



ECOLOGY

## Plant responses to fertilization experiments in lowland, species-rich, tropical forests

Journal:	<i>Ecology</i>
Manuscript ID	ECY17-1116.R1
Wiley - Manuscript type:	Articles
Date Submitted by the Author:	n/a
Complete List of Authors:	Wright, S. Joseph; Smithsonian Tropical Research Institute, ; Smithsonian Tropical Research Institute, Turner, Benjamin; Smithsonian, Yavitt, Joseph; Cornell University, Natural Resources Harms, Kyle; Louisiana State University, Biological Sciences Kaspari, Michael; University of Oklahoma, Dept of Biology Tanner, Edmund; University of Cambridge, Plant Sciences Bujan, Jelena; University of Oklahoma, Griffin, Eric; University of Pittsburgh, Biological Sciences Mayor, Jordan; Smithsonian Tropical Research Institute Pasquini, Sarah; University of California Sheldrake, Merlin; University of Cambridge, Plant Sciences Garcia, Milton; Smithsonian Tropical Research Institute
Substantive Area:	Chemical and Biochemical Control < Ecosystems < Substantive Area
Organism:	Other (specify type in field below) < Angiosperms < Plants
Habitat:	Seasonal Forest < Tropical Zone < Terrestrial < Habitat
Geographic Area:	Mainland < Central America < Geographic Area
Additional Keywords:	Barro Colorado Nature Monument, fertilization, fine litter production, foliar nutrient concentrations, meta-analysis, nitrogen, old-growth forest, phosphorus, potassium, secondary forest, tree growth rates
Abstract:	We present a meta-analysis of plant responses to fertilization experiments conducted in lowland, species-rich, tropical forests and, for a 15-year factorial nitrogen (N), phosphorus (P) and potassium (K) fertilization conducted in central Panama, update a key result and present the first species-level analyses of tree growth rates. The update concerns community-level tree growth rates, which responded significantly to the addition of N and K together after 10 years of fertilization (Wright et al. 2011) but not after 15 years (this study). Our experimental soils are infertile for the region, and species whose regional distributions are strongly associated with low soil P availability dominate the local tree flora. Under these circumstances, we expect muted responses to fertilization, and we predicted species associated with low-P soils would respond most

slowly. The data did not support this prediction – species-level tree growth responses to P addition were unrelated to species-level soil P associations. The meta-analysis demonstrated that nutrient limitation is widespread in lowland tropical forests and evaluated two directional hypotheses concerning plant responses to N addition and to P addition. The meta-analysis supported the hypothesis that tree (or biomass) growth rate responses to fertilization are weaker in old growth forests and stronger in secondary forests, where rapid biomass accumulation provides a nutrient sink. The meta-analysis found no support for the long-standing hypothesis that plant responses are stronger for P addition and weaker for N addition. We do not advocate discarding the latter hypothesis. There are only 14 fertilization experiments from lowland, species-rich, tropical forests, 13 of the 14 experiments added nutrients for five or fewer years, and responses vary widely among experiments. Potential fertilization responses should be muted when the species present are well adapted to nutrient-poor soils, as is the case in our experiment, and when pest pressure increases with fertilization, as it does in our experiment. The statistical power and especially the duration of fertilization experiments conducted in old growth, tropical forests might be insufficient to detect the slow, modest growth responses that are to be expected.

Note: The following files were submitted by the author for peer review, but cannot be converted to PDF. You must view these files (e.g. movies) online.

MS\_Paffinity\_20171005\_DataS1.csv

SCHOLARONE™  
Manuscripts

View Only

## Plant responses to fertilization experiments in lowland, species-rich, tropical forests

Running head: Nutrient limitation in tropical forests

S. Joseph Wright<sup>1,9</sup>, Benjamin L. Turner<sup>1</sup>, Joseph B. Yavitt<sup>2</sup>, Kyle E. Harms<sup>3</sup>, Michael Kaspari<sup>4</sup>,  
Edmund V. J. Tanner<sup>1,5</sup>, Jelena Bujan<sup>4</sup>, Eric A. Griffin<sup>1,6,7</sup>, Jordan R. Mayor<sup>1,8</sup>, Sarah C.  
Pasquini<sup>6</sup>, Merlin Sheldrake<sup>1,5</sup> and Milton N. Garcia<sup>1</sup>

1. Smithsonian Tropical Research Institute, Apartado 0843-03092, Balboa, Republic of Panama
2. Department of Natural Resources, Cornell University, Ithaca, New York 14853 USA
3. Department of Biological Sciences, Louisiana State University, Baton Rouge, Louisiana 70803 USA
4. Department of Biology, University of Oklahoma, Norman, OK, 73019-0235 USA
5. Department of Plant Sciences, University of Cambridge, Cambridge CB2 3EA United Kingdom
6. Department of Biological Sciences, University of Pittsburgh, 4249 Fifth Ave, Pittsburgh, PA 15260 USA
7. Smithsonian Environmental Research Center, 647 Contees Wharf Rd., Edgewater, MD, 21307 USA
8. ICF, PO Box 4495, Arcata, CA 95518 USA
9. Email: wrightj@si.edu

**Abstract** – We present a meta-analysis of plant responses to fertilization experiments conducted in lowland, species-rich, tropical forests and, for a 15-year factorial nitrogen (N), phosphorus (P) and potassium (K) fertilization conducted in central Panama, update a key result and present the first species-level analyses of tree growth rates. The update concerns community-level tree growth rates, which responded significantly to the addition of N and K together after 10 years of fertilization (Wright et al. 2011) but not after 15 years (this study). Our experimental soils are infertile for the region, and species whose regional distributions are strongly associated with low soil P availability dominate the local tree flora. Under these circumstances, we expect muted responses to fertilization, and we predicted species associated with low-P soils would respond most slowly. The data did not support this prediction – species-level tree growth responses to P addition were unrelated to species-level soil P associations. The meta-analysis demonstrated that nutrient limitation is widespread in lowland tropical forests and evaluated two directional hypotheses concerning plant responses to N addition and to P addition. The meta-analysis supported the hypothesis that tree (or biomass) growth rate responses to fertilization are weaker in old growth forests and stronger in secondary forests, where rapid biomass accumulation provides a nutrient sink. The meta-analysis found no support for the long-standing hypothesis that plant responses are stronger for P addition and weaker for N addition. We do not advocate discarding the latter hypothesis. There are only 14 fertilization experiments from lowland, species-rich, tropical forests, 13 of the 14 experiments added nutrients for five or fewer years, and responses vary widely among experiments. Potential fertilization responses should be muted when the species present are well adapted to nutrient-poor soils, as is the case in our experiment, and when pest pressure increases with fertilization, as it does in our experiment. The statistical

power and especially the duration of fertilization experiments conducted in old growth, tropical forests might be insufficient to detect the slow, modest growth responses that are to be expected.

**Key words:** Barro Colorado Nature Monument, fertilization, fine litter production, foliar nutrient concentrations, meta-analysis, nitrogen, old growth forest, phosphorus, potassium, secondary forest, tree growth rates

For Review Only

## 1 **Introduction**

2           Tropical forests cover just 7% of the Earth's land surface but store 25% of terrestrial  
3 carbon and account for 33% of terrestrial net primary productivity (Bonan 2008). The vast  
4 majority of these forests spread across the humid lowlands of tropical Africa, Asia and the  
5 Americas and support tremendous numbers of species. How these lowland, species-rich, tropical  
6 forests respond to atmospheric and climate change will have profound implications for future  
7 global carbon and hydrological cycles (Bonan 2008), with the potential for nutrient supplies to  
8 limit future carbon sequestration being a crucial uncertainty (Wieder et al. 2015).

9           Comparative nutrient cycling studies generated the hypothesis that phosphorus (P) is  
10 limiting in many lowland tropical forests while nitrogen (N) is not. Briefly, leaf N:P ratios  
11 increase dramatically before abscission in many lowland tropical forests, suggesting more  
12 efficient reabsorption of P than N (McGroddy et al. 2004). Partly as a result, P tends to cycle  
13 more efficiently than N in fine litter in tropical lowland forests relative to temperate, boreal and  
14 tropical montane forests (Vitousek 1984, Vitousek and Sanford 1986). In addition, gaseous and  
15 hydrological losses of N tend to be much larger in tropical lowland forests than in temperate and  
16 boreal forests, suggesting N supplies exceed plant demand in tropical lowland forests (Houlton et  
17 al. 2006, Hedin et al. 2009). Finally, foliar P concentrations correlate strongly with total soil P  
18 stocks (Cleveland et al. 2011), and fine litter production increases with litter P but not litter N  
19 concentration in lowland tropical forests (Vitousek 1984). These comparative studies are  
20 consistent with the hypothesis that P is limiting in many lowland tropical forests while N is not.

21           At least two mechanisms contribute to the nutrient cycling differences observed between  
22 tropical lowland forests versus temperate, boreal and tropical montane forests. Bedrock and  
23 biological fixation are the primary sources of P and N, respectively, and Walker and Syers

24 (1976) showed that P availability declines during pedogenesis as bedrock decomposes and P  
25 leaches away and N availability increases as biological fixation develops. Rapid erosion keeps  
26 many montane soils in early stages of pedogenesis (Porder et al. 2007), and Quaternary  
27 glaciation exposed fresh bedrock and spread dust and till from ground bedrock over a large  
28 portion of temperate and boreal but not tropical latitudes (Vitousek 1984). In addition, warm,  
29 moist conditions enhance decomposition rates in the lowland tropics, preventing accumulation of  
30 unavailable organic N in an organic horizon. These regional differences in soil age and  
31 decomposition rates and the consistent results of comparative nutrient cycling studies motivate  
32 the hypothesis that lowland tropical forests growing on old, highly weathered Ultisols and  
33 Oxisols tend to be limited by P (or another rock derived nutrient) and not by N (Vitousek 1984,  
34 Vitousek and Sanford 1986, Vitousek et al. 2010). This hypothesis is now widely accepted (Elser  
35 et al. 2007), but experimental tests are few.

36       Operational definitions of nutrient limitation of biological processes include positive  
37 responses to experimental fertilization (Vitousek and Howarth 1991, Vitousek et al. 2010).  
38 Classic fertilization experiments confirmed the hypothesis that N and P limit Hawaiian forests  
39 growing on young versus old soils, respectively (Harrington et al. 2001). Strong, positive  
40 responses to N and P fertilization indicate N and P both limit plants in most terrestrial biomes  
41 (Elser et al. 2007, LeBauer and Treseder 2008). These meta-analyses report reasonable numbers  
42 of experiments for tropical forests, but these mainly concern montane forests, Hawaiian forests  
43 dominated by a single tree species, species-poor mangrove and planted forests, and even potted  
44 seedlings (see *Discussion: Experimental evidence for nutrient limitation*). Just 14 fertilization  
45 experiments have been conducted in lowland, species-rich, tropical forests (LSRTF), with more  
46 than half published in the 10 years since the two meta-analyses (Appendix S1: Tables S1 and

47 S2). In a third global meta-analysis, temperate and boreal forest plants respond strongly to  
48 potassium (K) fertilization; however, the single tropical study concerned potted seedlings  
49 (Tripler et al. 2006). Experimental evidence of nutrient limitation of plants in LSRTF is scant  
50 and a systematic review is lacking.

51 Here, we update a key result from our own factorial NPK fertilization experiment  
52 conducted in central Panama and perform a meta-analysis of the 14 fertilization experiments  
53 conducted in LSRTF. In our experiment, each added nutrient increased plant tissue  
54 concentrations of that nutrient; K addition (henceforth +K) decreased fine root biomass and  
55 increased rates of seedling growth, fine root turnover and decomposition; +P increased fine litter  
56 production and rates of photosynthesis, stomatal conductance and decomposition; combined N  
57 and P addition (henceforth +NP) increased seedling growth rates; +PK further increased stomatal  
58 conductance; and +NK ameliorated declining tree growth rates (Kaspari et al. 2008, Wright et al.  
59 2011, Yavitt et al. 2011, Pasquini and Santiago 2012, Santiago et al. 2012, Mayor et al. 2014,  
60 Turner et al. 2014, Pasquini et al. 2015). The key result that has changed as the fertilization  
61 treatments continue concerns tree growth rates, which are no longer responsive to +NK.

62 We also use newly available information on species-level responses to a natural gradient  
63 of P availability (Condit et al. 2013) to inform the first species-level analysis of tree growth rates  
64 for our experiment. We test the hypothesis that species whose regional distributions are  
65 associated with P-rich soils respond more strongly to P addition than do species associated with  
66 P-poor soils.

67 Our meta-analysis focuses on two hypotheses. The first hypothesis is that P limitation is  
68 stronger and N limitation is weaker in LSRTF. The second hypothesis is that nutrient limitation  
69 is stronger in secondary forests (and in high light tree-fall gaps) and weaker in old growth



70 forests. The second hypothesis is motivated by the large nutrient sink imposed by rapid biomass  
71 accumulation in secondary forests and tree-fall gaps. The meta-analysis is limited to N and P  
72 because just two fertilization experiments have considered any other nutrient for LSRTF. The  
73 meta-analysis is also limited to four types of responses – foliar N and P concentrations, fine litter  
74 N and P concentrations, fine litter production, and tree or biomass growth rates – because too  
75 few experiments have considered other types of responses. The meta-analysis indicates that N  
76 and P are equally likely to be limiting in LSRTF and are more likely to limit tree/biomass growth  
77 rates in secondary forests than in old growth forests.

78

## 79 **Methods**

### 80 *Meta-analysis*

81 We compiled 18 published articles from 14 fertilization experiments conducted in  
82 LSRTF (Appendix S1). For each experiment, we extracted a soil description, forest type  
83 (secondary or old growth), forest age for secondary forests, fertilizer type, number of years of  
84 fertilization, plot area and number, and any special circumstances. We tallied the number of  
85 significant responses to any type of fertilizer for four types of responses – foliar nutrient  
86 concentrations, fine litter nutrient concentrations, fine litter production and tree or biomass  
87 growth rates. We are limited to these four response types because too few studies documented  
88 any other response.

89 We also conducted a formal meta-analysis for the subset of experiments that compared  
90 control versus +N and/or +P treatments (or in one case –N and –P treatments versus a complete  
91 fertilizer). We extracted community-level and/or species-level treatment means, standard  
92 deviations and sample sizes for each response. We treated analyses for different species and for

93 different tree size categories as separate tests. We excluded community-level analyses if analyses  
94 were also partitioned by species or size. Several studies did not report standard deviations for  
95 particular responses and these responses were excluded (tree growth, Mirmanto et al. 1999;  
96 species-level growth, Gehring et al. 1999; all responses, Newbery et al. 2002; foliar nutrient  
97 concentrations, Davidson et al. 2004). Appendix S1, Metadata S1 and Data S1 present the  
98 articles and extracted data included in the meta-analysis.

99 We conducted the meta-analysis with the ‘metafor’ package (version 2.0-0, Viechtbauer  
100 2010) in R version 3.3.2. We used the `escalc()` function to calculate Hedge’s  $g$  and random effect  
101 models to estimate 95% confidence intervals for each response. Plant performance improved  
102 significantly with fertilization when lower 95% confidence intervals were positive. We added  
103 moderators to random effect models to test two directional hypotheses. Fertilization responses  
104 are (1) stronger for P addition than for N addition and (2) stronger for secondary forests than for  
105 old growth forests (see *Introduction* for rationale). Just two experiments conducted in secondary  
106 forests reported litter production and litter nutrient concentrations (Appendix S1: Table S2), so  
107 we could not isolate forest type for litter responses. We evaluated the first hypothesis with  
108 secondary and old growth forests pooled for all four types of response and for each forest type  
109 separately for foliar nutrient concentrations and tree/biomass growth. We also evaluated the  
110 second hypothesis for foliar nutrient concentrations and tree/biomass growth. We report Wald  
111 chi-square and one-tailed  $p$ -values for directional hypotheses. We followed the recommendations  
112 of Jennions et al. (2013) to evaluate potential publication bias.

### 113 *Study site*

114 Our experiment is located on the mainland in the Barro Colorado Nature Monument in  
115 central Panama (9° 06' 31" N, 79° 50' 37" W). Tree species composition and stature (canopy

116 heights up to 43 m) are characteristic of old growth (>200 years) forest. Aerial photographs  
117 confirm the presence of tall forest in 1927 (SJW, personal observation). The soils developed on  
118 Miocene basalt and transition from Oxisols (Typic Hapludox in Soil Taxonomy; Soil Survey  
119 Staff 1999) in the upper northeast corner of the 38.4-ha experimental plot to poorly drained  
120 Inceptisols (Aeric Epiaquepts) in the low lying, southwest corner (B. L. Turner, *unpublished*  
121 *data*). Although our experimental forest is fertile relative to many Amazonian forests (Wright et  
122 al. 2011), it is infertile for central Panama, with very low concentrations of exchangeable  
123 phosphate ( $< 1 \text{ mg P kg}^{-1}$  by resin extraction) and moderately low exchangeable K (Yavitt et al.  
124 2009, Condit et al. 2013, Mirabello et al. 2013, Turner et al. 2013, 2015).

### 125 *Experimental design*

126 We replicated the eight treatments of a 2x2x2 factorial NPK experiment four times.  
127 Within each replicate, we blocked the N, P, K and NPK treatments versus the NP, NK, PK and  
128 control treatments (see App. Fig. 1 in Wright et al. 2011). This balanced, incomplete-block  
129 design minimizes uncontrolled error associated with spatial variation, enables evaluation of main  
130 effects and two-way interactions, but limits power to evaluate the three-way interaction (Winer  
131 1971). The 32 experimental plots are each 40 by 40 m and are separated by a minimum distance  
132 of 40 m, with the exception of two plots separated by 20 m and a 2-m deep streambed.

133 Beginning in 1998, we added fertilizer by hand in four equal doses each wet season with  
134 6–8 weeks between applications (approximate dates May 15–30, July 1–15, September 1–15 and  
135 October 15–30). Nitrogen was added as coated urea ( $(\text{NH}_2)_2\text{CO}$ ), P as triple superphosphate  
136 ( $\text{Ca}(\text{H}_2\text{PO}_4)_2 \cdot \text{H}_2\text{O}$ ), and K as potassium chloride (KCl). Annual doses were  $125 \text{ kg N ha}^{-1} \text{ yr}^{-1}$ ,  
137  $50 \text{ kg P ha}^{-1} \text{ yr}^{-1}$  and  $50 \text{ kg K ha}^{-1} \text{ yr}^{-1}$ , which equals 69%, 470% and 88% of annual inputs from  
138 fine litter at a site 3-km to the north, respectively (Yavitt et al. 2004). Similar large additions of P

139 relative to annual litter inputs are standard practice in tropical nutrient addition experiments (see  
140 studies in Appendix S1: Tables S1 and S2) because tropical soils, including the soils at our site,  
141 tend to sequester large amounts of added P in forms believed to be inaccessible to plants  
142 (Mirabello et al. 2013).

143 We identified trees and measured diameter at breast height (DBH defined as 1.3 m) in  
144 1997, 1998, 1999, 2000, 2001 (DBH  $\geq$  100 mm only), 2003, 2008 and 2013, using the methods  
145 of Condit (1998). We recorded measurement height if buttresses or deformities prevented  
146 measurement at 1.3 m. We censused all trees with DBH  $\geq$  100 mm and, for a central 20-by-30 m  
147 subset of each plot, all trees with DBH  $\geq$  10 mm. We used the 1998, 2003, 2008 and 2013  
148 censuses to calculate relative growth rates (RGR) for three 5-yr census intervals as:

$$149 \quad \text{RGR} = (\ln(\text{DBH}_f) - \ln(\text{DBH}_i)) / (\text{DOC}_f - \text{DOC}_i) / 365.25,$$

150 where DOC refers to day of century of the corresponding DBH measurement and the subscripts  $f$   
151 and  $i$  refer to final and initial values, respectively. We excluded palms because diameter growth  
152 is absent or limited and dicots with broken main trunks, changes in measurement height, or  
153 multiple stems at the measurement height.

#### 154 *Analyses of community-level growth rates*

155 We used repeated measures ANOVA and a linear mixed effects analysis to analyze RGR.  
156 Repeated measures ANOVA is the appropriate analysis for our designed experiment. The  
157 response variable was the average RGR value for each plot-census interval combination,  
158 repeated measures were on census interval and main effects were nutrient treatments, their two-  
159 way interactions, blocks within replicates and all interactions with census interval. RGR tends to  
160 decline with DBH ( $r = -0.11$ ,  $n = 17,824$  in our data). To minimize this source of variation, we  
161 performed repeated measures ANOVAs for five relatively narrow size classes as follows: shrubs

162 and saplings ( $10 \text{ mm} \leq \text{DBH}_i < 25 \text{ mm}$ ), small poles ( $25 \text{ mm} \leq \text{DBH}_i < 50 \text{ mm}$ ), large poles ( $50$   
163  $\text{mm} \leq \text{DBH}_i < 100 \text{ mm}$ ), small trees ( $100 \text{ mm} \leq \text{DBH}_i < 250 \text{ mm}$ ) and large trees ( $\text{DBH}_i \geq 250$   
164  $\text{mm}$ ). We performed repeated measures ANOVAs with the ‘aov’ command in R 3.3.2. We  
165 repeated each analysis in SYSTAT© 11 (Richmond, CA) to evaluate the compound symmetry  
166 assumption, which was satisfied for all five size classes (Huynh-Feldt Epsilon close to 1).

167 The repeated measures ANOVAs partitioned by tree size suggested a possible  
168 fertilization-tree size interaction (see *Results: Community-level growth rates*). We used a linear  
169 mixed effects analysis to evaluate this possibility, incorporating initial size ( $\text{DBH}_i$ ) as a  
170 covariate. The response variable was the RGR value for each individual-census interval  
171 combination. Fixed effects were the N, P and K treatments, their two-way interactions and their  
172 interactions with  $\text{DBH}_i$ . Random effects were census interval, individual within species, and plot  
173 within block and replicate. We included species with 20 or more individuals. We compared AIC  
174 values for models that included all random effects and all possible combinations of fixed effects.

#### 175 *Analyses of species-level growth rates*

176 We used the P effect sizes of Condit et al. (2013) to characterize species-level P affinities  
177 for the regional species pool and the experimental forest. Condit et al. (2013) evaluated  
178 relationships between occurrence and soil fertility for 550 tree species, using 72 tree and soil  
179 surveys conducted within 50 km of our site. Strong effect sizes, with absolute values  $> 0.5$ ,  
180 characterized relationships between occurrence probability and soil resin P availability for 57.6%  
181 of the 550 species (Condit et al. 2013). To describe P affinity for the regional species pool, we  
182 examined the distribution of the 550 central Panama species among P effect sizes. To describe P  
183 affinity for the experimental forest, we examined the distribution of individual trees among P  
184 effect sizes.

185 We used a linear mixed effects analysis to evaluate relationships between species-level  
186 RGR, P addition and P affinity. We treated P effect sizes as a covariate representing species-  
187 level P affinity, P addition as a fixed effect, and census interval, individual within species and  
188 plot within block and replicate as random effects. We did not consider the N and K treatments  
189 because they were insignificant in the previous analysis (see *Results: Community-level growth*  
190 *rates*) and N and K effect sizes were consistently small in the analysis of Condit et al. (2013).  
191 We included species with four or more individuals in each P treatment. We compared AIC  
192 values for models that included all random effects and all combinations of P addition, P affinity  
193 and the interaction between P addition and P affinity. To avoid the compounding number of  
194 interactions associated with a second covariate, we performed this analysis for the four smaller  
195 size classes described previously (see *Methods: Analyses of community-level growth rates*). We  
196 excluded the largest size class because too few species had four or more large individuals in each  
197 P treatment.

#### 198 *Power analysis*

199 We conducted two simulations of our repeated measures ANOVA to evaluate statistical  
200 power. We used simulations because, to the best of our knowledge, a standard power analysis is  
201 not available for our incomplete block design (see *Methods: Experimental Design*). The first  
202 simulation used a common overall plot mean RGR value and its standard deviation (SD). The  
203 second simulation used plot mean RGR and SD values observed for each block and census  
204 interval. To simulate positive responses to P addition, we increased means observed for no-P  
205 plots by 1%, 3%, 5%, ... and 41% for +P plots. We then drew random values from normal  
206 distributions with the appropriate means and SDs for all 32 plots and performed the repeated

207 measures ANOVA. We repeated these steps 1,000 times for each percentage increase in +P  
208 means and tallied the number of times the main effect of P was significant.

209

## 210 **Results**

### 211 *Community-level growth rates*

212 The number of RGR values ranged from 330 to 6,633 for the five tree size classes  
213 (Appendix S2: Table S1). The main effects of N, P and K addition and their two-way interactions  
214 were insignificant for all five size classes in the repeated measures ANOVAs, although the main  
215 effect of K addition was marginally insignificant ( $p = 0.057$ ) for the largest trees (Appendix S2:  
216 Table S2). RGR varied significantly among census intervals for shrubs and saplings, with lower  
217 RGR in the second census interval (2003 to 2008), but did not vary significantly among census  
218 intervals for the four larger size classes (Appendix S2: Table S2). The repeated measures  
219 ANOVAs provide little to no statistical evidence that fertilizers affected RGR.

220 The relative growth rates of shrubs, saplings and small poles ( $< 50$  mm DBH) tended to  
221 be larger in the control treatment than in any of the nutrient addition treatments (Fig. 1). This  
222 insignificant tendency was absent for large poles and tended to be reversed for small and large  
223 trees (Fig. 1). This suggested a possible interaction between nutrient treatments and tree size. To  
224 evaluate this possibility, we performed a linear mixed effects analysis of RGR that included trees  
225 of all sizes and treated initial size ( $DBH_i$ ) as a covariate. This analysis included species  
226 represented by 20 or more individuals, 13,688 RGR measurements and 5,510 individual trees.  
227 The number of RGR values ranging from 73 to 208 (mean = 143) among plot-census interval  
228 combinations. The model that included  $DBH_i$  as a covariate minimized AIC, with  $\Delta AIC = 12$  for  
229 the next best model and  $\Delta AIC = 183$  for the null model that included just random effects

230 (Appendix S2: Table S3). The linear mixed effects analysis provides no statistical evidence that  
231 fertilizers affected RGR.

### 232 *Species-level growth rates*

233 The 550 species for which Condit et al. (2013) determined P effect sizes include 93.1%  
234 of the species and 98.3% of the individuals in our experimental forest. Species with strong  
235 positive P effect sizes ( $>0.5$ ) are associated with P-rich soils (high-P affinity) and comprised  
236 20% of the species in the regional species pool (Fig. 2A), but just 6% of the individuals in the  
237 experimental forest (Fig. 2B). Species associated with P-rich soils are underrepresented in the  
238 experimental forest relative to the regional species pool.

239 The number of RGR values ranged from 747 to 5,851 for the four tree size classes for the  
240 linear mixed effects analysis that included P affinity (Appendix S2: Table S4). The main effects  
241 of P addition, P affinity and their interaction were insignificant for all size classes (Fig. 2C,  
242 Appendix S2: Table S5). As in the repeated measures ANOVAs, RGR varied significantly  
243 among census intervals for shrubs and saplings, with lower RGR in the second census interval  
244 (2003 to 2008), but not for the three larger size classes (Appendix S2: Table S5).

### 245 *Power analysis*

246 Our repeated measures ANOVA had a 46% chance of detecting a 20% increase in RGR  
247 (Appendix S2: Fig. S1). Relaxing control of spatial and temporal variation associated with  
248 blocks and census intervals had little effect on power (Appendix S2: Fig. S1). This is consistent  
249 with the uniformly insignificant effects of blocks and replicates in the repeated measures  
250 ANOVAs (Appendix S2: Table S2).

251

252



253 *Meta-analysis*

254           Seven and nine of the 14 fertilization experiments enable isolation of N and/or P  
255 responses, respectively (Appendix S1: Tables S1 and S2). Sample sizes can be larger when  
256 single studies reported responses for multiple species or tree size classes. Sample sizes can also  
257 be smaller when only a subset of studies documented a particular response. There was no  
258 evidence for publication bias after accounting for heterogeneity between secondary and old  
259 growth forests (Appendix S1: Table S3).

260           All mean effect sizes were positive, indicating plants tend to be limited by N and by P in  
261 LSRTF (Fig. 3). Ten of the 16 mean effect sizes were individually significant ( $p < 0.05$ ). Four of  
262 the six insignificant effect sizes concerned responses to both +N and +P treatments for  
263 tree/biomass growth rates in old growth forests (Fig. 3F) and for litter concentrations of the same  
264 element (Fig. 3B).

265           We evaluated the interaction between forest type (secondary versus old growth) and  
266 fertilizer type (+N versus +P) and the directional hypothesis that responses are stronger in  
267 secondary forests and weaker in old growth forests for foliar nutrient concentrations and  
268 tree/biomass growth rates. Forest\*fertilizer interactions were insignificant for foliar nutrient  
269 concentrations (Fig. 3E;  $QM_1 = 0.287$ ,  $p = 0.296$ ) and for tree/biomass growth rates (Fig. 3F;  
270  $QM_1 = 0.255$ ,  $p = 0.307$ ). For foliar nutrient concentrations, the null hypothesis that fertilization  
271 responses were similar in secondary and old growth forests was accepted (Fig. 3E;  $QM_1 = 1.64$ ,  
272  $p = 0.100$  for pooled responses;  $QM_1 = 1.44$ ,  $p = 0.116$  for N responses to +N;  $QM_1 = 0.327$ ,  $p = 0.284$   
273 for P responses to +P). For tree/biomass growth rates, the null hypothesis that fertilization  
274 responses were similar in secondary and old growth forests was rejected (Fig. 3F;  $QM_1 = 8.41$ ,  
275  $p = 0.0019$  for responses to pooled fertilizers;  $QM_1 = 5.59$ ,  $p = 0.0091$  for responses to +N;

276  $QM_1=2.97, p=0.0423$  for responses to +P). Tree/biomass growth responses were significantly  
277 stronger in secondary forests and weaker in old growth forests (Fig. 3F).

278 We evaluated the directional hypothesis that responses are stronger for P addition and  
279 weaker for N addition for all four responses with secondary and old growth forests pooled and  
280 for foliar nutrient concentrations and tree/biomass growth rates for each forest type. The null  
281 hypothesis that responses are similar for +P and +N could never be rejected. Figures 3A through  
282 3D present tests with secondary and old growth forests pooled (Fig. 3A,  $QM_1=2.66, p=0.051$  for  
283 foliar nutrient concentrations; Fig. 3B,  $QM_1=0.153, p=0.348$  for fine litter nutrient  
284 concentrations; Fig. 3C,  $QM_1=0.111, p=0.370$  for fine litter production; Fig. 3D,  $QM_1=0.430,$   
285  $p=0.256$  for tree/biomass growth rates). Figure 3E presents foliar nutrient concentrations for each  
286 forest type ( $QM_1=0.0957, p=0.378$  for secondary and  $QM_1=2.35, p=0.063$  for old growth  
287 forests). Figure 3F presents tree/biomass growth rates for each forest type ( $QM_1=0.0253,$   
288  $p=0.437$  for secondary and  $QM_1=0.758, p=0.192$  for old growth forests). The meta-analysis  
289 provides no evidence that plant responses differ for +P versus +N treatments.

290

## 291 Discussion

292 When just two census intervals were available, our experiment included significant  
293 N\*K\*census interval interactions for growth for the three smaller tree size classes (Wright et al.  
294 2011). Growth rates were larger in the first census interval, and +NK ameliorated the decline to  
295 the second census interval (see Fig. 1 in Wright et al. 2011). Now, with three census intervals  
296 available, all N\*K\*interval interactions are insignificant and the second census interval has  
297 significantly slower growth rates for the smallest size class only (Appendix S2: Table S2). Our  
298 new analysis includes 50% more RGR values and years of fertilization and must replace the

299 earlier analysis. After 15 years of chronic nutrient additions, our experiment provides virtually  
300 no evidence that nutrients limit tree growth (Fig. 1, Appendix S2: Tables S2 and S3).

301 Our treatments have significantly increased soil nutrient availability and many measures  
302 of plant performance. Phosphorus addition increased soil phosphate availability by 2800%; +K  
303 increased K availability by 91%; and +N increased nitrate availability by 120% (Yavitt et al.  
304 2011, Mirabello et al. 2013, Turner et al. 2013). Species- and community-level plant responses,  
305 ranging from increases in metabolism (photosynthesis, stomatal conductance) to increases in  
306 production (fine litter), standing biomass and tissue turnover rates (fine roots) were reviewed  
307 earlier (see *Introduction*). We now consider why these increases in soil nutrient availability and  
308 plant performance do not lead to increased tree growth. We begin with a meta-analysis of 14  
309 fertilization experiments conducted in lowland, species-rich tropical forests (LSRTF).

#### 310 *Experimental evidence for nutrient limitation*

311 All 14 fertilization experiments address the hypothesis that some combination of  
312 nutrients limit plants in LSRTF, and a tally of statistically significant effects indicates that  
313 nutrient limitation is widespread (Appendix S1: Tables S1 and S2). Foliar and fine litter  
314 concentrations of at least one nutrient increased significantly in seven of eight and five of five  
315 experiments, respectively. Fine litter production increased significantly in five of seven  
316 experiments. Tree/biomass growth responses varied with forest environment. Tree/biomass  
317 growth rates increased significantly in seven of eight experiments conducted in secondary forests  
318 or with saplings in high light microsites in old growth forests. In contrast, six experiments  
319 conducted in old growth forests documented tree growth responses without finding significant  
320 responses for trees larger than 100-mm DBH and with a significant increase for smaller trees in  
321 just one study. The contrast between forest environments is significant (Fisher Exact Test,

322  $p=0.0256$  for trees  $< 100$ -mm DBH and  $p=0.00466$  for trees  $> 100$ -mm DBH). To summarize,  
323 fertilization is often associated with significant increases in tissue nutrient concentrations and  
324 fine litter production and, in secondary forests and tree-fall gaps, with significant increases in  
325 tree/biomass growth rates. There is, however, no experimental evidence for nutrient limitation of  
326 growth rates for trees larger than 100 mm DBH in old growth LSRTF.

327 Our formal meta-analysis of the subset of these experiments with +N or +P treatments  
328 produced broadly similar results. All 16 mean effect sizes were positive and 10 were strong  
329 (mean Hedge's  $g > 0.5$ ), indicating widespread limitation by N and by P (Fig. 3). The directional  
330 hypothesis that N and P limitation of tree/biomass growth rates is stronger in secondary forests  
331 and weaker in old growth forests was also supported (Fig. 3F), and once again there was no  
332 evidence for N or P limitation of tree growth rates in old growth forests (open circles in Fig. 3F).

333 Our meta-analysis produced two related surprises. Nitrogen limits foliar N concentrations  
334 and fine litter production in LSRTF (Figs. 3A and 3C), and there is no evidence that P limitation  
335 is stronger than N limitation in LSRTF (all tests in Fig. 3). This is consistent with a meta-  
336 analysis of terrestrial N addition experiments in which “The degree of N limitation in the  
337 remainder of the tropical forest studies [when Hawaiian forests on recent lava flows were  
338 excluded]... was comparable to that of temperate forests...” (LeBauer and Treseder 2008). This  
339 conclusion of LeBauer and Treseder (2008) and our own meta-analysis are inconsistent with a  
340 second meta-analysis in which “... most fertilization experiments in forests were conducted in  
341 tropical latitudes, and this habitat type had a stronger response to added P than added N,  
342 suggesting support for the long-held belief that tropical ecosystems on old soils are  
343 predominantly P limited (Walker and Syers 1976).” (Elser et al. 2007). To reconcile these

344 contrasting conclusions concerning N limitation, we examined the tropical forest studies in both  
345 earlier meta-analyses.

346 Both meta-analyses include experiments conducted in a wide range of tropical forest  
347 environments. Elser et al. (2007) include mangrove forests (3 studies), montane forests (8), a  
348 monospecific *Eucalyptus* stand (1) and seedlings planted into pots (2), abandoned land (2) and  
349 forest understory (1). Several of these studies are of questionable relevance to their conclusion  
350 concerning P limitation on old soils because the experimental soils are artificial (pot  
351 experiments) or relatively young (many montane forests, Porder et al. 2007). These 17 studies  
352 are also irrelevant to our interest in LSRTF. LeBauer and Treseder (2008) also include seven  
353 studies conducted in montane forests. Just six and three experiments conducted in LSRTF remain  
354 in the compilations of Elser et al. (2007) and LeBauer and Treseder (2008), respectively. Our  
355 meta-analysis included these experiments plus eight additional experiments conducted in  
356 LSRTF, and we believe the conclusions of our meta-analysis stand for LSRTF.

357 To summarize those conclusions, both N and P addition are associated with strong  
358 increases in foliar nutrient concentrations (Figs. 3A and 3E), fine litter production (Fig. 3C) and  
359 fine litter nutrient concentrations (Fig. 3B) in LSRTF. The increases in fine litter nutrient  
360 concentrations are highly variable, however, suggesting variation in concentrations and/or  
361 resorption among tissues and studies (Fig. 3B, Schreag et al. 2014, Alvarez-Clare et al. 2015).  
362 Both N and P addition are also associated with strong increases in tree/biomass growth in  
363 secondary forests, where rapid biomass accumulation ensures a nutrient sink (filled circles in  
364 Fig. 3F). There is, however, no evidence for the long-standing hypothesis that P limitation is  
365 stronger and N limitation is weaker in lowland tropical forests (Figs. 3A through 3F) nor for  
366 nutrient limitation of tree growth rates in old growth forests (open circles in Fig. 3F).

367 *Why is tree growth unresponsive to fertilization in old growth forests?*

368           At least four mutually compatible mechanisms might contribute to the absence of tree  
369 growth responses to fertilization in old growth LSRTF (open circles in Fig. 3F). The first  
370 concerns local species composition and potential growth responses. Species adapted to low  
371 resource levels tend to have limited potential to increase growth rates in response to increased  
372 resource levels (Coley et al. 1985). This could limit fertilization responses until species adapted  
373 to high nutrient soils arrive changing species composition (Chapin et al. 1986). At our  
374 experimental forest, species whose regional distributions are strongly associated with P-poor and  
375 P-rich soils comprise 47% and just 6% of the individual trees, respectively (Fig. 2B). We should  
376 expect modest and slow responses to P addition when species associated with P-poor soils  
377 dominate local species composition (Chapin et al. 1986, Kitayama 2005, Dalling et al. 2016).

378           A second possible mechanism for muted growth responses to fertilization concerns plant  
379 enemies. Fertilization often increases tissue nutrient concentrations (Figs. 3A, 3B and 3E),  
380 making fertilized plants more attractive to herbivores and possibly other pests. Two fertilization  
381 experiments conducted in LSRTF considered herbivory. Herbivory increased with +P in 10-year  
382 old forests in Mexico (Campo and Dirzo 2003) and with +K and +P in our experiment (Santiago  
383 et al. 2012). Potassium addition also reduced net adverse effects of foliar bacteria in our  
384 experiment (Griffin et al. 2016). Spatial scale becomes important if fertilized plants attract pests.  
385 Fertilizers applied to individual plants or small plots might create nutrient hotspots that recruit  
386 nearby pests. Most fertilization experiments are conducted at spatial scales of 10 to 50 m in  
387 LSRTF with measurements limited to a central core area (see plot sizes in Appendix S1: Tables  
388 S1 and S2). This might limit problems posed by immigration; however, enemy populations  
389 might still increase in large fertilized plots if their local demography changes. As an example,

390 forest floor arthropod abundance increased with +K and +P in the central area of our 1,600-m<sup>2</sup>  
391 experimental plots (Kaspari et al. 2017, also see Bujan et al. 2016). If fertilization increases pest  
392 pressure, those pests might consume increased primary production, limiting potential tree growth  
393 responses (Anderson et al. 2010).

394 As an aside, pest pressure might also contribute to a striking difference between growth  
395 responses to +P in our experiment versus a growing house experiment conducted with a subset of  
396 our species. In the growing house experiment, species-specific growth responses to +P increase  
397 steadily with the P effect sizes of Condit et al. (2013) (Zalamea et al. 2017). In our forest  
398 experiment, sapling growth responses were unrelated to these same P effect sizes (Fig. 2C).  
399 Species adapted to high resource levels tend to be poorly defended against herbivores and other  
400 pests (Coley et al. 1985). Pests that are absent from the growing house experiment might prevent  
401 species associated with P-rich soils from achieving their potential growth responses in the forest  
402 experiment. Of course, with so few individuals of species associated with P-rich soils present  
403 (Fig. 2B), our statistical power to evaluate their responses is also limited (Fig. 2C).

404 A third possible mechanism for muted growth responses to fertilization concerns time.  
405 Thirteen of the 14 fertilization experiments added nutrients for five or fewer years (Appendix S1:  
406 Tables S1 and S2) while tropical trees can live for centuries (Chambers 1998, Worbes and Junk  
407 1999). There is evidence for size-dependent responses to fertilization. Three studies fertilized *in*  
408 *situ* seedlings, and seedling growth rates increased in all three studies (Hättenschwiler 2002,  
409 Yavitt et al. 2008, Santiago et al. 2012). Five studies fertilized saplings only (Villagra et al. 2013,  
410 Chou et al. 2017) or partitioned growth analyses by tree size (Fisher et al. 2013, Alvarez-Clare et  
411 al. 2013, this study), and growth rates of saplings or the smallest tree size class increased in three  
412 of the five studies (Alvarez-Clare et al. 2013, Villagra et al. 2013, Chou et al. 2017). In contrast,

413 N and P fertilization had no effect on the growth rates of trees larger than 100 mm DBH in the  
414 six fertilization experiments that evaluated larger trees in old growth LSRTF (Appendix S1:  
415 Table S1). Large trees can accumulate large reserves of nutrients and carbohydrates, and many  
416 years might be required to capture their growth responses to nutrient addition.

417 The final reason for insignificant growth responses concerns statistical power. Our  
418 experiment has a reasonable chance of detecting a 20% increase in RGR (Appendix S2: Fig. S1).  
419 Sample sizes (see numbers of plots in Appendix S1: Tables S1 and S2) suggest power is likely to  
420 be similar or lower for 12 of the 13 remaining experiments conducted in LSRTF unless plot-to-  
421 plot variation is unexpectedly low. To summarize, the potential fertilization response of tropical  
422 forest trees will be limited if the species present are well adapted to nutrient-poor soils, as in our  
423 experiment (Fig. 2B), and if pest pressure increases with fertilization, as in our experiment  
424 (Santiago et al. 2012). The statistical power and especially the duration of fertilization  
425 experiments conducted in old growth, tropical forests might also be insufficient to detect the  
426 slow, modest growth responses that are to be expected.

#### 427 *Conclusions*

428 Our review of 14 fertilization experiments conducted in LSRTF indicates that nutrient  
429 limitation is widespread (Appendix S1: Tables S1 and S2). Nutrient availability is already likely  
430 to be limiting the ability of these forests to sequester carbon despite rising atmospheric CO<sub>2</sub>  
431 concentrations (Wieder et al. 2015). Our formal meta-analysis of the subset of these experiments  
432 that include +N and/or +P treatments supports the hypothesis that nutrient limitation is stronger  
433 in secondary forest and weaker in old growth forest, but does not support the hypothesis that P  
434 limitation is stronger and N limitation is weaker. As an aside, evidence for P limitation is also  
435 suspect because every experiment that included a +P treatment used simple or triple super



436 phosphate fertilizer. Super phosphate fertilizers supply calcium (Ca) and P at a 1:2 ratio of Ca to  
437 P. Bedrock is the primary source for Ca as well as P, and Ca like P might limit tropical forests  
438 growing on highly weathered soils (Vitousek 1984, Sanford and Vitousek 1986, Baillie et al.  
439 1987, Cuevas and Medina 1988). Nonetheless, we believe it would be premature to discard the  
440 hypothesis that P limitation is stronger than N limitation in lowland tropical forests for two  
441 reasons.

442 First, the number of fertilization experiments conducted in LSRTF remains small  
443 (Appendix S1: Tables S1 and S2) and most of the experiments share modest sample sizes and  
444 short durations (see *Discussion: Why is tree growth unresponsive to fertilization in old growth*  
445 *forests?*). Effect sizes tend to be larger for plant responses to P addition than to N addition;  
446 however, the difference is small and insignificant (Fig. 3). In contrast, a recent meta-analysis of  
447 microbial responses to fertilization experiments conducted in tropical forests found strong  
448 evidence for P limitation over all tropical forests and evidence for N limitation in montane but  
449 not lowland tropical forests (Camezind et al., in press). The contrasting generation times and  
450 fertilization responses of microbes and plants suggest that the responses of long-lived plants  
451 might strengthen as the duration of fertilization experiments increases.

452 The second reason we believe it would be premature to discard the hypothesis that plant  
453 limitation by P is stronger than limitation by N in lowland tropical forests concerns evidence  
454 from our own experiment after 15 years of chronic nutrient additions. Soil Ca availability is  
455 extraordinarily high in our control plots (averaging 1,690 mg kg<sup>-1</sup>, Yavitt et al. 2009), and the Ca  
456 added with the triple super phosphate fertilizer is not an issue. In contrast to the results of our  
457 meta-analysis, our own experiment provides much more evidence for P (and K) limitation than  
458 for N limitation. The evidence includes a wide range of bacterial, fungal, arthropod and plant

459 responses (summarized in Table S2 in Kaspari et al. 2017; additional responses in Schreeg et al.  
460 2014; Pasquini et al. 2015; Wurzburger and Wright 2015; Griffin et al. 2016, 2017; Bujan et al.  
461 2016; Sheldrake et al. 2017). Statistically significant plant responses are roughly equally divided  
462 between the +P and +K treatments, with just one significant response to the +N treatment (an  
463 increase in tissue N concentrations). We conclude that N rarely limits plant function at our site,  
464 and N addition is unlikely to affect tree growth in the future. We predict that the many significant  
465 plant responses to +P and +K will, with time, lead to significant increases in tree growth and net  
466 primary production. After 15 years, a marginally insignificant trend ( $p = 0.057$ ) suggests that  
467 growth responses might be developing first in response to K addition among trees larger than  
468 249 mm DBH (Fig. 1, Appendix S2: Table S2). Lloyd et al. (2015) recently hypothesized that K  
469 availability plays a key role determining tropical forest structure. We are now in the 20<sup>th</sup> year of  
470 our chronic nutrient addition treatments, and we plan to continue indefinitely to test these and  
471 other predictions.

472

### 473 **Acknowledgements**

474 We thank two anonymous reviewers for thoughtful comments. The Smithsonian Tropical  
475 Research Institute and three grants from the Smithsonian Institution Scholarly Studies and  
476 Restricted Endowment Programs provided funding.

477 **Literature Cited**

- 478 Alvarez-Clare, S., M. C. Mack, and M. Brooks. 2013. A direct test of nitrogen and phosphorus  
479 limitation to net primary productivity in a lowland tropical wet forest. *Ecology* 94:1540-  
480 1551.
- 481 Alvarez-Clare, S., and M. C. Mack. 2015. Do foliar, litter, and root nitrogen and phosphorus  
482 concentrations reflect nutrient limitation in a lowland tropical wet forest? *Plos One*  
483 10:e0123796.
- 484 Andersen, K. M., M. D. Corre, B. L. Turner, and J. W. Dalling. 2010. Plant–soil associations in a  
485 lower montane tropical forest: physiological acclimation and herbivore-mediated  
486 responses to nitrogen addition. *Functional Ecology* 24:1171-1180.
- 487 Baillie, I. C., P. S. Ashton, M. N. Court, J. A. R. Anderson, E. A. Fitzpatrick, and J. Tinsley.  
488 1987. Site characteristics and the distribution of tree species in mixed Dipterocarp forest  
489 on Tertiary sediments in central Sarawak, Malaysia. *Journal of Tropical Ecology* 3:201-  
490 220.
- 491 Bonan, G. B. 2008. Forests and climate change: forcings, feedbacks, and the climate benefits of  
492 forests. *Science* 320:1444-1449.
- 493 Bujan, J., S. J. Wright, and M. Kaspari. 2016. Biogeochemical drivers of Neotropical ant activity  
494 and diversity. *Ecosphere* 7.
- 495 Camenzind, T., S. Hattenschwiler, K. Treseder, A. Lehmann, and M. C. Rillig. In press. Nutrient  
496 limitation of soil microbial processes in tropical forests. *Ecological Monographs*.  
497 Doi:10.1002/ecm.1279.

- 498 Campo, J., and R. Dirzo. 2003. Leaf quality and herbivory responses to soil nutrient addition in  
499 secondary tropical dry forests of Yucatan, Mexico. *Journal of Tropical Ecology* 19:525-  
500 530.
- 501 Chambers, J. Q., N. Higuchi, and J. P. Schimel. 1998. Ancient trees in Amazonia. *Nature*  
502 391:135-136.
- 503 Chapin, F. S., P. M. Vitousek, and K. Vancleve. 1986. The nature of nutrient limitation in plant  
504 communities. *American Naturalist* 127:48-58.
- 505 Chou, C. B., L. O. Hedin, and S. W. Pacala. 2017. Functional groups, species, and light interact  
506 with nutrient limitation during tropical rainforest sapling bottleneck. *Journal of Ecology*.
- 507 Cleveland, C. C., A. R. Townsend, P. Taylor, S. Alvarez-Clare, M. M. C. Bustamante, G.  
508 Chuyong, S. Z. Dobrowski, P. Grierson, K. E. Harms, B. Z. Houlton, A. Marklein, W.  
509 Parton, S. Porder, S. C. Reed, C. A. Sierra, W. L. Silver, E. V. J. Tanner, and W. R.  
510 Wieder. 2011. Relationships among net primary productivity, nutrients and climate in  
511 tropical rain forest: a pan-tropical analysis. *Ecology Letters* 14:1313-1317.
- 512 Coley, P. D., J. P. Bryant, and F. S. Chapin III. 1985. Resource availability and plant  
513 antiherbivore defense. *Science* 230:895-900.
- 514 Condit, R. 1998. *Tropical forest census plots*. Springer-Verlag and RG Landes Company, Berlin.
- 515 Condit, R., B. M. J. Engelbrecht, D. Pino, R. Perez, and B. L. Turner. 2013. Species distributions  
516 in response to individual soil nutrients and seasonal drought across a community of  
517 tropical trees. *Proceedings of the National Academy of Sciences of the United States of*  
518 *America* 110:5064-5068.
- 519 Cuevas, E., and E. Medina. 1988. Nutrient dynamics within Amazonian forests II. Fine root-  
520 growth, nutrient availability and leaf litter decomposition. *Oecologia* 76:222-235.

- 521 Dalling, J. W., K. Heineman, O. R. Lopez, S. J. Wright, and B. L. Turner. 2016. Nutrient  
522 availability in tropical rain forests: the paradigm of phosphorus limitation. Pages 261-273  
523 in G. Goldstein and L. S. Santiago, editors. *Tropical tree physiology*. Springer.
- 524 Davidson, E. A., C. J. R. de Carvalho, I. C. G. Vieira, R. D. Figueiredo, P. Moutinho, F. Y.  
525 Ishida, M. T. P. dos Santos, J. B. Guerrero, K. Kalif, and R. T. Saba. 2004. Nitrogen and  
526 phosphorus limitation of biomass growth in a tropical secondary forest. *Ecological  
527 Applications* 14:S150-S163.
- 528 Elser, J. J., M. E. S. Bracken, E. E. Cleland, D. S. Gruner, W. S. Harpole, H. Hillebrand, J. T.  
529 Ngai, E. W. Seabloom, J. B. Shurin, and J. E. Smith. 2007. Global analysis of nitrogen  
530 and phosphorus limitation of primary producers in freshwater, marine and terrestrial  
531 ecosystems. *Ecology Letters* 10:1135-1142.
- 532 Fisher, J. B., Y. Malhi, I. C. Torres, D. B. Metcalfe, M. J. van de Weg, P. Meir, J. E. Silva-  
533 Espejo, and W. H. Huasco. 2013. Nutrient limitation in rainforests and cloud forests  
534 along a 3,000-m elevation gradient in the Peruvian Andes. *Oecologia* 172:889-902.
- 535 Gehring, C., M. Denich, M. Kanashiro, and P. L. G. Vlek. 1999. Response of secondary  
536 vegetation in Eastern Amazonia to relaxed nutrient availability constraints.  
537 *Biogeochemistry* 45:223-241.
- 538 Griffin, E. A., M. B. Traw, P. J. Morin, J. N. Pruitt, S. J. Wright, and W. P. Carson. 2016. Foliar  
539 bacteria and soil fertility mediate seedling performance: a new and cryptic dimension of  
540 niche differentiation. *Ecology* 97:2998-3008.
- 541 Griffin, E. A., S. J. Wright, P. J. Morin, and W. P. Carson. 2017. Pervasive interactions between  
542 foliar microbes and soil nutrients mediate leaf production and herbivore damage in a  
543 tropical forest. *New Phytologist* 216:99-112.

- 544 Harrington, R. A., J. H. Fownes, and P. M. Vitousek. 2001. Production and resource use  
545 efficiencies in N-and P-limited tropical forests: a comparison of responses to long-term  
546 fertilization. *Ecosystems* 4:646-657.
- 547 Hattenschwiler, S. 2002. Liana seedling growth in response to fertilisation in a neotropical forest  
548 understorey. *Basic and Applied Ecology* 3:135-143.
- 549 Hedin, L. O., E. N. J. Brookshire, D. N. L. Menge, and A. R. Barron. 2009. The Nitrogen  
550 Paradox in Tropical Forest Ecosystems. *Annual Review of Ecology Evolution and*  
551 *Systematics* 40:613-635.
- 552 Houlton, B. Z., D. M. Sigman, and L. O. Hedin. 2006. Isotopic evidence for large gaseous  
553 nitrogen losses from tropical rainforests. *Proceedings of the National Academy of*  
554 *Sciences of the United States of America* 103:8745-8750.
- 555 Jennions, M. D., C. J. Lortie, M. S. Rosenberg, and H. R. Rothstein. 2013. Publication and  
556 related biases. Pages 207-236 in J. Koricheva, J. Gurevitch, and K. Mengersen, editors.  
557 *Handbook of meta-analysis in ecology and evolution*. Princeton University Press,  
558 Princeton, NJ.
- 559 Kaspari, M., M. N. Garcia, K. E. Harms, M. Santana, S. J. Wright, and J. B. Yavitt. 2008.  
560 Multiple nutrients limit litterfall and decomposition in a tropical forest. *Ecology Letters*  
561 11:35-43.
- 562 Kaspari, M., J. Bujan, M. D. Weiser, D. Ning, S. T. Michaletz, Z. L. He, B. J. Enquist, R. B.  
563 Waide, J. Z. Zhou, B. L. Turner, and S. J. Wright. 2017. Biogeochemistry drives diversity  
564 in the prokaryotes, fungi, and invertebrates of a Panama forest. *Ecology* 98:2019-2028.
- 565 Kitayama, K. 2005. Comment on "Ecosystem properties and forest decline in contrasting long-  
566 term chronosequences". *Science* 308.

- 567 LeBauer, D. S., and K. K. Treseder. 2008. Nitrogen limitation of net primary productivity in  
568 terrestrial ecosystems is globally distributed. *Ecology* 89:371-379.
- 569 Lloyd, J., T. F. Domingues, F. Schrodt, F. Y. Ishida, T. R. Feldpausch, G. Saiz, C. A. Quesada,  
570 M. Schwarz, M. Torello-Raventos, M. Gilpin, B. S. Marimon, B. H. Marimon-Junior, J.  
571 A. Ratter, J. Grace, G. B. Nardoto, E. Veenendaal, L. Arroyo, D. Villarroel, T. J. Killeen,  
572 M. Steininger, and O. L. Phillips. 2015. Edaphic, structural and physiological contrasts  
573 across Amazon Basin forest-savanna ecotones suggest a role for potassium as a key  
574 modulator of tropical woody vegetation structure and function. *Biogeosciences* 12:6529-  
575 6571.
- 576 Mayor, J. R., S. Joseph Wright, and B. L. Turner. 2014. Species-specific responses of foliar  
577 nutrients to long-term nitrogen and phosphorus additions in a lowland tropical forest.  
578 *Journal of Ecology* 102:36-44.
- 579 McGroddy, M. E., T. Daufresne, and L. O. Hedin. 2004. Scaling of C : N : P stoichiometry in  
580 forests worldwide: Implications of terrestrial redfield-type ratios. *Ecology* 85:2390-2401.
- 581 Mirabello, M. J., J. B. Yavitt, M. Garcia, K. E. Harms, B. L. Turner, and S. J. Wright. 2013. Soil  
582 phosphorus responses to chronic nutrient fertilisation and seasonal drought in a humid  
583 lowland forest, Panama. *Soil Research* 51:215-221.
- 584 Mirmanto, E., J. Proctor, J. Green, L. Nagy, and Suriantata. 1999. Effects of nitrogen and  
585 phosphorus fertilization in a lowland evergreen rainforest. *Philosophical Transactions of*  
586 *the Royal Society of London Series B-Biological Sciences* 354:1825-1829.
- 587 Newbery, D. M., G. B. Chuyong, J. J. Green, N. C. Songwe, F. Tchuenteu, and L. Zimmermann.  
588 2002. Does low phosphorus supply limit seedling establishment and tree growth in

- 589 groves of ectomycorrhizal trees in a central African rainforest? *New Phytologist* 156:297-  
590 311.
- 591 Pasquini, S., and L. Santiago. 2012. Nutrients limit photosynthesis in seedlings of a lowland  
592 tropical forest tree species. *Oecologia* 168:311-319.
- 593 Pasquini, S. C., S. J. Wright, and L. S. Santiago. 2015. Lianas always outperform tree seedlings  
594 regardless of soil nutrients: results from a long-term fertilization experiment. *Ecology*  
595 96:1866-1876.
- 596 Porder, S., P. M. Vitousek, O. A. Chadwick, C. P. Chamberlain, and G. E. Hilley. 2007. Uplift,  
597 erosion, and phosphorus limitation in terrestrial ecosystems. *Ecosystems* 10:158-170.
- 598 Santiago, L. S., S. J. Wright, K. E. Harms, J. B. Yavitt, C. Korine, M. N. Garcia, and B. L.  
599 Turner. 2012. Tropical tree seedling growth responses to nitrogen, phosphorus and  
600 potassium addition. *Journal of Ecology* 100:309-316.
- 601 Schreeg, L. A., L. S. Santiago, S. J. Wright, and B. L. Turner. 2014. Stem, root, and older leaf  
602 N:P ratios are more responsive indicators of soil nutrient availability than new foliage.  
603 *Ecology* 95:2062-2068.
- 604 Sheldrake, M., N. P. Rosenstock, D. Revillini, P. A. Olsson, S. J. Wright, and B. L. Turner.  
605 2017. A phosphorus threshold for mycoheterotrophic plants in tropical forests.  
606 *Proceedings of the Royal Society B-Biological Sciences* 284.
- 607 Soil Survey Staff. 1999. *Soil Taxonomy: A Basic System of Soil Classification for Making and*  
608 *Interpreting Soil Surveys*. United States Department of Agriculture - Natural Resources  
609 Conservation Service.
- 610 Tripler, C. E., S. S. Kaushal, G. E. Likens, and M. T. Walter. 2006. Patterns in potassium  
611 dynamics in forest ecosystems. *Ecology Letters* 9:451-466.



- 612 Turner, B. L., J. B. Yavitt, K. E. Harms, M. N. Garcia, T. E. Romero, and S. J. Wright. 2013.  
613 Seasonal Changes and Treatment Effects on Soil Inorganic Nutrients Following a Decade  
614 of Fertilizer Addition in a Lowland Tropical Forest. *Soil Science Society of America*  
615 *Journal* 77:1357-1369.
- 616 Turner, B. L., and S. J. Wright. 2014. The response of microbial biomass and hydrolytic  
617 enzymes to a decade of nitrogen, phosphorus, and potassium addition in a lowland  
618 tropical rain forest. *Biogeochemistry* 117:115-130.
- 619 Turner, B. L., J. B. Yavitt, K. E. Harms, M. N. Garcia, and S. J. Wright. 2015. Seasonal changes  
620 in soil organic matter after a decade of nutrient addition in a lowland tropical forest.  
621 *Biogeochemistry* 123:221-235.
- 622 Viechtbauer, W. 2010. Conducting meta-analyses in R with the metafor package. *Journal of*  
623 *Statistical Software* 36:1-48.
- 624 Villagra, M., P. I. Campanello, L. Montti, and G. Goldstein. 2013. Removal of nutrient  
625 limitations in forest gaps enhances growth rate and resistance to cavitation in subtropical  
626 canopy tree species differing in shade tolerance. *Tree Physiology* 33:285-296.
- 627 Vitousek, P. M. 1984. Litterfall, nutrient cycling and nutrient limitation in tropical forests.  
628 *Ecology* 65:285-298.
- 629 Vitousek, P. M., and R. W. Howarth. 1991. Nitrogen limitation on land and in the sea: how can it  
630 occur? *Biogeochemistry* 13:87-115.
- 631 Vitousek, P. M., S. Porder, B. Z. Houlton, and O. A. Chadwick. 2010. Terrestrial phosphorus  
632 limitation: mechanisms, implications, and nitrogen-phosphorus interactions. *Ecological*  
633 *Applications* 20:5-15.

- 634 Vitousek, P. M., and R. L. Sanford. 1986. Nutrient cycling in moist tropical forest. *Annual*  
635 *Review of Ecology and Systematics* 17:137-167.
- 636 Walker, T. W., and J. K. Syers. 1976. Fate of phosphorus during pedogenesis. *Geoderma* 15:1-  
637 19.
- 638 Wieder, W. R., C. C. Cleveland, W. K. Smith, and K. Todd-Brown. 2015. Future productivity  
639 and carbon storage limited by terrestrial nutrient availability. *Nature Geoscience* 8:441-  
640 445.
- 641 Winer, B. J. 1971. *Statistical principles in experimental design*. McGraw-Hill Book Company,  
642 New York.
- 643 Worbes, M., and W. Johannes Junk. 1999. How old are tropical trees? The persistence of a myth.  
644 *Iawa Journal* 20:255-260.
- 645 Wright, S. J., J. B. Yavitt, N. Wurzburger, B. L. Turner, E. V. J. Tanner, E. J. Sayer, L. S.  
646 Santiago, M. Kaspari, L. O. Hedin, K. E. Harms, M. N. Garcia, and M. D. Corre. 2011.  
647 Potassium, phosphorus, or nitrogen limit root allocation, tree growth, or litter production  
648 in a lowland tropical forest. *Ecology* 92:1616-1625.
- 649 Wurzburger, N., and S. J. Wright. 2015. Fine-root responses to fertilization reveal multiple  
650 nutrient limitation in a lowland tropical forest. *Ecology* 96:2137-2146.
- 651 Yavitt, J. B., K. E. Harms, M. N. Garcia, M. J. Mirabello, and S. J. Wright. 2011. Soil fertility  
652 and fine root dynamics in response to 4 years of nutrient (N, P, K) fertilization in a  
653 lowland tropical moist forest, Panama. *Austral Ecology* 36:433-445.
- 654 Yavitt, J. B., K. E. Harms, M. N. Garcia, S. J. Wright, F. He, and M. J. Mirabello. 2009. Spatial  
655 heterogeneity of soil chemical properties in a lowland tropical moist forest, Panama.  
656 *Australian Journal of Soil Research* 47:674-687.

- 657 Yavitt, J. B., and S. J. Wright. 2008. Seedling growth responses to water and nutrient  
658 augmentation in the understorey of a lowland moist forest, Panama. *Journal of Tropical*  
659 *Ecology* 24:19-26.
- 660 Yavitt, J. B., S. J. Wright, and R. K. Wieder. 2004. Seasonal drought and dry-season irrigation  
661 influence leaf-litter nutrients and soil enzymes in a moist, lowland forest in Panama.  
662 *Austral Ecology* 29:177-188.
- 663 Zalamea, P. C., B. L. Turner, K. Winter, F. A. Jones, C. Sarmiento, and J. W. Dalling. 2016.  
664 Seedling growth responses to phosphorus reflect adult distribution patterns of tropical  
665 trees. *New Phytologist* 212:400-408.

## Figure Captions

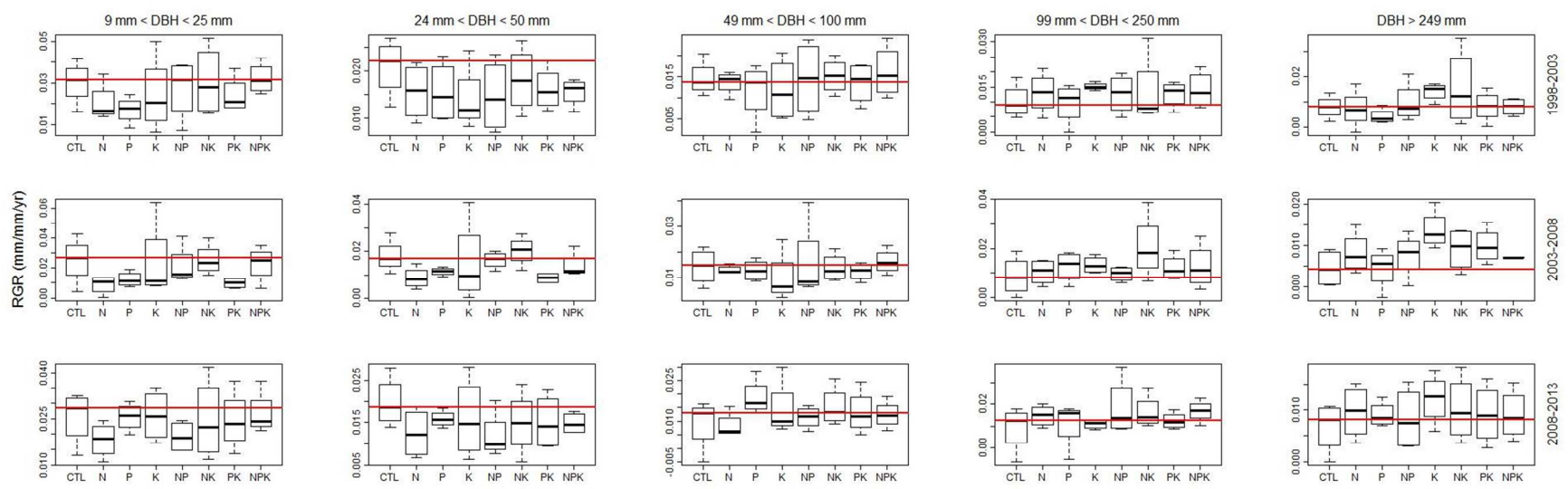
**Figure 1.** Bar-and-whisker plots of mean relative growth rates (RGR) for five size classes (columns) and three census intervals (rows). The solid red lines represent median RGR for the control treatment for each size class and census interval. The three census intervals are 1998-2003 (top row), 2003-2008 (middle row) and 2008-2013 (final row). The five size classes are shrubs and saplings (first column,  $10 \text{ mm} \leq \text{DBH}_i < 25 \text{ mm}$ ), small poles (second column,  $25 \text{ mm} \leq \text{DBH}_i < 50 \text{ mm}$ ), large poles (third column,  $50 \text{ mm} \leq \text{DBH}_i < 100 \text{ mm}$ ), small trees (fourth column,  $100 \text{ mm} \leq \text{DBH}_i < 250 \text{ mm}$ ) and large trees (final column,  $\text{DBH}_i \geq 250 \text{ mm}$ ). In the first four columns, treatments order is control, one nutrient (+N, +P and +K), two nutrients (+NP, +NK and +PK) and three nutrients (+NPK). In the final column, treatment order groups –K (control, +N, +P and +NP) versus +K (+K, +NK, +PK and +NPK) treatments to illustrate the marginally significant effect of K addition. Thick horizontal lines represent medians, boxes represent the interquartile range (25% to 75%) and whiskers represent extreme values.

**Figure 2.** Histograms of the distributions of species in the regional species pool (panel A) and individuals in the 38.4-ha experimental plot (panel B) with respect to species-level phosphorus affinity and the relationship between species-level phosphorus affinity and the ratio of mean relative growth rates (RGR) of conspecifics with versus without added phosphorus ( $\text{RGR}_{+P} : \text{RGR}_{-P}$ ) for shrubs and saplings ( $10 \text{ mm} \leq \text{DBH} < 25 \text{ mm}$ ) (panel C). In panel C, each species is represented by four or more individuals in each phosphorus treatment, and the horizontal dashed line represents equal RGR values in both phosphorus treatments. The orange and blue vertical lines represent strong positive and negative phosphorus affinity thresholds, respectively.

**Figure 3.** Meta-analysis of nitrogen (N) and phosphorus (P) responses for fertilization experiments conducted in lowland, species-rich, tropical forests (see Data S1). Points are mean

effect sizes, with 95% confidence intervals. Filled and empty points identify significant (lower 95% CI > 0) and insignificant effect sizes, respectively. The vertical axis label identifies the type of response (foliar or fine litter nutrient concentrations, fine litter production, or tree/biomass growth (RGR)), the added nutrient in square brackets (+N or +P), the type of forest (secondary, old growth or pooled), and sample size in parentheses. The first three gray horizontal lines separate four tests of the directional hypothesis that responses are stronger for P addition and weaker for N addition, with secondary and old growth forests pooled (sections A, B, C and D). The final gray horizontal line separates tests for interactions between fertilizer type (+N or +P) and forest type (secondary or old growth) and for the directional hypothesis that responses are stronger for secondary forests and weaker for old growth forests for foliar nutrient concentrations (E) and tree/biomass growth rates (F). Asterisks identify significant (\*  $p < 0.05$ , \*\*  $p < 0.01$ ) and 'ns' insignificant outcomes.

Figure 1



Only

Figure 2

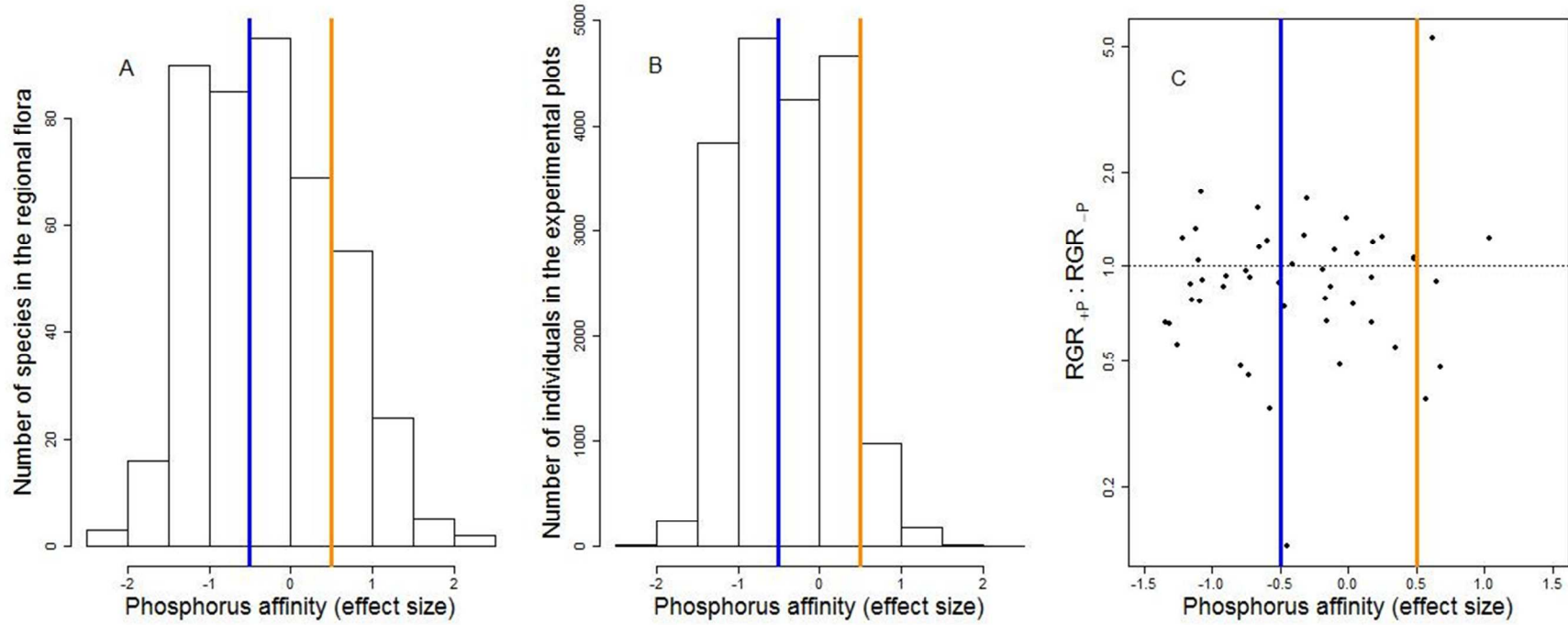
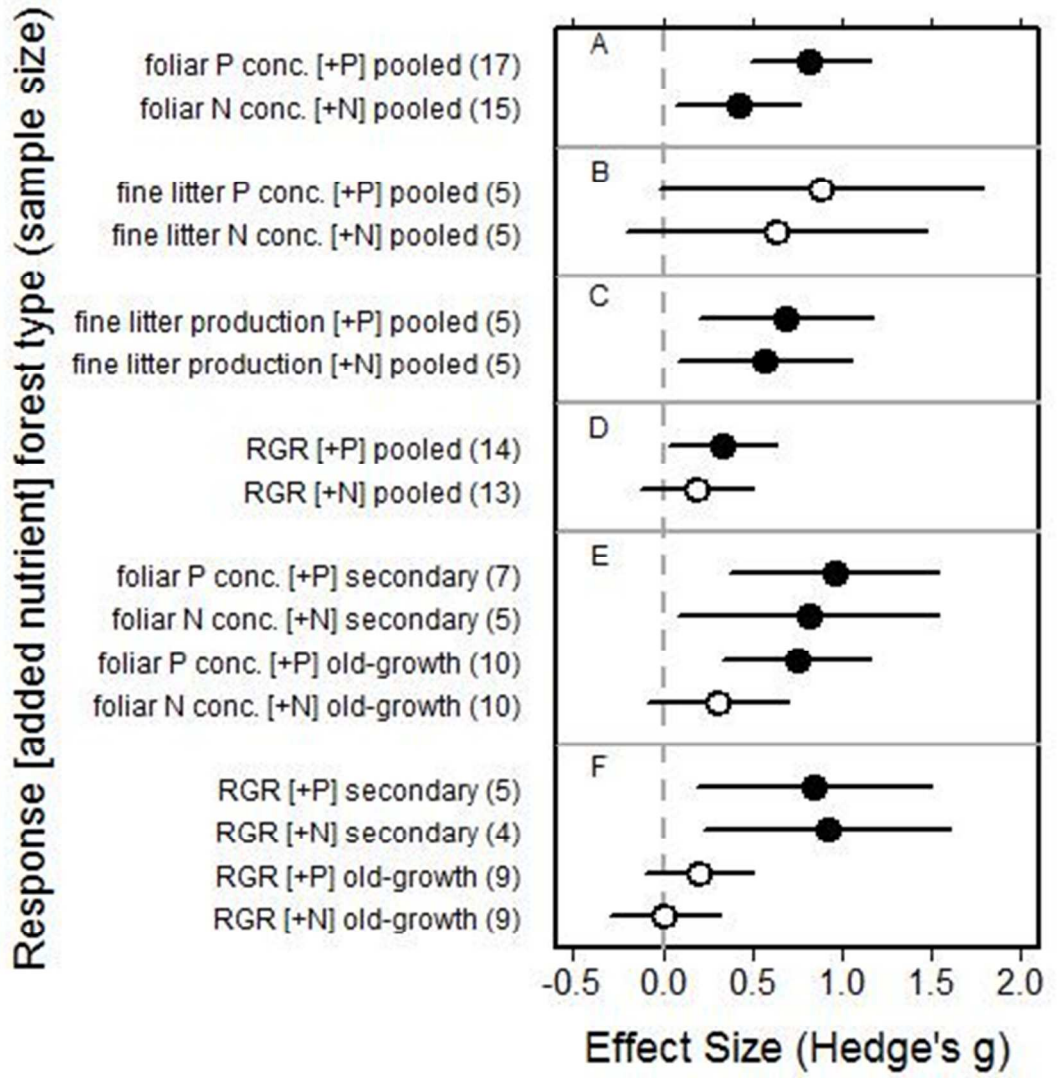


Figure 3





Appendix S1 presents summaries of 18 articles that contributed data to the meta-analysis and analyses for potential publication bias. Table S1 summarizes eight experiments conducted in old-growth forests. Table S2 summarizes six experiments conducted in secondary forests. Table S3 presents the analyses for potential publication bias. Table S4 presents full references and a key to the articles used in Data S1.

For Review Only

Appendix S1: Table S1. Summary of fertilization experiments conducted in old growth, lowland (&lt;400 m elevation), species-rich tropical forests.

Site	Soils [Special circumstances]	Fertilizer	Plot size (m) [and number]	Years fertilizer applied	Statistically Significant Effects			Reference
					Tissue nutrient concentrations	Litter Production	Tree Growth	
El Verde, Puerto Rico	Oxisols & Ultisols	complete	20 by 20 [8]	4	Not studied	Increases with complete fertilizer	Insignificant	Walker et al. 1996; Li et al. 2006
La Selva, Costa Rica	Ultisols [Saplings]	complete	Individual saplings	2.5	Not studied	Not studied	Increases in high light with complete fertilizer.	Chou et al. 2017
Iguazú, Argentina	Ultisols [forest gap saplings]	N and P together	15 by 15 [10]	5	Not studied	Not studied	Increases in high light with +NP.	Villagra et al. 2013
Korup, Cameroon	No soil type [Dominant trees ectomycorrhizal]	P	50 by 50 [14]	2	+P increases foliar and litter P.	Insignificant	Insignificant	Newbery et al. 2002
Kalimantan, Borneo	Yellow, sandy	Factorial N and P	50 by 50 [20]	4	+N, +P & +NP increase litter P & N.	Increases with +N, +P & +NP	Insignificant	Mirmanto et al. 1999
Tombopata, Peru	Alluvial terrace	Factorial N and P	Individual trees	4	Insignificant	Not studied	Insignificant <sup>@</sup>	Fisher et al. 2013
Limón, Costa Rica	Clayey, volcanic origin	Factorial N and P	30 by 30 [24]	3	+N (+P) increases foliar N (P) in selected species.	Insignificant	Increases with +P for small trees. Trees >100 mm DBH unaffected.	Alvarez-Clare et al. 2013, 2015
BCNM, Panama	Oxisols & Inceptisols	Factorial N, P & K	40 by 40 [32]	15	+P increases foliar & litter P.	Increases with +P	Insignificant	This study <sup>^</sup>

Appendix S1: Table S1. Continued.

@ We discount a marginally significant increase reported for +NP for all trees because (1) results were insignificant when partitioned by tree size, (2) the all-tree analysis did not control variation in tree size, (3) an outlier was excluded from the all-tree analysis to attain marginal significance ( $p=0.05$ ), and (4) methods used to identify the outlier were not presented.

^ Sources for the BCNM experiment are Kaspari et al. (2008), Wright et al. (2011), Mayor et al. (2014) and this study.

For Review Only

Appendix S1: Table S2. Summary of fertilization experiments conducted in secondary, lowland (&lt;400 m elevation), species-rich tropical forests.

Site	Soils	Fertilizer	Years fertilizer applied	Plot size (m) [and number]	Forest age (yrs)	Statistically Significant Effects			Reference
						Tissue nutrient concentrations	Litter production	Tree/biomass growth	
San Carlos de Rio Negro, Venezuela	Oxisol	NPK	0.33 <sup>@</sup>	1.5 by 1.5 [8]	0	Not studied	Not studied	Increases with +NPK	Uhl 1987
Igarapé Açu, Pará, Brazil	Typic Kandiuult ~70% sand by mass	Complete fertilizer combined with -1 treatments	2	6 by 7 [80]	0	+P increases foliar P. Litter not studied.	Not studied	Increases with +P	Gehring et al. 1999
Yucatan, Mexico	Shallow, organic rich directly over limestone	Factorial N and P	3	12 by 12 [16]	10	+P increases foliar & litter P	Increases with +NP	Increases with +N, +P & +NP	Campo & Dirzo 2003; Campo & Vázquez-Yanes 2004; Campo et al. 2007
				12 by 12 [16]	60	+NP increases foliar & litter P	Increases with +NP	Increases with +N, +P & +NP	
Paragominas, Pará, Brazil	Kaolinitic yellow Latosols	Factorial N and P	2	20 by 20 [12]	6	+P (+N) increases foliar P (N). Litter not studied.	Not studied	Increases with +N	Davidson et al. 2004
Paragominas, Pará, Brazil	Oxisol	P	2	20 by 20 [6]	24	Not studied	Not studied	Insignificant	Markewitz et al. 2012

Appendix S1: Table S3. Analyses for publication bias. Significant and marginally significant rank and regression tests (bolded and italicized p-values, respectively) indicate potential publication bias. All significant tests pool forest types; however, and significance disappears when forest type is included as a moderator (see rows with forest type recorded as “SF vs. OG”). We conclude that study heterogeneity associated with stronger responses in secondary forests (SF) and weaker responses in old-growth forests (OG) causes significant tests with SF and OG forest types pooled and that publication bias is not a problem.

Response	Treatment	Forest type	Panel in Fig. 3	Fail-safe number	Rank test		Regression test	
					Kendal $\tau$	p-value	z-value	p-value
Foliar P conc.	+P	Pooled	A	107	0.31	<i>0.091</i>	0.23	0.82
Foliar P conc.	+P	SF vs. OG	E	-	-	-	-0.32	0.75
Foliar N conc.	+N	Pooled	A	11	0.51	<b>0.0083</b>	1.83	<i>0.067</i>
Foliar N conc.	+N	SF vs. OG	E	-	-	-	1.52	0.13
Litter P conc.	+P	Pooled	B	-	0.60	0.23	1.22	0.22
Litter N conc.	+N	Pooled	B	-	0.20	0.82	0.86	0.39
Litter production	+P	Pooled	C	5	0.20	0.82	0.060	0.95
Litter production	+N	Pooled	C	2	0.60	0.23	0.58	0.56
RGR	+P	Pooled	D	5	0.49	<b>0.014</b>	1.97	<b>0.049</b>
RGR	+P	SF vs. OG	F	-	-	-	1.15	0.25
RGR	+N	Pooled	D	-	0.44	<b>0.042</b>	2.32	<b>0.020</b>
RGR	+N	SF vs. OG	F	-	-	-	1.37	0.17

*Notes:* The fail-safe number is the number of studies with a mean effect size of zero required to negate the significance of an observed effect size. We calculated fail-safe numbers for responses with significant effect sizes (Fig. 3) using the method of Rosenberg (Jennions et al. 2013). The rank test is the nonparametric correlation across studies between standardized effect sizes and standard errors. The regression test is a linear regression across studies between standard normal deviates of effect sizes and standard errors. Significant relationships between these measures of effect size and standard errors might indicate publication bias (Jennions et al. 2013).

Appendix S1: Table S4. References cited in Appendix S1: Tables S1 and S2. The bold numbers in parentheses after each reference refer to Supplement 1.

Alvarez-Clare, S., and M. C. Mack. 2015. Do foliar, litter, and root nitrogen and phosphorus concentrations reflect nutrient limitation in a lowland tropical wet forest? *Plos One* 10:e0123796. **(7)**

Alvarez-Clare, S., M. C. Mack, and M. Brooks. 2013. A direct test of nitrogen and phosphorus limitation to net primary productivity in a lowland tropical wet forest. *Ecology* 94:1540-1551. **(6)**

Campo, J., and R. Dirzo. 2003. Leaf quality and herbivory responses to soil nutrient addition in secondary tropical dry forests of Yucatan, Mexico. *Journal of Tropical Ecology* 19:525-530. **(12)**

Campo, J., E. Solis, and M. G. Valencia. 2007. Litter N and P dynamics in two secondary tropical dry forests after relaxation of nutrient availability constraints. *Forest Ecology and Management* 252:33-40. **(11)**

Campo, J., and C. Vazquez-Yanes. 2004. Effects of nutrient limitation on aboveground carbon dynamics during tropical dry forest regeneration in Yucatan, Mexico. *Ecosystems* 7:311-319. **(10)**

Chou, C. B., L. O. Hedin, and S. W. Pacala. 2017. Functional groups, species, and light interact with nutrient limitation during tropical rainforest sapling bottleneck. *Journal of Ecology*.

Davidson, E. A., C. J. R. de Carvalho, I. C. G. Vieira, R. D. Figueiredo, P. Moutinho, F. Y. Ishida, M. T. P. dos Santos, J. B. Guerrero, K. Kalif, and R. T. Saba. 2004. Nitrogen and phosphorus limitation of biomass growth in a tropical secondary forest. *Ecological Applications* 14:S150-S163. **(13)**

Fisher, J. B., Y. Malhi, I. C. Torres, D. B. Metcalfe, M. J. van de Weg, P. Meir, J. E. Silva-Espejo, and W. H. Huasco. 2013. Nutrient limitation in rainforests and cloud forests along a 3,000-m elevation gradient in the Peruvian Andes. *Oecologia* 172:889-902. **(5)**

- Gehring, C., M. Denich, M. Kanashiro, and P. L. G. Vlek. 1999. Response of secondary vegetation in Eastern Amazonia to relaxed nutrient availability constraints. *Biogeochemistry* 45:223-241. **(9)**
- Kaspari, M., M. N. Garcia, K. E. Harms, M. Santana, S. J. Wright, and J. B. Yavitt. 2008. Multiple nutrients limit litterfall and decomposition in a tropical forest. *Ecology Letters* 11:35-43. **(15)**
- Li, Y. Q., M. Xu, and X. M. Zou. 2006. Effects of nutrient additions on ecosystem carbon cycle in a Puerto Rican tropical wet forest. *Global Change Biology* 12:284-293. **(2)**
- Markewitz, D., R. D. Figueiredo, C. J. R. de Carvalho, and E. A. Davidson. 2012. Soil and tree response to P fertilization in a secondary tropical forest supported by an Oxisol. *Biology and Fertility of Soils* 48:665-678. **(14)**
- Mayor, J. R., S. Joseph Wright, and B. L. Turner. 2014. Species-specific responses of foliar nutrients to long-term nitrogen and phosphorus additions in a lowland tropical forest. *Journal of Ecology* 102:36-44. **(16)**
- Mirmanto, E., J. Proctor, J. Green, L. Nagy, and Suriantata. 1999. Effects of nitrogen and phosphorus fertilization in a lowland evergreen rainforest. *Philosophical Transactions of the Royal Society of London Series B-Biological Sciences* 354:1825-1829. **(4)**
- Newbery, D. M., G. B. Chuyong, J. J. Green, N. C. Songwe, F. Tchuenteu, and L. Zimmermann. 2002. Does low phosphorus supply limit seedling establishment and tree growth in groves of ectomycorrhizal trees in a central African rainforest? *New Phytologist* 156:297-311.
- Uhl, C. 1987. Factors controlling succession following slash-and-burn agriculture in Amazonia. *Journal of Ecology* 75:377-407. **(8)**
- Villagra, M., P. I. Campanello, L. Montti, and G. Goldstein. 2013. Removal of nutrient limitations in forest gaps enhances growth rate and resistance to cavitation in subtropical canopy tree species differing in shade tolerance. *Tree Physiology* 33:285-296. **(3)**

Walker, L. R., J. K. Zimmerman, D. J. Lodge, and S. GuzmanGrajales. 1996. An altitudinal comparison of growth and species composition in hurricane-damaged forests in Puerto Rico. *Journal of Ecology* 84:877-889. **(1)**

Wright, S. J., J. B. Yavitt, N. Wurzburger, B. L. Turner, E. V. J. Tanner, E. J. Sayer, L. S. Santiago, M. Kaspari, L. O. Hedin, K. E. Harms, M. N. Garcia, and M. D. Corre. 2011. Potassium, phosphorus, or nitrogen limit root allocation, tree growth, or litter production in a lowland tropical forest. *Ecology* 92:1616-1625. **(17)**

For Review Only



## Appendix S2

This appendix presents sample sizes and results of analyses of tree growth responses after 15 years of factorial nitrogen, phosphorus and potassium fertilization of an old-growth forest in central Panama. This appendix also presents a power analysis for the repeated measures analysis of variance.

For Review Only

Appendix S2: Table S1. Samples sizes for repeated measures ANOVA analyses of RGR. Initial diameter at breast height (DBH<sub>i</sub>) is between 10 and 24 mm for shrubs and saplings, 25 and 49 mm for small poles, 50 and 99 mm for large poles, 100 and 249 mm for small trees, and larger than 249 mm for large trees. The analysis included all RGR values.

		Shrubs and saplings	Small poles	Large poles	Small trees	Large trees
Totals	Individual trees	3,147	2,000	1,034	451	137
	RGR values	6,633	4,424	2,353	1,044	330
RGR values per plot	Minimum	30	21	11	3	1
	Maximum	116	72	41	23	7
	Mean	69.1	46.1	24.5	10.9	3.44

Appendix S2: Table S2. Repeated measures analyses of mean, plot-level RGR for five size classes. Entries are F values. Initial diameter at breast height (DBH<sub>i</sub>) is between 10 and 24 mm for shrubs and saplings, 25 and 49 mm for small poles, 50 and 99 mm for large poles, 100 and 249 mm for small trees, and larger than 249 mm for large trees.

Between Subjects (or plots) Effects						
Effects <sup>@</sup>	Df	Shrubs & saplings	Small poles	Large poles	Small trees	Large trees
N	1	0.07	0.19	0.20	1.43	0.06
P	1	0.33	1.27	0.59	0.00	1.59
K	1	1.64	0.01	0.11	1.02	4.12 <sup>°</sup>
N*P	1	3.42	0.75	0.00	0.06	0.05
N*K	1	2.25	2.62	0.89	0.00	1.35
P*K	1	0.07	0.03	0.16	0.29	1.32
Repl	3	2.58	2.12	2.63	0.48	0.93
Repl:block	4	1.41	1.39	0.30	0.26	1.24
Residuals	18					
Within Subjects (or plots) Effects						
Intrvl	2	5.54**	1.48	0.18	0.63	1.11
N*Intrvl	2	1.34	1.60	0.96	2.09	0.38
P*Intrvl	2	0.55	0.19	0.18	1.09	1.06
K*Intrvl	2	0.02	0.16	0.30	1.00	0.88
N*P*Intrvl	2	1.65	1.05	0.50	1.30	0.31
N*K*Intrvl	2	0.12	0.14	0.05	1.45	1.15
P*K*Intrvl	2	0.95	2.10	1.53	0.72	0.46
Repl*Intrvl	6	1.01	0.44	0.43	1.93	1.19
Repl:block*Intrvl	8	1.11	0.60	0.36	0.39	2.17
Residuals	36					

<sup>@</sup> Repl refers to the four replicates of the factorial NPK experiment. Intrvl refers to the three census intervals. <sup>°</sup> p = 0.057; \*\* p < 0.01

Appendix S2: Table S3. Linear mixed effects analysis of RGR, with initial DBH as a covariate.

Entries are fixed effects and  $\Delta$ AIC values. All models include random effects for census interval, individual within species, and plot within block and treatment.

Fixed Effects <sup>@</sup>	$\Delta$ AIC
Null (no fixed effects)	183
$N * P * DBH_i + N * K * DBH_i + P * K * DBH_i$	194
$N * P + N * K * DBH_i + P * K * DBH_i$	172
$N * P + N * DBH_i + N * K + P * K * DBH_i$	150
$N * P + N * DBH_i + N * K + P * DBH_i + K * DBH_i + P * K$	129
$N * P + N * DBH_i + N * K + P * DBH_i + P * K$	105
$N * P + N * DBH_i + N * K + P * DBH_i$	94
$N * P + N * K + P * DBH_i$	71
$N * K + P * DBH_i$	62
$N + K + P * DBH_i$	53
$K + P * DBH_i$	40
$P * DBH_i$	28
$P + DBH_i$	12
$DBH_i$	0

@ Models include main effects and lower level interactions for each variable included in higher level interactions.

Appendix S2: Table S4. Samples sizes for the linear mixed effects analyses involving species-level phosphorus affinity. We excluded species with < 4 individuals with added P or without added P. Therefore, sample sizes are smaller than for the repeated measures ANOVAs (Appendix S2: Table 1). Initial diameter at breast height (DBH<sub>i</sub>) is between 10 and 24 mm for shrubs and saplings, 25 and 49 mm for small poles, 50 and 99 mm for large poles and 100 and 249 mm for small trees.

		Shrubs and saplings	Small poles	Large poles	Small trees
Total	Individual trees	2,754	1,627	650	313
	RGR values	5,851	3,631	1,524	747
RGR values per plot	Minimum	25	13	5	0 <sup>@</sup>
	Maximum	98	60	26	14
	Mean	61.0	37.8	15.9	7.8

<sup>@</sup> One plot lacked small trees with known species-level P affinity and was excluded from the analysis.

Appendix S2: Table S5. Linear mixed effects analyses of RGR, with species-level phosphorus affinity as a covariate. Entries are fixed effects and  $\Delta$ AIC values. All models included random effects for census interval, individual within species, and plot within block and treatment. Initial diameter at breast height (DBH<sub>i</sub>) is between 10 and 24 mm for shrubs and saplings, 25 and 49 mm for small poles, 50 and 99 mm for large poles and 100 and 249 mm for small trees.

Fixed Effects <sup>@</sup>	$\Delta$ AIC			
	Shrubs and saplings	Small poles	Larger poles	Small trees
Null (no fixed effects)	53	0	0	0
P affinity	66	13	12	10
P*P affinity	85	37	37	24
INT	0	10	25	27
INT + P affinity	13	23	38	38
INT + P*P affinity	35	47	63	62

<sup>@</sup> Models include main effects for each variable included in an interaction term.

## Appendix S2: Figure S1

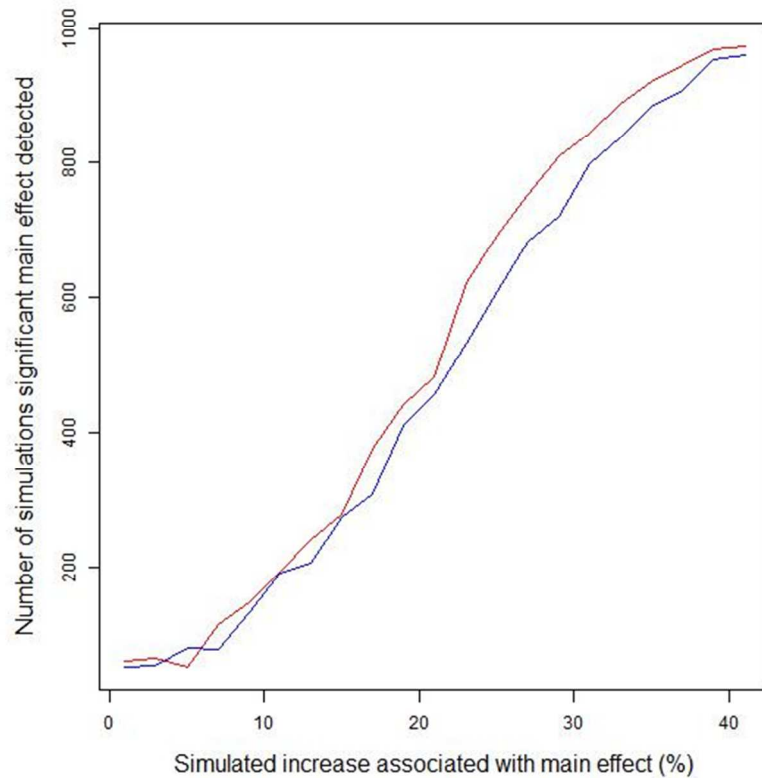


Figure S1. Power analysis for our factorial, incomplete blocks experimental design. The power analysis simulated a percentage increase for one nutrient main effect. Simulated values for plots where the nutrient was not added were random draws from normal distributions with the mean and standard deviation (SD) observed for plots where the nutrient was not added. Simulated values for plots where the nutrient was added were random draws from normal distributions with means increased by 1%, 3%, ... 41% and the same observed SDs. Means and SDs were calculated over all relevant plots (blue line) or over the relevant plots in each block and replicate (red line).

## Metadata S1. Metadata for the data file Data S1.

Variable	Definitions
Paper	An integer between 1 and 18. The key to references for the published articles can be found in Appendix S1: Table S4.
Age	OG or SF for old-growth or secondary forest, respectively.
Yrs	NA for Age=="OG"; forest age in years for Age=="SF"
Fertilizer	The nutrient treatment compared with the control. Values are 'complete', 'N', 'NP', 'NPK' and 'P'. Just 'N' and 'P' are used in the formal meta-analysis.
Response	The four response types are:
	1. foliar nutrient concentrations (values 'leaf.N' and 'leaf.P')
	2. fine litter nutrient concentrations (values 'litter.N' and 'litter.P')
	3. fine litter production (values 'litter.production' or 'leaf.litter.production')
	4. tree or biomass growth rates (values 'RGR.saplings', 'RGR.poles', 'RGR.small.trees', 'RGR.large.trees', 'RGR.alltrees', 'biomass.total' and 'biomass.woody')
units	Units provided by author.
Genus	NA for community-level responses. Genus name for species-level responses.
Species	NA for community-level responses. Species name for species-level responses.
m1	Plot-level mean for the control treatment.
sd1	Plot-level standard deviation for the control treatment (NA for missing values).
n1	Number of control plots.
m2	Plot-level mean for the fertilizer treatment (see Fertilizer above).
sd2	Plot-level standard deviation for the fertilizer treatment (see Fertilizer above) (NA



	for missing values).
n2	Number of fertilizer plots (see Fertilizer above).
Notes	The figure or table from which m1 through n2 were taken. Standard errors were converted to standard deviations.

## Notes:

1. The following articles lacked standard deviations for the responses in parentheses: Mirmanto et al. (1999) (RGR), Gehring et al. (1999) (species-level RGR), Newbery et al. (2002) (all responses), Davidson et al. (2004) (foliar N and P concentrations) and Chou et al. (2017) (all responses).
2. Gehring et al. (1999) combined a complete fertilizer treatment with minus single nutrient treatments. M1, sd1 and n1 refer to the minus N or minus P treatment. M2, sd2 and n2 refer to the complete fertilizer treatment.
3. For our factorial NPK experiment, nutrient interactions were insignificant, and we calculated m1, sd1, n1, m2, sd2 and n2 for all 16 plots fertilized with N (or P) and for all 16 plots NOT fertilized with N (or P).
4. For our factorial NPK experiment, Mayor et al. (2014) excluded the +K plots, leaving a factorial NP experiment. Therefore, n1 and n2 equal 8, not 16.