with C2 photosynthetic |
| 950 | nothway C2U C2 metabolism barby S shruby BLT broad looved trees NUT poodle looved tree. All |
| 951 | pathway; C3H, C3 metabolism nero; S, shrub; BLT, broad-leaved tree; NLT, needle-leaved tree. All |
| 952 | values are shown on a log ₁₀ scale. See Table S5 for standardized major axis (SMA) regression |
| 953 | Outputs. |
| 954 | Log-log relationships between (a) area-based dark respiration (R _{Dark,a}) and light-saturated |
| 955 | photosynthesis and (b) area-based mitochondrial respiration in the light (R _{Light,a}) showing individual |
| 956 | observations within herbaceous (herb) and woody species. See Table 5 for standardized major axis |
| 957 | (SMA) regression outputs. |
| 958 | |
| 959 | Fig. S4 Log-log relationships between (a) -N-based dark respiration (R_{Dark,N}) and (b) photosynthetic |
| 960 | nitrogen use efficiency (AN) as a function of N:P ratio for six plant functional types . |
| 961 | |
| 962 | Fig. S5 Log-log relationships of nitrogen-use efficiency (via photosynthesis-N relationships) |
| 963 | between <u>p</u> Proteoid <mark>B</mark> broadleaved species (yellow) and Broadleaved trees (BIT, blue) in each |
| 964 | nutrient treatment : H_NH_P (high N – high P, H_NL_P (High N – Low P), L_NH_P (Low N – High P) and L_NL_P |
| 965 | (Low N – Low P). Nitrogen-use efficiency was significantly (P < 0.05) higher for Proteoids when |
| 966 | grown in low P conditions (right panels) but not when P supply was high (left panels). |
| | |
| 967 | Table S1 Species averages per nutrient treatment for leaf mass per area ratio (<i>M</i> _a), area-based leaf |
| 968 | nitrogen and leaf phosphorus, light-saturated photosynthesis, leaf respiration in darkness and in |
| 969 | the light. |
| 970 | Table S2 Means for several leaf structural/chemical composition and metabolic traits on either an |
| 971 | area- or mass basis |
| P' 1 | |

- 972 **Table S3** Means of mass-based total soluble sugars, starch and total non-structural carbohydrates
- 973 <u>on 16 woody species for each nutrient treatment.</u>
- 974 **Table S4** Standardized Major Axis regression slopes and their confidence intervals for mass-based
- 975 log transformed bivariate relationships in each nutrient treatment.
- 976 **Table S5** Standardized Major Axis regression slopes and their confidence intervals for area-based
- 977 log transformed bivariate relationships in each PFT or between herbaceous and woody PFT groups.

| 979 | | | | | | | | | | | | | | | | | |
|---------------|-----|--------------------|-----|----------------|-----|--------------------|-----|----------------------|-----|---------|-----|--------------------|-----|----------------------------|-----|--------------------|--|
| | | Ma | | N _m | | P _m | | Na | | Pa | | Aa | | R _{Dark,a} | | $R_{Light,a}$ | |
| Source | DF | F | DF | F | DF | F | DF | F | DF | F | DF | F | DF | F | DF | F | |
| PFT | 5 | 204 ^{***} | 5 | 42.8*** | 5 | 43.7*** | 5 | 142*** | 5 | 38.8*** | 5 | 12.4*** | 5 | 50.7*** | 5 | 10.8*** | |
| Treatm | 3 | 1.9 ^{ns} | 3 | 83.2*** | 3 | 253 ^{***} | 3 | 78.9 ^{****} | 3 | 181*** | 3 | 7.2 ^{***} | 3 | 8.1*** | 3 | 3.5* | |
| PFT x | 15 | 1.4 ^{ns} | 15 | 2.4** | 15 | 8.3*** | 15 | 2.38 ^{**} | 15 | 6.3*** | 15 | 1.1 ^{ns} | 15 | 1.9 [*] | 15 | 0.34 ^{ns} | |
| Treatm | | | | | | | | | | | | | | | | | |
| Error | 777 | | 779 | | 779 | | 768 | | 768 | | 744 | | 735 | | 566 | | |
| <u>Within</u> | | | | | | | | | | | | | | | | | |
| Total | 800 | 45.7*** | 802 | 23.4*** | 802 | 58.4*** | 791 | 43.9*** | 791 | 42.9*** | 767 | 4.7*** | 758 | 14.1*** | 589 | 4.1*** | |

978 Table 1 Two-way ANOVA testing for differences between PFT and Nutrient treatments (Treatm).

981 Variables shown are: leaf mass per unit area (M_a), mass-based leaf nitrogen (N_m) and leaf phosphorus (P_m), area-based leaf nitrogen values (N_a)

982 and phosphorus (Pa), area-based rates of light-saturated photosynthesis (Aa) and area-based rates of leaf respiration in darkness (R_{Dark,a}) and in

983 the light (*R*_{Light,a}). Degrees of freedom (DF) and F-ratio (F) are indicated for each variable with statistical significance indicated as follows: *** *P*

984 < 0.0001, ** *P* < 0.01, * *P* < 0.05, ns is non-significant. All models were highly significant with error and total degrees of freedom varying

985 depending on the variable.

| 987 | Table 2 Means and standard error | or leaf traits and metabolic fluxes measu | red on individual plants across |
|-----|----------------------------------|---|---------------------------------|
|-----|----------------------------------|---|---------------------------------|

plantPFTs-functional groups.

| | | Treatment | | | | | | | | | |
|-----------------------------|--------------------------------------|-----------------------------|-----------------------------|-----------------------------|----------------------------|--|--|--|--|--|--|
| | | High N | High N | Low N | Low N | | | | | | |
| Trait | Units | High P | Low P | High P | Low P | | | | | | |
| N _m | mg g⁻¹ | 26.69 ± 0.85 ^a | 22.21 ± 0.64 ^b | 16.42 ± 0.65 ^c | 15.31 ± 0.55 ^c | | | | | | |
| P _m | mg g ⁻¹ | 3.88 ± 0.20 ^a | 1.07 ± 0.09 ^b | 5.52 ± 0.30 ^c | 1.69 ± 0.12 ^d | | | | | | |
| N:P | dimensionless | 8.57 ± 0.29 ^a | 34.56 ± 1.55 ^b | 4.72 ± 0.26 ^c | 15.00 ± 0.87 ^d | | | | | | |
| MA | g m ⁻² | 113.18 ± 7.22 ^a | 118.82 ± 5.77 ^a | 109.75 ± 5.71 ^a | 119.96 ± 6.16 ^a | | | | | | |
| Aa | µmol m⁻² s⁻¹ | 15.59 ± 0.48 ^a | 13.01 ± 0.48 ^b | 12.17 ± 0.43 ^b | 12.13 ± 0.41 ^b | | | | | | |
| A _m | nmol g ⁻¹ s ⁻¹ | 245.42 ± 17.77 ^a | 168.02 ± 12.56 ^b | 162.00 ± 9.83 ^b | 149.59 ± 8.15 ^b | | | | | | |
| A _N | µmol <u>g</u> _N ₦ s⁻¹ | 8.65 ± 0.41 ^a | 6.92 ± 0.39 ^b | 9.56 ± 0.36 ^c | 9.55 ± 0.36 ^c | | | | | | |
| A _P | µmol g <mark>₽</mark> ₽ s⁻¹ | 72.99 ± 3.80 ^a | 195.35 ± 8.98 ^b | 43.84 ± 2.72 ^c | 131.42 ± 6.90 ^d | | | | | | |
| R _{Dark,a} | µmol m ⁻² s ⁻¹ | 1.07 ± 0.05 ^a | 0.90 ± 0.04 ^b | 0.78 ± 0.04 ^{bc} | 0.74 ± 0.03 ^c | | | | | | |
| $R_{Dark,m}$ | nmol g ⁻¹ s ⁻¹ | 14.67 ± 0.87 ^a | 10.68 ± 0.66 ^b | 9.94 ± 0.61^{b} | 8.79 ± 0.51 ^c | | | | | | |
| $R_{Dark,N}$ | µmol <u>g</u> _N ₦ s⁻¹ | 0.56 ± 0.03 ^b | 0.47 ± 0.03 ^a | 0.60 ± 0.03^{c} | 0.57 ± 0.03 ^{bc} | | | | | | |
| $R_{Dark,P}$ | µmol <u>g</u> ₽₽ s⁻¹ | 4.87 ± 0.30 ^a | 14.24 ± 0.87 ^b | 2.50 ± 0.15 ^c | 7.62 ± 0.45 ^d | | | | | | |
| $R_{ m Light, a}$ | µmol m ⁻² s ⁻¹ | 0.86 ± 0.05 ^a | 0.76 ± 0.05 ^{ab} | 0.67 ± 0.05 ^{bc} | 0.61 ± 0.05 ^c | | | | | | |
| $R_{Light,m}$ | nmol g ⁻¹ s ⁻¹ | 11.83 ± 1.12 ^a | 9.41 ± 1.06 ^b | 9.18 ± 1.10^{b} | 8.31 ± 1.14 ^b | | | | | | |
| $R_{Light, N}$ | µmol <u>g</u> _N ₦ s⁻¹ | 0.43 ± 0.03 ^a | 0.39 ± 0.03 ^a | 0.52 ± 0.06 ^a | 0.49 ± 0.06 ^a | | | | | | |
| R Light, P | µmol <u>g</u> ₽ s⁻¹ | 3.77 ± 0.30 ^c | 11.31 ± 0.91 ^a | 2.01 ± 0.05 ^d | 6.10 ± 0.53^{b} | | | | | | |
| R _{Dark} :A | dimensionless | 0.076 ± 0.004 ^{ab} | 0.086 ± 0.007 ^a | 0.070 ± 0.004 ^{ab} | 0.064 ± 0.002 ^b | | | | | | |
| R _{Light} :A | | 0.056 ± 0.004 ^{ab} | 0.061 ± 0.006 ^a | 0.052 ± 0.005 ^{bc} | 0.046 ± 0.004 ^c | | | | | | |
| $R_{Light}:R_{Dark}$ | | 0.701 ± 0.03 ^a | 0.787 ± 0.06 ^a | 0.741 ± 0.04 ^a | 0.731 ± 0.05 ^a | | | | | | |

The sample size for each leaf trait varied between 217-220 (171-183 for R_{Light} variables). Leaf traits presented here are mass-based nitrogen (N_m) and phosphorus (P_m) concentrations, their ratio (N:P) and leaf mass per area ratio (M_A) Metabolic traits are light saturated photosynthesis (A) and dark respiration (R_{Dark}), presented on an area-, mass, Nand P-basis as well as the R/A ratio. Lowercase letters indicated significant differences between different nutrient treatments at P < 0.05 in a least square means Student-t post-hoc test.

995 Table 3 Means and standard error of the mean across Plant Functional Type (PFT) for soluble sugars,

996 starch and non-structural carbohydrates (TNC) both on a mass-basis (upper panel) and an area-basis

997 (bottom panel), including sample size (n).

| Treatment | Units | H _N H _P | H _N L _P | L _N H _P | L _N L _P |
|-----------------------------------|--------------------|-------------------------------|-------------------------------|-------------------------------|-------------------------------|
| n | | 217 | 218 | 218 | 220 |
| MASS-BASED | | | | | |
| Soluble Sugars | mg g ⁻¹ | 51.85 ± 3.35 ^a | 43.13 ± 3.62 ^{ab} | 39.34 ± 2.76 ^b | 39.81 ± 2.68 ^b |
| Starch | mg g⁻¹ | 35.20 ± 4.57 ^b | 56.74 ± 7.87 ^a | 41.99 ± 7.27 ^b | 62.41 ± 10.21 ^{ab} |
| TNC | mg g⁻¹ | 87.05 ± 6.53 ^a | 99.88 ± 8.84 ^a | 81.32 ± 8.21 ^a | 102.22 ± 10.44 ^a |
| AREA-BASED | | | | | |
| Soluble Sugars | g m ⁻² | 7.44 ± 1.07 ^a | 6.20 ± 0.68 ^a | 5.29 ±0.56 ^a | 6.20 ± 0.57 ^a |
| Starch | g m ⁻² | 3.80 ± 0.44^{b} | 6.89 ± 1.08 ^a | 5.12 ± 0.99 ^b | 8.52 ± 1.46 ^a |
| TNC | g m ⁻² | 11.24 ± 1.32 ^{bc} | 13.09 ± 1.26 ^{ab} | 10.41 ± 1.19 ^c | 14.72 ± 1.53 ^a |
| <i>M</i> a (TNC _{corr}) | g m⁻² | 137.7 ± 16.2 ^a | 130.6 ± 7.8 ^a | 128.0 ± 11.4^{a} | 143.8 ± 11.2 ^a |

998

999 Different letters indicate statistical differences between treatments at P < 0.05 in a Least square means 900 Student-t post-hoc test after log transformation. Also shown are M_a corrected for area-based TNC (TNC_{corr}). 901 Nutrient treatments are defined as follows: High N–High P (H_NH_P), High N–Low P (H_NL_P), Low N–High P 902 (L_NH_P) and Low N–Low P (L_NL_P).

Table 4 Standardized Major Axis coefficients for area-based log-transformed bivariate relationships for Figs 3, 4 and 5.

| | | | | | | | Starting model | | | | Simpler | nodel, comm | on slope | es | |
|------|--|-------------------------------|-----|-------|-----------------|--------------------------------|--------------------------------|--------------|----------|-------------------------|--------------------------------|-------------------|----------|-------------------------|---------------|
| | | | | | | | | H0: no diff. | | | | HO: no diff. | | | H0: no diff. |
| | | | | - 2 | | Scaling exponent (β_1) | Proportionality | in β_1 | Pairwise | | Proportionality | in β _o | Pairwise | 2 | in axis shift |
| Fig. | Model | Treat. | n | Rź | <i>p</i> -value | (±CIs) | coefficient (β_0) (±Cls) | (p-value) | comp. | Common β_1 (±CIs) | coefficient (β_0) (±Cls) | (p-value) | comp. | Common β_0 (±Cls) | (p-value) |
| 3a | N _a - <i>M</i> _a | H_NH_P | 182 | 0.675 | < 0.0001 | 0.901 (0.829, 0.98) | -1.424 (-1.571, -1.277) | 0.4955 | | 0.854 (0.814, 0.897) | -1.334 (-1.417, -1.251) | < 0.0001 | А | | < 0.0001 |
| | | H_NL_P | 177 | 0.603 | < 0.0001 | 0.838 (0.763, 0.921) | -1.356 (-1.514, -1.198) | | | | -1.388 (-1.473, -1.302) | | В | | |
| | | L_NH_P | 186 | 0.443 | < 0.0001 | 0.835 (0.749, 0.93) | -1.518 (-1.697, -1.339) | | | | -1.557 (-1.642, -1.471) | | С | | |
| | | L _N L _P | 184 | 0.466 | < 0.0001 | 0.822 (0.739, 0.915) | -1.513 (-1.689, -1.337) | | | | -1.577 (-1.663, -1.49) | | С | | |
| 3b | P _a - <i>M</i> _a | $H_{\rm N}H_{\rm P}$ | 182 | 0.475 | < 0.0001 | 1.053 (0.947, 1.171) | -2.608 (-2.827, -2.39) | < 0.0001 | А | | | | | | |
| | | H_NL_P | 177 | 0.025 | 0.0361 | 0.95 (0.821, 1.101) | -3.034 (-3.316, -2.752) | | A | | | | | | |
| | | L_NH_P | 186 | 0.369 | < 0.0001 | 1.463 (1.304, 1.642) | -3.288 (-3.623, -2.953) | | В | | | | | | |
| | | L _N L _P | 184 | 0.147 | < 0.0001 | 1.143 (0.999, 1.308) | -3.21 (-3.52, -2.899) | | А | | | | | | |
| 3c | N _a - P _a | $H_{\rm N}H_{\rm P}$ | 182 | 0.624 | < 0.0001 | 0.856 (0.782, 0.936) | 0.808 (0.756, 0.861) | < 0.0001 | А | | | | | | |
| | | H_NL_P | 177 | 0.057 | 0.0014 | 0.882 (0.764, 1.019) | 1.321 (1.167, 1.474) | | А | | | | | | |
| | | $L_{\rm N}H_{\rm P}$ | 186 | 0.265 | < 0.0001 | 0.57 (0.504, 0.646) | 0.358 (0.312, 0.404) | | В | | | | | | |
| | | L _N L _P | 184 | 0.119 | < 0.0001 | 0.719 (0.627, 0.825) | 0.795 (0.693, 0.898) | | А | | | | | | |
| 4a | A _a - N _a | $H_{\rm N}H_{\rm P}$ | 182 | 0.094 | < 0.0001 | 0.747 (0.649, 0.858) | 0.926 (0.878, 0.975) | 0.0008 | С | | | | | | |
| | | $H_{N}L_{P}$ | 177 | 0.070 | 0.0004 | 1.135 (0.983, 1.31) | 0.702 (0.633, 0.77) | | А | | | | | | |
| | | $L_{\rm N}H_{\rm P}$ | 186 | 0.250 | < 0.0001 | 0.921 (0.813, 1.045) | 0.929 (0.894, 0.963) | | В | | | | | | |
| | | L _N L _P | 184 | 0.211 | < 0.0001 | 0.895 (0.786, 1.019) | 0.935 (0.899, 0.97) | | B,C | | | | | | |
| 4b | A _a - P _a | $H_{\rm N}H_{\rm P}$ | 182 | 0.026 | 0.0306 | 0.639 (0.553, 0.738) | 1.53 (1.462, 1.598) | < 0.0001 | В | | | | | | |
| | | $H_{N}L_{P}$ | 177 | 0.049 | 0.0030 | 1.001 (0.866, 1.157) | 2.2 (2.026, 2.375) | | А | | | | | | |
| | | $L_{N}H_{P}$ | 186 | 0.057 | 0.0010 | 0.526 (0.457, 0.605) | 1.258 (1.207, 1.309) | | В | | | | | | |
| | | L _N L _P | 184 | 0.017 | 0.0745 | | | | | | | | | | |
| 4c | A _a - M _a | $H_{\rm N}H_{\rm P}$ | 183 | 0.001 | 0.6419 | | | | | | | | | | |
| | | $H_{N}L_{P}$ | 177 | 0.003 | 0.4969 | | | | | | | | | | |
| | | $L_{N}H_{P}$ | 189 | 0.023 | 0.0379 | 0.77 (0.668, 0.887) | -0.469 (-0.687, -0.251) | 0.9540 | | 0.767 (0.694, 0.848) | -0.464 (-0.621, -0.308) | 0.4150 | | -0.475 (-0.629, -0.321) | 0.5442 |
| | | L _N L _P | 190 | 0.020 | 0.0505 | 0.765 (0.664, 0.882) | -0.485 (-0.705, -0.264) | | | | -0.489 (-0.648, -0.33) | | | | |
| 4d | R _{dark,a} - N _a | $H_{\rm N}H_{\rm P}$ | 182 | 0.140 | < 0.0001 | 1.03 (0.899, 1.18) | -0.363 (-0.427, -0.299) | 0.2024 | | 1.142 (1.068, 1.221) | -0.397 (-0.453, -0.341) | < 0.0001 | В | | < 0.0001 |
| | | H_NL_P | 177 | 0.080 | 0.0001 | 1.271 (1.102, 1.466) | -0.508 (-0.585, -0.432) | | | | -0.469 (-0.524, -0.414) | | С | | |
| | | L_NH_P | 186 | 0.223 | < 0.0001 | 1.121 (0.987, 1.274) | -0.296 (-0.34, -0.253) | | | | -0.299 (-0.34, -0.257) | | А | | |
| | | L _N L _P | 184 | 0.188 | < 0.0001 | 1.171 (1.027, 1.336) | -0.347 (-0.394, -0.299) | | | | -0.343 (-0.388, -0.299) | | A,B | | |
| 4e | R _{dark,a} - P _a | $H_N H_P$ | 182 | 0.049 | 0.0027 | 0.881 (0.764, 1.017) | 0.47 (0.378, 0.561) | < 0.0001 | В | | | | | | |
| | | H _N L _P | 177 | 0.035 | 0.0127 | 1.121 (0.969, 1.298) | 1.17 (0.973, 1.367) | | А | | | | | | |
| | | L _N H _P | 186 | 0.208 | < 0.0001 | 0.64 (0.562, 0.728) | 0.105 (0.051, 0.159) | | С | | | | | | |
| | | L _N L _P | 184 | 0.054 | 0.0015 | 0.842 (0.731, 0.971) | 0.585 (0.46, 0.71) | | В | | | | | | |
| 4f | R _{dark,a} - M _a | H _N H _P | 183 | 0.074 | 0.0002 | 0.929 (0.807, 1.069) | -1.834 (-2.091, -1.577) | 0.5230 | | 0.969 (0.903, 1.04) | -1.912 (-2.053, -1.77) | < 0.0001 | А | | 0.0563 |
| | | H _N L _P | 177 | 0.045 | 0.0047 | 1.066 (0.922, 1.232) | -2.232 (-2.544, -1.919) | | | | -2.041 (-2.186, -1.895) | | В | | |
| | | L _N H _P | 189 | 0.081 | < 0.0001 | 0.938 (0.817, 1.076) | -2.002 (-2.259, -1.744) | | | | -2.063 (-2.205, -1.922) | | В | | |
| | | LNLD | 190 | 0.047 | 0.0026 | 0.958 (0.833. 1.102) | -2.116 (-2.3881.844) | | | | -2.137 (-2.2831.992) | | С | | |

006 Table 4 contd.

| International difference of the conditional difference of the condit difference of the conditional difference of the con | | | | | | | Starting model | | | Simpler model, common slopes | | | | | | | |
|--|------|---|-------------------------------|-----|-------|-----------------|------------------------------------|----------------------------------|--------------|------------------------------|-------------------------|----------------------------------|--------------|----------|-------------------------|---------------|--|
| Staling requences ([b] Proportionality in Proportionality | | | | | | | | | H0: no diff. | | | | H0: no diff. | | | H0: no diff. | |
| $ \begin{array}{c c c c c c c c c c c c c c c c c c c $ | | | | | 2 | | Scaling exponent (β ₁) | Proportionality | in β_1 | Pairwise | | Proportionality | in β_0 | Pairwise | | in axis shift | |
| | Fig. | Model | Treat. | n | R² | <i>p</i> -value | (±CIs) | coefficient (β_0) (±Cls) | (p-value) | comp. | Common β_1 (±Cls) | coefficient (β_0) (±Cls) | (p-value) | comp. | Common β_0 (±CIs) | (p-value) | |
| $ \begin{array}{ c c c c c c c c c c c c c c c c c c c$ | 4g | <i>R_{light,a}</i> - N _a | $H_{\rm N}H_{\rm P}$ | 152 | 0.082 | 0.0004 | 1.299 (1.114, 1.515) | -0.629 (-0.726, -0.531) | 0.0471 | А | | | | | | | |
| $ \begin{array}{ c c c c c c } & c c c c c c c c c c c c c c c c c c $ | | | H_NL_P | 138 | 0.041 | 0.0176 | 1.572 (1.332, 1.854) | -0.757 (-0.876, -0.639) | | A,B | | | | | | | |
| $ \begin{array}{ c c c c c c c c c c c c c c c c c c c$ | | | $L_{\rm N}H_{\rm P}$ | 148 | 0.044 | 0.0107 | 1.738 (1.482, 2.038) | -0.586 (-0.68, -0.492) | | В | | | | | | | |
| $ \begin{array}{c c c c c c c c c c c c c c c c c c c $ | | | L _N L _P | 134 | 0.040 | 0.0209 | 1.679 (1.42, 1.986) | -0.588 (-0.685, -0.492) | | В | | | | | | | |
| $ \begin{array}{ c c c c c c c c c c c c c c c c c c c$ | 4h | <i>R_{light,a}</i> - P _a | $H_{\rm N}H_{\rm P}$ | 152 | 0.039 | 0.0150 | 1.082 (0.925, 1.267) | 0.425 (0.3, 0.55) | 0.0016 | B,C | | | | | | | |
| $ \begin{array}{c c c c c c c c c c c c c c c c c c c $ | | | $H_{\rm N}L_{\rm P}$ | 138 | 0.053 | 0.0068 | 1.455 (1.235, 1.715) | 1.399 (1.115, 1.683) | | А | | | | | | | |
| $ \begin{array}{c c c c c c c c c c c c c c c c c c c $ | | | $L_{\rm N}H_{\rm P}$ | 148 | 0.109 | < 0.0001 | 0.969 (0.831, 1.131) | 0.06 (-0.04, 0.16) | | С | | | | | | | |
| | | | L _N L _P | 134 | 0.032 | 0.0397 | 1.317 (1.113, 1.559) | 0.891 (0.658, 1.125) | | A,B | | | | | | | |
| $ \begin{array}{c c c c c c c c c c c c c c c c c c c $ | 4i | R _{light,a} - M _a | $H_{\rm N}H_{\rm P}$ | 152 | 0.024 | 0.0573 | | | | | 1.35 (1.244, 1.464) | | | | -2.938 (-3.158, -2.718) | | |
| $ \begin{array}{c c c c c c c c c c c c c c c c c c c $ | | | H_NL_P | 138 | 0.019 | 0.1077 | | | | | | | | | | | |
| $ \begin{array}{ c c c c } \hline \ \ \ \ \ \ \ \ \ \ \ \ \$ | | | L_NH_P | 151 | 0.003 | 0.4968 | | | | | | | | | | | |
| 5a R _{daba} A _a HNHP 187 0.128 < 0.0001 | | | L _N L _P | 138 | 0.000 | 0.8346 | | | | | | | | | | | |
| $ \begin{array}{c c c c c c c c c c c c c c c c c c c $ | 5a | R _{dark,a} - A _a | HNHP | 187 | 0.128 | < 0.0001 | 1.373 (1.2, 1.572) | -1.635 (-1.854, -1.416) | 0.1637 | | 1.234 (1.163, 1.311) | -1.476 (-1.573, -1.379) | 0.0600 | | -1.45 (-1.532, -1.368) | < 0.0001 | |
| $ \begin{array}{c c c c c c c c c c c c c c c c c c c $ | | | HNLP | 187 | 0.191 | < 0.0001 | 1.109 (0.974, 1.263) | -1.284 (-1.441, -1.126) | | | | -1.415 (-1.506, -1.324) | | | | | |
| $ \begin{array}{c c c c c c c c c c c c c c c c c c c $ | | | LNHP | 191 | 0.387 | < 0.0001 | 1.222 (1.093, 1.368) | -1.435 (-1.581, -1.29) | | | | -1.448 (-1.532, -1.364) | | | | | |
| 5b R _{light,a} - A _a HNP 154 0.204 < 0.001 1.75 (1.517, 2.017) -2.28 (-2.524, -1.931) 0.0035 B,C INP 145 0.094 0.002 1.504 (1.286, 1.759) -1.89 (-2.157, -1.623) C INP 151 0.190 < 0.0001 | | | LNLP | 190 | 0.428 | < 0.0001 | 1.253 (1.124, 1.397) | -1.509 (-1.654, -1.364) | | | | -1.49 (-1.574, -1.406) | | | | | |
| HNLP 145 0.094 0.0002 1.504 (1.286, 1.759) -1.89 (-2.157, -1.623) C LNHP 151 0.190 <0.0001 | 5b | R _{light,a} - A _a | HNHP | 154 | 0.204 | < 0.0001 | 1.75 (1.517, 2.017) | -2.228 (-2.524, -1.931) | 0.0035 | B,C | | | | | | | |
| $ \begin{array}{c c c c c c c c c c c c c c c c c c c $ | | | HNLP | 145 | 0.094 | 0.0002 | 1.504 (1.286, 1.759) | -1.89 (-2.157, -1.623) | | С | | | | | | | |
| $ \begin{array}{c c c c c c c c c c c c c c c c c c c $ | | | LNHP | 151 | 0.190 | < 0.0001 | 2.186 (1.89, 2.527) | -2.712 (-3.064, -2.359) | | А | | | | | | | |
| $ \begin{array}{c c c c c c c c c c c c c c c c c c c $ | | | LNLP | 138 | 0.191 | < 0.0001 | 2.018 (1.734, 2.35) | -2.553 (-2.892, -2.214) | | A,B | | | | | | | |
| $ \begin{array}{c c c c c c c c c c c c c c c c c c c $ | 5c | R _{light,a} - R _{dark,a} | HNHP | 154 | 0.525 | < 0.0001 | 1.615 (1.447, 1.804) | -0.239 (-0.284, -0.195) | 0.4906 | | 1.613 (1.518, 1.714) | -0.239 (-0.284, -0.195) | 0.0948 | | -0.188 (-0.216, -0.161) | < 0.0001 | |
| $ \begin{array}{c c c c c c c c c c c c c c c c c c c $ | | | HNLP | 145 | 0.402 | < 0.0001 | 1.536 (1.352, 1.744) | -0.181 (-0.236, -0.125) | | | | -0.177 (-0.233, -0.121) | | | | | |
| $ \begin{array}{c c c c c c c c c c c c c c c c c c c $ | | | LNHP | 151 | 0.425 | < 0.0001 | 1.742 (1.541, 1.969) | -0.155 (-0.217, -0.093) | | | | -0.169 (-0.225, -0.113) | | | | | |
| 5d R _{dark,a} - A _a Herb 361 0.165 < 0.001 | | | LNLP | 138 | 0.412 | < 0.0001 | 1.554 (1.365, 1.769) | -0.165 (-0.231, -0.099) | | | | -0.157 (-0.219, -0.094) | | | | | |
| $ \begin{array}{c c c c c c c c c c c c c c c c c c c $ | 5d | R _{dark,a} - A _a | Herb | 361 | 0.165 | < 0.0001 | 1.122 (1.021, 1.234) | -1.408 (-1.521, -1.295) | 0.4462 | | 1.864 (1.663, 2.089) | -2.359 (-2.592, -2.127) | < 0.0001 | | | < 0.0001 | |
| $ \begin{array}{c c c c c c c c c c c c c c c c c c c $ | | | Woody | 394 | 0.354 | < 0.0001 | 1.177 (1.087, 1.275) | -1.334 (-1.44, -1.227) | | | | -2.253 (-2.459, -2.047) | | | | | |
| Woody - - 1.154 (1.086, 1.227) -1.308 (-1.39, -1.227) 5e R _{light,a} - A _a Herb 267 0.104 <0.001 | | | Herb | - | - | - | 1.154 (1.086, 1.227) | -1.441 (-1.519, -1.363) | | | | | | | | | |
| Se R light,a - A a Herb 267 0.104 < 0.0001 1.864 (1.663, 2.089) -2.359 (-2.592, -2.127) 0.5959 1.9 (1.715, 2.105) -0.059 (-0.119, 0.001) 0.5008 -2.318 (-2.472, -2.165) < 0.0001 Woody 321 0.198 < 0.0001 | | | Woody | - | - | - | 1.154 (1.086, 1.227) | -1.308 (-1.39, -1.227) | | | | | | | | | |
| Woody 321 0.198 < 0.001 1.789 (1.621, 1.975) -2.253 (-2.459, -2.047) -0.248 (-0.277, -0.219) 5f R light, a - R dark, a Herb 267 0.281 < 0.0001 | 5e | R _{light,a} - A _a | Herb | 267 | 0.104 | < 0.0001 | 1.864 (1.663, 2.089) | -2.359 (-2.592, -2.127) | 0.5959 | | 1.9 (1.715, 2.105) | -0.059 (-0.119, 0.001) | 0.5008 | | -2.318 (-2.472, -2.165) | < 0.0001 | |
| 5f R light,a - R dark,a Herb 267 0.281 < 0.0001 1.9 (1.715, 2.105) -0.059 (-0.119, 0.001) 0.0005 Woody 321 0.576 < 0.0001 | | - | Woody | 321 | 0.198 | < 0.0001 | 1.789 (1.621, 1.975) | -2.253 (-2.459, -2.047) | | | | -0.248 (-0.277, -0.219) | | | | | |
| Woody 321 0.576 < 0.0001 1.521 (1.416, 1.634) -0.248 (-0.277, -0.219) | 5f | R _{light,a} - R _{dark,a} | Herb | 267 | 0.281 | < 0.0001 | 1.9 (1.715, 2.105) | -0.059 (-0.119, 0.001) | 0.0005 | | | | | | | | |
| | | | Woody | 321 | 0.576 | < 0.0001 | 1.521 (1.416, 1.634) | -0.248 (-0.277, -0.219) | | | | | | | | | |

Coefficients of determination (R²) and significance values (P-value) of each log-log trait-trait relationship are shown in addition to 95% confidence intervals (CI) of SMA slopes (scaling-exponent) and elevation

(proportionality-coefficient). When scaling-exponents were significantly different between nutrient treatments (P < 0.05) in the starting model, a post hoc test showed pairwise comparisons between

treatments, indicated with capital letters. When scaling-exponents were not significantly different, SMA tests for proportionality-coefficient differences were conducted using a common scaling-exponent

(simpler model). Where there was a significant difference in proportionality-coefficient of the common scaling-exponent SMA regressions, values for the proportionality-coefficient are pairwise comparisons

were indicated via different letters in post-hoc testing. When elevation using a common scaling-exponent showed no significant differences, a test investigating potential shifts along a relationship was

018 <u>conducted (last column) indicated via a P-value.</u>

1014 Figure Legends[0A5][0A6]

- 1015 **Fig. 1** Theoretical <u>log-log</u> relationships linking metabolic rates to leaf nitrogen <u>concentration</u>
- 1016 <u>tent</u> among plant functional types. Changes associated with Hypothesis <u>2 and 3</u>2 are illustrated and
- 1017 show expected changes in slopes (scaling exponent) and/or y axis intercepts elevation
- 1018 (proportionality coefficient proportionality-coefficient) of log-log relationships. Ellipses represent
- 1019 anticipated metabolic rate-N concentration combinations in each of the four treatments (high N-
- 1020 high P, high N-low P, low N-high P and low N-low P) used in our study, with their standardized
- 1021 major axes (SMAs) shown. Note that as a consequence of Hypothesis 1, we anticipate low P supply
- 1022 to reduce rates of metabolism when N is abundant, but with less sonot when N is limiting. When N
- 1023 is limiting, <u>average</u> rates of metabolism are predicted to be <u>slightly lower similar</u> in <u>high and the low</u>
- 1024 P-grown plants, resulting in <u>but with a shallower slope</u>lower scaling proportionality-coefficient
- 1025 <u>exponent in the low P-grown plants of the SMA regressions compared to high-P grown plants</u>,
- 1026 Irrespective of P supply, we anticipate low N supply to result in an increase in the y-axis
- 1027 intercepts<u>elevation</u> of log log relationships linkinglowerhigher metabolic rates to for a given level of
- 1028 leaf N, reflecting increased relative allocation of leaf N to photosynthesis when N is limiting.
- 1029

1030 Fig. 2 Box plots showing the variation of leaf structural and chemical traits and leaf metabolism of carbon fluxes for six different plant functional type (PFT) classifications across all nutrient 1031 1032 treatments. Traits shown are: (a) mass-based leaf nitrogen concentration (Nm); (b) mass-based leaf 1033 <u>phosphorus concentration (P_m); (c)</u> leaf mass per unit leaf area (M_a); (b) area-based leaf nitrogen 1034 concentration (N_a); and (c) area based leaf phosphorus concentration (P_a); (d) area-based lightsaturated photosynthesis (A_a) ; (e) area-based leaf mitochondrial respiration in darkness $(R_{Dark,a})$; 1035 1036 and (f) area-based leaf mitochondrial respiration in the absence of photorespiration (R_{Light,a}).- Data 1037 shown are for individual observations to give an indication of underlying data distribution. The boxes indicate the interquartile range and median. Observations outside the whiskers are shown as 1038 1039 individual points. PFT classifications: C4G, grass species with C4 photosynthetic pathway; C3G, grass 1040 species with C3 photosynthetic pathway; C3H, C3 metabolism herb; S, shrub; BITBLT, broad-1041 leaved tree; NITNLT, needle-leaved tree. Post-hoc differences among PFT are shown in Table S2 1042 (across treatments column).

Fig. 3 Box plots showing the variation of leaf metabolism of carbon fluxes for six different plant
 functional type (PFT) classifications. Traits shown are: (a) area-based light saturated photosynthesis
 (Aa); (b) area-based leaf mitochondrial respiration in darkness (*R*_{Dark,a}); and (c) area-based leaf
 mitochondrial respiration in the absence of photorespiration (*R*_{Light,a}). Data shown are for individual
 observations to give an indication of underlying data distribution. PFT classifications: C4G, grass
 with C4-photosynthetic pathway; C3G, grass species with C3-photosynthetic pathway; C3H, C3
 metabolism herb; S, shrub; BIT<u>BLT</u>, broad-leaved tree; NIT<u>NLT</u>, needle-leaved tree.

1051

Fig. 4–3_Relationships between area-based leaf nitrogen levels (N_a) and area-based leaf
phosphorus concentration (P_a) concentration as a function of area-based leaf mass per unit leaf
area (M_a) (panels (a) and (b) respectively) and the relationship between N_a versus P_a (panel (c)). All
values are shown on a log₁₀ scale. SMA analyses were done on individual observations within each
nutrient combination treatment: H_NH_P (high N – high P, redblack), H_NL_P (High N – Low P, green),
L_NH_P (Low N – High P, blue) and L_NL_P (Low N – Low P, vellow orange). See Table 4 for standardized
major axis (SMA) regression outputs.

1059

1060 Fig. 5-4 Relationships between: (1) area-based rates of leaf CO₂ exchange light-saturated 1061 photosynthesis (A_a) as a function of (a, d and g) area-based leaf nitrogen concentration (N_a), (b, e, and h) area-based leaf phosphorus concentration (P_a), -and-(c, f and i) leaf mass per unit leaf area 1062 1063 (M_a) . Upper, middle and lower panels show light-saturated photosynthesis (A_a) , area-based dark respiration (R_{Dark,a}) and mitochondrial respiration in the light (R_{Light,a}), respectively. – All values are 1064 1065 shown on a log₁₀ scale. SMA analyses were done on individual observations within each nutrient 1066 combination treatment: H_NH_P (high N – high P, redblack), H_NL_P (High N – Low P, green), L_NH_P (Low N 1067 – High P, blue) and L_NL_P (Low N – Low P, yelloworange). See Table 4 for standardized major axis 1068 (SMA) regression outputs.

1069

1D70Fig. 6 - 5 Relationships between area-based dark respiration ($R_{\text{Dark},a}$) (upper panels) as a function of1D71(a) area-based leaf nitrogen concentration (N_a), (b) area-based leaf phosphorus concentration (P_a)1D72and (c) leaf mass per unit leaf area (M_a), while lower panels show mitochondrial respiration in the1D73light ($R_{\text{Light},a}$) as a function of (d) area-based leaf nitrogen concentration (N_a), (e) area-based leaf1D74phosphorus concentration (Pa) and (f) leaf mass per unit leaf area (M_a). All values are shown on a

1075 log₁₀ scale. SMA analyses were done on individual observations within each nutrient combination treatment: H_NH_P (high N – high P, redblack), H_NL_P (High N – Low P, green), L_NH_P (Low N – High P, 1076 1077 blue) and L_NL_P (Low N – Low P, yelloworange). See Table 4 for standardized major axis (SMA) regression outputs. Regression lines are not shown when relationships were not significant. 1078 1079 Fig. 7-65 Log-log trait-trait rRelationships on a log₁₀ scale among nutrient treatments (upper 1080 1081 panels) between: (a) area-based dark respiration ($R_{\text{Dark},a}$) and light-saturated photosynthesis (A_a) 1082 across nutrient treatments, with the single regression line showing the overall relationship, given 1083 the lack of slope differences among nutrient treatments (Table 4)[OA7]; and (b) area-based mitochondrial respiration in the light (R_{Light,a}) and A_a; and (c) area-based R_{Light} and area-based R_{Dark} 1084 1085 showing SMA regression lines for each treatment.- Values shown are for individual observations within each nutrient combination treatment: H_NH_P (high N – high P, black), H_NL_P (High N – Low P, 1086 green), L_NH_P (Low N – High P, blue) and L_NL_P (Low N – Low P, orange). See Table 4 for standardized 1087 major axis (SMA) regression outputs. All values are shown on a log₁₀ scale. The bottom panels (d, e, 1088 1089 f) reflect the same relationships for individual observations within herbaceous (herb, blue) and woody species (orange) with SMA outputs in Table S5. In (a) and (b), values shown are for individual 1090 observations within each nutrient combination treatment: H_NH_P (high N – high P, red), H_NL_P (High N 1091 - Low P, green), L_NH_P (Low N - High P, blue) and L_NL_P (Low N - Low P, yellow). See Table 4 for 1092 standardized major axis (SMA) regression outputs. 1093 1094 1095 Fig. 8 Relationships between (a) area-based dark respiration (R_{Darka}) and light-saturated 1096 photosynthesis showing individual observations within each plant functional type across all nutrient 1097 treatments: C4G, grass with C4 photosynthetic pathway; C3G, grass species with C3 photosynthetic pathway; C3H, C3 metabolism herb; S, shrub; BITBLT, broad-leaved tree; NITNLT, needle-leaved 1098 tree. All values are shown on a log₁₀ scale. See Table 5 for standardized major axis (SMA) regression 1099 1100 outputs. 1101 1102 Fig. 9- Relationships between area based mitochondrial respiration in the light (R_{Light,a}) and areabased dark respiration (R_{Dark,a}) showing individual observations within (a) each nutrient treatment 1103

1104 being H_NH_P (high N – high P, red), H_NL_P (High N – Low P, green), L_NH_P (Low N – High P, blue) and L_NL_P

1105 (Low N – Low P, yellow) and (b) within woody and herbaceous plant species. All values are shown

| 1106 | on a log ₁₀ scale. | See Table 4 | for standardiz | ed major axis | (SMA) re | gression οι | itputs for | each nutrient |
|------|-------------------------------|-------------|----------------|---------------|----------|------------------------|-----------------------|---------------|

- 1107 treatment and Table 5 for standardized major axis (SMA) regression outputs for woody versus
- 1108 herbaceous species
- 109







117 Figure 3



Figure 4



