1	Lianas Have a Seasonal Growth Advantage Over Co-occurring Trees
2	
3	Stefan A. Schnitzer ^{1,2*} and Geertje M.F. van der Heijden ³
4	
5	¹ Marquette University, Department of Biological Sciences, PO Box 1881, Milwaukee, WI 53201, USA
6	² Smithsonian Tropical Research Institute, Apartado Postal 0843-03092, Balboa, República de Panamá
7	³ School of Geography, University of Nottingham, Nottingham, NG7 2RD, UK
8	
9	* Corresponding author: S1@mu.edu
10	
11	Running Head: Lianas Have a Seasonal Growth Advantage

12 **Abstract:** The seasonal growth advantage hypothesis posits that plant species that grow well 13 during seasonal drought will increase in abundance in forests with increasing seasonality of 14 rainfall both in absolute numbers and also relative to co-occurring plant species that grow poorly 15 during seasonal drought. That is, seasonal drought will give some plant species a growth 16 advantage that they lack in aseasonal forests, thus allowing them attain higher abundance. For 17 tropical forest plants, the seasonal growth advantage hypothesis may explain the distribution of 18 drought-adapted species across large-scale gradients of rainfall and seasonality. We tested the 19 seasonal growth advantage hypothesis with lianas and trees in a seasonal tropical forest in central 20 Panama. We measured the dry-season and wet-season diameter growth of 1117 canopy trees and 21 648 canopy lianas from 2011-2016. We also evaluated how lianas and trees responded to the 22 2015-2016 el Niño, which was the third strongest el Niño drought on record in Panama. We 23 found that liana growth rate was considerably higher during the dry-season months than the wet-24 season months in each of the five years. Lianas achieved a full half of their annual growth during 25 the 4-month dry season. By contrast, trees grew far more during the wet season; they realized 26 only a quarter of their annual growth during the dry season. During the strong 2015-2016 el Niño 27 dry season, trees essentially stopped growing, whereas lianas grew unimpeded and as well as 28 during any of the previous four dry seasons. Our findings support the hypothesis that seasonal 29 growth gives lianas a decided growth advantage over trees in seasonal forests compared to 30 aseasonal forests, and may explain why lianas peak in both absolute and relative abundance in 31 highly seasonal tropical forests. Furthermore, the ability of lianas to grow during a strong el Niño 32 drought suggests that lianas will benefit from the predicted increasing drought severity, whereas 33 trees will suffer, and thus lianas are predicted to increase in relative abundance in seasonal 34 tropical forests.

35 Keywords: El niño, Lianas, Seasonal growth advantage, Tropical forests, Plant distribution

36 INTRODUCTION

37 Determining the mechanisms responsible for the abundance and distribution of organisms is 38 one of the central goals in ecology (Krebs 1972, Brown 1984). Within the tropics, the abundance 39 (density) of most plant groups (e.g., trees, palms, herbs, and epiphytes) tends to increase with 40 increasing mean annual precipitation (Gentry 1991, Schnitzer 2005). By contrast, the abundance 41 of lianas, a common tropical plant group that is both taxonomically and functionally diverse 42 (Schnitzer & Bongers 2002, Wyka et al. 2013, Gianoli 2015, Schnitzer et al. 2015), deviates 43 from this common trend. Specifically, the density of lianas in tropical forests, both in absolute 44 terms and relative to trees, increases with the strength of seasonal drought and decreases with 45 increasing mean annual precipitation and the availability of soil moisture (e.g., Schnitzer 2005, 46 DeWalt et al. 2010, 2015, Manzané-Pinzón et al. 2018).

47 The seasonal growth advantage hypothesis may explain the unique distribution of lianas and 48 other species that increase in abundance with the intensity of seasonal drought. The seasonal 49 growth advantage hypothesis states that species that grow best during seasonal drought, when 50 solar radiation is high (due to the absence of thick cloud cover) and water availability is low, will 51 realize more annual growth than those same species in aseasonal areas, where they lack a season 52 of high growth (Schnitzer 2005, 2015a, 2018). In interspecific comparisons, there is a tradeoff 53 between growth rate and survivorship (e.g., Wright et al. 2010); however, intraspecific variation 54 in growth rate is positively correlated with higher survivorship (e.g., Camac et al. 2018) and 55 presumably fecundity, which we hypothesize culminates in higher densities. For liana species, 56 the period of high dry-season growth in seasonal tropical forests (commonly ranging from two to 57 six months) is hypothesized to increase liana size and survival, ultimately resulting in higher 58 density (in both absolute terms and relative to co-occurring tree species that grow poorly during 59 seasonal drought) compared to aseasonal forests (Schnitzer 2005, 2015a, 2018). Even a slight

Page 4 of 30

60 seasonal growth advantage could, over decades, allow liana species to increase in relative 61 abundance with increasing forest seasonality. Thus, the seasonal growth advantage hypothesis 62 may explain the increase in liana density (both in absolute terms and relative to trees) in forests 63 with increasing seasonality across the tropics (Schnitzer 2005, 2018). 64 The seasonal growth advantage hypothesis has been tested primarily in studies that measured 65 the physiological responses of lianas and trees during wet and dry seasons. These studies 66 reported that lianas generally have better access to water, experience less water stress, have 67 higher levels of photosynthesis, and have greater water use efficiency and osmotic adjustment 68 than do co-occurring trees during the dry season relative to the wet season (Cai et al. 2009, Chen 69 et al. 2015, Maréchaux et al. 2017). For example, in a common garden study with six replicated 70 tree species and six replicated liana species in central Panama, Smith-Martin et al. (in review) 71 found that, compared to trees, lianas had 44% higher predawn leaf water potential, 61% higher 72 intrinsic water-use efficiency, and 28% higher photosynthesis in the dry season compared to the 73 wet season. In an examination of physiological traits of liana and tree saplings that were growing 74 along roadsides in wet and seasonal forests in Panama, van der Sande et al. (in press) reported 75 that trees had the expected tradeoff between hydraulic conductance and hydraulic safety, 76 whereas lianas did not, suggesting that lianas had the capacity to maintain high conductivity and 77 thus high growth rates while resisting cavitation. Collectively, these studies suggest that lianas 78 are better able to grow during the dry season than co-occurring trees, and thus are able to 79 capitalize on high dry season light availability. In addition, one study (Schnitzer 2005) measured 80 the height growth of liana and tree saplings in the understory during a wet and a dry season of a 81 seasonal forest in Panama and found that lianas grew proportionally more than trees during the 82 dry season than the wet season, supporting the hypothesis that lianas had a seasonal growth 83 advantage.

Page 5 of 30

Ecology

84	Previous studies, however, were limited in their ability to test the seasonal growth advantage
85	hypothesis for two main reasons. First, although studies on plant water status and the
86	physiological responses of lianas and trees to low water availability were consistent with
87	underlying physiological mechanisms that could confer a dry season growth advantage to lianas
88	over trees (e.g., Cai et al. 2009, Chen et al. 2015, Maréchaux et al. 2017, Smith-Martin et al.
89	unpublished), none of the physiology-oriented studies actually demonstrated that lianas had
90	higher growth during the dry season. Less negative water potential during the dry season (e.g.,
91	Cai et al. 2009, Chen et al. 2015, 2017) could indicate that lianas limit carbon assimilation during
92	the day to avoid water loss. Alternatively, to remain physiologically active during periods of low
93	water availability, it is possible that lianas allocate much of their extra carbon to non-structural
94	carbohydrates (NSC) rather than to growth, because NSC can help plants maintain a healthy
95	water balance during periods of water stress (Würth et al. 2005, Körner et al. 2015, Martinez-
96	Vilalta et al. 2016, de Baerdemaeker et al. 2017). If so, then higher dry-season photosynthesis
97	may not translate directly into higher growth. Second, in the one study that measured seasonal
98	growth of lianas and trees, the author measured height growth for juvenile plants in the forest
99	understory (Schnitzer 2005). However, the responses of juvenile understory plants may not
100	accurately reflect the growth of canopy individuals, which may be a stronger contributor to
101	population demographic rates. Further, many liana species grow like tree saplings in the
102	understory (Manzané-Pinzón et al. 2018), and thus sapling physiology and growth (e.g., van der
103	Sande et al. (in press) may not accurately represent the liana-tree comparison for canopy
104	individuals. Also, Schnitzer (2005) sampled liana and tree growth during one wet and one dry
105	season, and thus they could not remove the effects of that particular year on seasonal growth.
106	Therefore, while there is some evidence supporting the fundamental premise of the seasonal

107 growth advantage hypothesis, direct evidence that canopy lianas grow more than canopy trees 108 across multiple dry seasons relative to multiple wet seasons is lacking. 109 We tested the seasonal growth advantage hypothesis by comparing annual wet- and dry-110 season growth of canopy lianas and canopy trees over a consecutive 5-year period (2011-2016). 111 Each year we measured the seasonal diameter growth of 1117 large trees (\geq 10 cm diameter) and 112 648 large lianas (\geq 5 cm) in eight 80 x 80 m plots in central Panama (1765 total canopy plants). 113 During the final year of the study (2015–2016), Panama experienced the third-strongest el Niño 114 drought on record (S. Paton, Smithsonian Tropical Research Institute), and we predicted that the 115 intense drought would elicit an even stronger disparity in the ratio of dry to wet season growth 116 between lianas and trees, with trees suffering more during the dry season than lianas. This study 117 is the most comprehensive test of the seasonal growth advantage hypothesis to date, and the first 118 to use multi-year seasonal diameter growth of adult lianas and trees.

119

120 METHODS

121 We conducted the study on Gigante Peninsula, a 60-year old secondary forest that is part of 122 the Barro Colorado Natural Monument (BCNM) in the Republic of Panama. The forest on 123 Gigante Peninsula is classified as a semi-deciduous, seasonally moist forest (Leigh 1999). The 124 Gigante forest receives a mean annual rainfall of ~2600 mm and has a strong dry season from 125 January until May in which rainfall is less than 100 mm per month (Schnitzer & Carson 2010). 126 During the period of the experiment (from 2011 until 2016), annual rainfall on the BCNM varied 127 from 1807 to 3262 mm per year (S. Paton, Smithsonian Tropical Research Institute). The 2016 128 El Niño drought in year 5 of the experiment resulted in the third driest year in the 90-year 129 continuous record, and the BCNM received only \sim 1480 mm of rain during the wet season, 130 followed by a long and strong dry season (S. Paton, Smithsonian Tropical Research Institute).

Page 7 of 30

Ecology

131	In 2008, we established eight 80 x 80 m plots, which initially served as controls for an
132	ongoing liana removal experiment (e.g., Alvarez-Cansino et al. 2015, van der Heijden et al.
133	2015, Reid et al. 2015, Martinez-Izquierdo et al. 2016, Rodriguez-Ronderos et al. 2016, Garcia-
134	Leon et al. 2018). Because lianas were not experimentally removed and no other manipulations
135	were conducted in these plots, they were well suited to test the seasonal growth advantage
136	hypothesis. In each plot, we permanently tagged, mapped, measured the diameter (1.3 m along
137	the stem from the roots), and identified to species all trees and lianas ≥ 1 cm diameter within the
138	center 60 x 60 m portion of the plot. Plant surveys followed sampling protocols established by
139	Gerwing et al. (2006) and Schnitzer et al. (2008) for lianas and Condit et al. (1998) for trees.
140	For canopy lianas \geq 5 cm diameter and canopy trees \geq 10 cm diameter, which were the focus
141	of this study, we used a fabric diameter tape to precisely measure the stem diameter of each
142	individual 1.3 m along the stem from the roots (where we painted an orange mark on each stem
143	to facilitate repeated measurements) at the beginning and end of the wet and dry seasons. For the
144	canopy trees, we also installed manual dendrometer bands in mid-November 2010, prior to the
145	2011 dry season, and allowed the bands to settle on the trees for at least four months before
146	collecting measurements (follows Alvarez-Cansino et al. 2015). We used the dendrometer bands
147	to determine the seasonal diameter increment for trees; however, dendrometer bands did not
148	work well for the smaller liana stems, and thus seasonal diameter increment was based on
149	diameter tape measurements (van der Heijden et al. 2015). The patterns of tree diameter growth
150	were the same regardless of the measurement method (dendrometer band or diameter tape);
151	however, dendrometer measurements had lower variability than diameter tape measurements.
152	Tree and liana diameters were calculated seasonally each year from the beginning of the
153	2011 wet season (May, 2011) until the end of the 2016 dry season (May, 2016). Each year, we
154	started the wet season census in late April / early May, immediately after the dry season trade

155 winds had stopped and there was rain for five to seven consecutive days. In central Panama, the 156 beginning of the wet season is marked by a tapering of the trade winds and the onset of rains, 157 which correspond to the northward movement of the intertropical convergence zone in mid-158 April. We conducted a second census at the start of the dry season (and end of the wet season) in 159 late December / early January, after the dry season trade winds had gained strength and there was 160 no rainfall for five to seven consecutive days. In central Panama, the onset of the dry season can 161 occur abruptly, often in a single day, changing from cloudy and rainy to sunny and windy with 162 no rain. Thus, we were able to quantify dry season (January – May) and wet season (May – 163 January) diameter growth for canopy lianas and trees over five consecutive years (2011 - 2016).

164

165 *Data Analyses*

166 We calculated mean seasonal growth for lianas and trees as both a percentage of the initial 167 stem diameter (relative growth) and also as absolute diameter increase. Because the wet season 168 in Panama is twice as long as the dry season, we also annualized the data to compare a seasonal 169 growth rate for both the wet and dry seasons. We included individuals that were alive during the 170 entire 5-year census period to avoid aberrant growth rates attributed to dying or dead individuals. 171 To test whether seasonal growth patterns were driven primary by common species, we compared 172 the growth trends of the most common and the rare liana and tree species. For trees, there was a 173 total of 128 species, with 7 that we defined as common (n > 30 individuals) and 96 that we 174 defined as rare (n < 5 individuals). For lianas, there was a total of 54 species, with 4 that we 175 defined as common (n > 30 individuals) and 30 that we defined as rare (n < 5 individuals). On 176 nearby Barro Colorado Island, only 6.3% of the canopy trees are deciduous during the peak of 177 the dry season (Condit et al. 2000), indicating that relatively few of the canopy trees in this area

Page 9 of 30

Ecology

178 were deciduous, and none of the common tree or liana species in our study were deciduous179 (Croat 1978).

180 We tested for differences in the mean growth rates between seasons in each year of the study 181 for all measures of liana and tree growth (absolute, relative and annualized growth) using a 182 Monte Carlo bootstrapping approach. This method allowed us to compare liana and tree growth 183 between seasons in each year using a single analysis, without the need for post-hoc testing. The 184 Monte Carlo bootstrapping approach also allowed us to integrate measurement error into the 185 confidence intervals and the analyses (cf. van der Heijden et al, 2015). For this approach, we 186 varied the initial diameter of each tree in each plot at random using a normal distribution with a 187 standard deviation of 5%. Tree diameters in subsequent censuses were calculated by adding a 188 randomly selected value from the normal distribution of dendrometer increment measurements 189 with a standard deviation of 3% to the tree diameter of the previous census. For lianas, we varied 190 the diameter measurements in each census by adding a randomly value selected from a normal 191 distribution with a standard deviation of 5% (cf. van der Heijden et al, 2015). We used this 192 approach to calculate 100,000 realizations of mean liana and tree growth for each season and the 193 differences in growth between seasons. These data were then used to calculate the mean seasonal 194 growth for lianas and trees, as well as the mean difference in liana and tree growth between the 195 wet and dry season for each year of the study and their 90% and 95% confidence intervals. The 196 confidence intervals for the difference between mean wet and dry season growth for each of the 197 growth forms were used to determine whether these differences were significant for each year of study. Differences in tree or liana growth between seasons were considered significant or 198 199 marginally significant when the 95%-confidence interval (P<=0.05) or 90%-confidence interval 200 $(0.05 < P \le 0.10)$, respectively, did not overlap with zero. The actual and modeled mean growth 201 values per season were nearly identical.

202

203 **RESULTS**

204 Lianas grew as much or more during the 4-month dry season than they did during the entire 205 8-month wet season (Fig. 1a). Liana growth rate (controlling for the length of the season) was far 206 higher during the dry season months than the wet season months (Fig. 1c). By contrast, trees had 207 the opposite pattern. Trees realized the vast majority of their growth during the wet season (Fig. 208 1b), and tree growth rate was more than two-times higher during the wet season than the dry 209 season (Fig. 1d). The 2015-2016 El Niño dry season was particularly devastating for trees, and 210 they essentially stopped growing during this period (Figs. 1b, 1d). Lianas, however, maintained 211 their growth during the strong El Niño dry season, growing as well as the previous four dry 212 seasons (Figs. 1a, 1c).

213 For both lianas and trees, the seasonal growth patterns of the most common species were 214 similar to that of the least common species (Figs. 2 & 3). Therefore, the patterns of vigorous dry 215 season growth for lianas and vigorous wet season growth for trees were not driven solely by the 216 common species, but they was also shared, on average, by the rare species. For one common 217 liana species (Bauhinia guianensis), dry and wet season relative growth were similar (Fig. 2). 218 Nonetheless, the seasonal growth of *B. guianensis* was not weighted as heavily towards wet 219 season growth as was the majority of the tree species (Fig. 3). The seasonal growth responses of 220 lianas and trees were the same regardless of whether we examined growth relative to the initial 221 stem size (Figs. 1, 2 & 3) or absolute growth over time (AppendixS1: Figures S1, S2, & S3). 222

223 DISCUSSION

224 Evidence for the seasonal growth advantage hypothesis

Page 11 of 30

Ecology

225 Our data strongly support the hypothesis that lianas have a growth advantage in seasonal 226 forests. Canopy lianas achieved more than half of their annual growth during the 4-month dry 227 season, with the rest of their growth occurring during the remaining 8-month wet season. By 228 contrast, canopy trees grew two- to three-times more during the 8-month wet season than during 229 the 4-month dry season. The ability of lianas to maintain higher growth rates during the dry 230 season than during the wet season gives them approximately four months of relatively high 231 growth in this forest - a growth advantage that would be lacking in ever-wet forests. The 232 seasonal growth advantage for lianas is even greater relative to co-occurring trees, since trees 233 grew relatively poorly during the dry season and, instead, concentrated their growth during the 234 wet season. Presumably, lianas would benefit even more relative to co-occurring trees in forests 235 with an even stronger dry season, which was the case for the extremely dry El Niño dry season, 236 when lianas grew well and trees essentially stopped growing.

237 Over decades, high dry-season growth could result in greater annual liana growth and 238 survival, with a greater number of liana stems accumulating in seasonal forests compared to 239 aseasonal forests (Schnitzer 2005, 2015a, 2018). This phenomenon could explain why liana 240 density tends to increase in tropical forests with relatively high seasonality and low annual 241 rainfall and soil moisture availability (e.g., Schnitzer 2005, Swaine & Grace 2008, DeWalt et al. 242 2010, 2015, Manzané-Pinzón et al. 2018). By contrast, trees should be favored over lianas in 243 forests that lack a dry season, since trees grow well during rainy periods. Further, the ability of 244 canopy lianas to grow well even during a strong el Niño dry season, while canopy trees largely 245 stopped growing, indicates that stronger droughts, which are both now observed and predicted to 246 increase in the future (e.g., Fu et al. 2013), may further favor lianas over trees in seasonal forests. 247 Our findings were similar to those of Schnitzer (2005), who measured juvenile lianas and 248 trees (< 2 m tall) in a seasonal forest in Panama for one year. In that study, both lianas and trees

249 grew better during the wet season than the dry season, but lianas grew relatively more during the 250 dry season than the wet season compared to trees. In the current study, we found a far higher 251 growth rates for canopy lianas during the dry season than during the wet season, with canopy 252 trees displaying the opposite growth pattern. The slight disparity between the two studies may be 253 due to the focus on canopy lianas and trees in the current study versus juvenile plants in the 254 previous study. Another difference is the much longer duration of the current study (5 years 255 versus 1 year), along with the larger sample size in this study (1765 canopy lianas and trees 256 versus 384 juvenile lianas and trees in the previous study). Collectively, these studies indicate that both canopy and understory lianas have a seasonal growth advantage compared to co-257 258 occurring trees, thus explaining why liana density peaks in seasonal forests.

259

260 *Why lianas grow more than trees during season drought*

261 High seasonal growth for lianas may be due to their ability to capitalize on the high-light 262 conditions that are present during seasonal drought (Schnitzer 2005, 2018). During the dry 263 season, the lack of clouds results in intense solar radiation that is largely absent during the rest of 264 the year. For example, in Central Panama, light availability above the forest canopy can increase 265 50% from dry season to wet season (Wright & van Schaik 1994, Graham et al. 2003). Plants that 266 can manage water and avoid severe water stress can capitalize on high dry-season light 267 availability through increased photosynthesis and carbon fixation. Furthermore, lianas appear to 268 have a higher photosynthetic capacity than trees (Cai et al. 2009, Chen et al. 2015, Smith-Martin 269 et al. in review), and by maintaining healthy water status and maintaining high hydraulic 270 conductivity during the dry season (e.g., van der Sande et al. in press), lianas are particularly 271 well suited to take advantage of high dry-season light availability. By contrast, trees appear to 272 suffer more water stress (Smith-Martin et al. in review) and have a more conservative hydraulic

1:

Page 13 of 30

Ecology

conductivity strategy than lianas (van der Sande et al. in press), and thus trees may not be able to
capitalize as well as lianas on the high dry season light. The ability of lianas to capitalize on high
solar radiation while maintaining healthy water status may also explain their extremely high
abundance in such high-light areas as treefall gaps, forest edges, and young tropical forests
(reviewed by Schnitzer 2018).

278 The ability to maximize photosynthesis and carbon fixation with high light availability 279 requires some combination of access to sufficient quantities of water and the ability to use water 280 efficiently. Several studies have shown that lianas have access to water during the dry season, 281 which may allow them to maintain healthy water status during seasonal drought (e.g., Cai et al. 282 2009, Chen et al. 2015, 2017, Smith-Martin et al. in review). Lianas may be able to further 283 maintain healthy water status by minimizing the tradeoff between photosynthesis and water loss 284 (Schnitzer 2018). Smith-Martin et al. (in review) found that during the dry season in Panama, 285 lianas had 63% higher intrinsic water use efficiency than co-occurring trees. During the wet 286 season, however, water use efficiency between trees and lianas was similar. Also in Panama, van 287 der Sande et al. (in press) suggested that lianas, but not trees, could maintain high vascular 288 conductivity and presumably growth under dry conditions while resisting vascular cavitation. 289 Studies in SW China also reported that lianas used water and nitrogen more efficiently than did 290 trees during the dry season (e.g., Cai et al. 2009, Chen et al. 2015). Therefore, lianas appear to be 291 able to acquire and efficiently use soil moisture, which allows them to capitalize on the high-292 light environment of the dry season while simultaneously maintaining healthy water status.

293

294 *Potential alternative explanations*

It is possible that the seasonal growth advantage was driven more by a release from intense wet season tree competition rather than a dry season growth advantage. That is, vigorous canopy

297 tree growth during the wet season may have suppressed liana growth, and lianas may appear to 298 have a seasonal growth advantage because they are released from competition during the dry 299 season, when trees are largely dormant and some are deciduous. The available data, however, 300 appear to support the seasonal growth advantage hypothesis more than the competitive release 301 hypothesis. Lianas performed better than trees during the dry season even when individuals were 302 grown separately and there was no possibility for competitive release. For example, lianas and 303 trees that were grown separately in a common garden for five years performed similarly during 304 the wet season, but lianas performed better during the dry season in terms of water status, 305 photosynthesis, and water use efficiency (Smith-Martin et al. in review). Also, trees in the 306 common gardens grew far better when exposed to dry season irrigation (compared to non-307 irrigated controls), whereas lianas did not respond positively to dry season irrigation, suggesting 308 that trees, not lianas, suffered from low soil availability during the dry season (Smith-Martin et 309 al. in review). In terms of aboveground competition, liana foliage is typically deployed on top of 310 their tree hosts (e.g., Rodriguez et al. 2016); thus, by restricting our study to sun-exposed canopy 311 lianas and trees, we limited the effect of competition for light from canopy trees, and thus limited 312 the amount of competitive release that was possible. Canopy trees could possibly suppress 313 understory lianas during the wet season by decreasing light; however, understory lianas actually 314 grew more during the wet season than they did during the dry season, even though lianas grew 315 proportionally more than trees during the dry season than the wet season (Schnitzer 2005). 316 Therefore, the available data support the seasonal growth advantage hypothesis. Nonetheless, 317 little is known about the competitive effects of trees on lianas (Stewart & Schnitzer 2017), and 318 fully factorial experimental plant removal experiments would allow us to more definitively test 319 between these two alternative hypotheses.

1.

Page 15 of 30

Ecology

320 Our diameter growth estimates may have been influenced by the swelling or shrinking of 321 liana and tree stems during the seasons. The diameter of trees and presumably lianas can shrink 322 when stem storage tissues are dehydrated due to low soil moisture availability and when bark is 323 dehydrated due to low relative humidity (e.g., Borchert 1999, Stahl et al. 2010). For the census 324 that began at the beginning of the dry season, soil moisture was high after the previous eight 325 months of wet season rainfall (Reid et al. 2015), so stem storage tissues should not have been 326 dehydrated; however, there may have been bark shrinkage due to a drop in relative humidity 327 (Stahl et al. 2010). For the census that began at the beginning of the wet season, soil moisture 328 may have been low after the long dry season (Reid et al. 2015), so stem storage tissues could 329 have been dehydrated; however, stem shrinkage may have been minimized or absent because we 330 waited until we had received five to seven days of rainfall before we began the wet season 331 census. Furthermore, there may have been bark swelling due to the increase in relative humidity 332 after the start of the wet season. Whether seasonal stem changes not attributable to growth biased 333 our results would depend on the relative importance of stem storage tissue and bark swelling and 334 shrinking to stem size, which would give insight into whether we potentially underestimated or 335 overestimated seasonal stem growth for either lianas and trees. Importantly, there is little 336 evidence that lianas and trees differ systematically in stem storage tissues and bark 337 characteristics, and thus there is no evidence that they would shrink or swell differently during 338 the wet and dry seasons. Therefore, based on the available evidence, a likely explanation for our 339 findings is that lianas grew more than trees during seasonal drought.

340

341 *Does seasonal water partitioning explain liana and tree coexistence?*

342 Lianas and trees have coexisted in tropical forests for millions of years and the ability to

343 climb was an early innovation in terrestrial plants (Burnham 2015). However, the factors that

344 determine the relative abundance of lianas and trees, and whether liana and tree densities are 345 inherently stable, is the subject of recent investigation (Stewart & Schnitzer 2017, Visser et al. 346 2018a, 2018b, Muller-Landau & Pacala in press). One potential explanation is that liana and tree 347 fitness and demographic rates are controlled by factors other than liana-tree competition, and 348 thus a modest change in the abundance of one group does not necessarily reduce the abundance 349 of the other. However, lianas have strong negative effects on tree growth (Schnitzer et al. 2014, 350 van der Heijden et al. 2015, Toledo-Aceves 2015, Estrada-Villegas & Schnitzer 2018), 351 reproduction (Kainer et al. 2014, Garcia-Leon et al. 2018), and recruitment (Grauel & Putz 2004, 352 Schnitzer & Carson 2010), and thus it seems unlikely that lianas would have little effect on tree 353 demography (Visser et al. 2018a).

354 Another explanation for liana and tree coexistence is that lianas and trees have diverged (or 355 not fully converged) in their seasonal resource use, which theoretically could permit stable 356 coexistence. That is, the ability of lianas to grow more during the dry season while trees grow 357 more during the wet season may be a form of temporal resource partitioning (sensu Hutchinson 358 1961), which may explain long-term stable coexistence between lianas and trees. If lianas and 359 trees had completely overlapping resource requirements, the removal of lianas should result in a 360 strong response in trees during the dry season - the period when lianas grow most and thus 361 presumably compete the most. However, trees in the Gigante Peninsula forest did not experience 362 a greater competitive release after removing lianas from eight experimentally manipulated plots 363 during the dry season compared to the wet season (van der Heijden et al. in review). Nor did lianas appear to have a stronger dry-season effect on trees in a forest fragment in Brazil 364 365 (Venegas-Gonzalez et al. in review). These recent experimental findings support the possibility 366 of ecological divergence between lianas and trees during their long history of coexistence, which 367 may allow them to coexist rather than for either growth-form to ultimately displace the other. In

368	effect, the temporal partitioning of resources between lianas and trees during the year may
369	represent a ghost of competition past (sensu Connell 1980).
370	
371	Implications for increasing liana abundance in tropical forests
372	The ability of lianas to grow well in dry conditions with high evaporative demand (e.g.,
373	seasonal droughts, forest gaps, regenerating forests, and highly seasonal forests; Schnitzer 2018)
374	may explain documented increases in lianas in neotropical forests (Phillips et al. 2002, Schnitzer
375	& Bongers 2011, Schnitzer 2015b). Many tropical areas are now experiencing more intense
376	droughts with increasing global climate change (Lewis et al. 2011, Fu et al. 2013). Our findings
377	suggest that more intense droughts would favor liana growth over that of trees, especially in
378	seasonal tropical forests, which would presumably increase liana survival and fecundity,
379	ultimately resulting in higher liana densities relative to trees. Furthermore, extended droughts
380	increase tree mortality, which results in greater forest disturbance and increases the availability
381	of high light areas that favor liana proliferation (Schnitzer et al. 2000, Dalling et al. 2012, Ledo
382	& Schnitzer 2014, Schnitzer 2018). While there may be other factors that favor lianas in a
383	changing environment, including elevated atmospheric CO ₂ (Phillips et al. 2002; but see Marvin
384	et al. 2015) and nitrogen deposition (Schnitzer & Bongers 2011, Schnitzer 2015b; but see
385	Pasquini et al. 2015), increasing drought and disturbance may be co-conspirators that are
386	responsible for increasing liana abundance in many tropical forests.
387	

388 Summary

This is the first study to demonstrate unequivocally that canopy lianas gain a large proportion of their annual growth during the dry season; whereas canopy trees grow mostly during the wet season. Vigorous dry-season growth may result in greater survival and fecundity, allowing liana

392 stems to accumulate over time, thus explaining the relatively high abundance of lianas in 393 seasonal tropical forests compared to aseasonal forests. Furthermore, this study supports the 394 seasonal growth advantage hypothesis to explain the pan-tropical distribution of lianas, which 395 peak in abundance in highly seasonal tropical forests. The ability of lianas to grow well even 396 during a particularly strong el Niño dry season, when co-occurring trees essentially stopped 397 growing, indicates that the frequency and intensity of drought, which are predicted to increase 398 with global climate change, will favor lianas over trees and may explain observed increases in 399 liana abundance in tropical forests.

400

401 Acknowledgements

We thank Truman Young and two anonymous reviewers for helpful comments on this
manuscript. We also thank Maria García-León, who coordinated the logistics, as well as Boris
Bernal, Salomé Pérez, Abelino Valdés, Oldemar Valdés, and Severino Valdés, who provided
valuable assistance in the field. Guadalupe Alvarado entered data from this project from 2011
until 2018. Financial support was provided by NSF-DEB 0845071, NSF-DEB 1019436, NSFDEB 1822473, and NSF-IOS 1558093. Logistical support was provided by Marquette University
and the Smithsonian Tropical Research Institute.

409	Literature Cited
410	Alvarez-Cansino L, Schnitzer SA, Reid J, Powers JS. 2015. Liana competition with tropical trees
411	varies with seasonal rainfall and soil moisture, but not tree species identity. Ecology 96: 39-45.
412	Andrade JL, Meinzer FC, Goldstein G, Schnitzer SA. 2005. Water uptake and transport in lianas
413	and co-occurring trees of a seasonally dry tropical forest. Trees-Structure & Function 19: 282-
414	289.
415	Asner GP, Martin RE. 2015. Canopy chemistry expresses the life-history strategies of lianas and
416	trees. Pages 299-308 in The Ecology of Lianas, Schnitzer, S.A., F. Bongers, R.J. Burnham, F.E.
417	Putz, editors. Wiley-Blackwell, Oxford.
418	Borchert R. 1999. Climatic periodicity, phenology, and cambium activity in tropical dry forest
419	trees. IAWA Journal 20: 239-247
420	Brown JH. 1984. On the relationship between abundance and distribution of species. The
421	American Naturalist 124: 255–279.
422	Burnham RJ. 2015. Climbing plants in the fossil record: Paleozoic to present. Pages 205-220 in
423	The Ecology of Lianas, Schnitzer, S.A., F. Bongers, R.J. Burnham, F.E. Putz, editors. Wiley-
424	Blackwell, Oxford.
425	Cai ZQ, Schnitzer SA, Bongers F. 2009. Seasonal differences in leaf-level physiology give lianas
426	a competitive advantage over trees in a tropical forest. Oecologia 161: 25-33.
427	Camac JS, Condit R, Fitzjohn RG, McCalman L, Steinberg D, Westoby M, Wright SJ, Falster DS.
428	2018. Partitioning mortality into growth-dependent and growth-independent hazards across 203
429	tropical tree species. Proceedings of the National Academy of Sciences 49: 12459-12464.
430	Chen Y-J, Cao K-F, Schnitzer SA, Fan Z-X, Zhang J-L, Bongers F. 2015. Water-use advantage of
431	lianas over trees in seasonal tropical forests. New Phytologist 205: 128-136.

1

- 432 Chen Y-J, Schnitzer SA, Zhang, Fan Z-X, Goldstein G, Tomlinson KW, Cao K-F, Zhang J-L.
- 433 2017. Stomatal regulation and efficient xylem water transport regulate diurnal water and carbon
- 434 balances of tropical lianas. *Functional Ecology* 31: 306-317.
- 435 Condit R. 1998. Tropical Forest Census Plots: Methods and Results from Barro Colorado Island,
- 436 Panama and a Comparison with Other Plots. Springer-Verlag, Berlin.
- 437 Condit R, Watts K, Bohlman SA, Pérez R, Foster RB, Hubbell SP. 2000. Quantifying the
- 438 deciduousness of tropical canopies under varying climates. *Journal of Vegetation Science* 11:
 439 649-658.
- 440 Connell JH. 1980. Diversity and the Coevolution of Competitors, or the Ghost of Competition
 441 Past. *Oikos* 35: 131-138.
- 442 Croat TB. 1978. The Flora of Barro Colorado Island. Stanford University Press, Stanford.
- 443 Dalling JW, Schnitzer SA, Baldeck C, Harms K, John R, Mangan SA, Lobo E, Yavitt JB, Hubbell
- 444 SP. 2012. Resource-based habitat associations in a neotropical liana community. Journal of
- 445 *Ecology* 100: 1174-1182.
- 446 De Baerdemaeker NJF, Salomón RL, De Roo L, Steppe K. 2017. Sugars from woody tissue
- 447 photosynthesis reduce xylem vulnerability to cavitation. *New Phytologist* 216: 720-727.
- 448 DeWalt SJ, Schnitzer SA, Añves Ñ-F, Bongers F, Burnham RJ et al. 2015. Biogeographical
- 449 patterns of liana abundance and diversity. Pages 131-1466 in The Ecology of Lianas, Schnitzer,
- 450 S.A., F. Bongers, R.J. Burnham, F.E. Putz, *editors*. Wiley-Blackwell, Oxford.
- 451 DeWalt SJ, Schnitzer SA, Chave J, Bongers F, Burnham RJ, Cai Z-Q, Chuyong G, Clark DB,
- 452 Ewango CEN, Gerwing JJ, Gortaire E, Hart T, Ibarra-Manríquez G, Ickes K, Kenfack D, Macía
- 453 MJ, Makana JR, Mascaro J, Martínez-Ramos M, Moses S, Muller-Landau HC, Parren MPE,
- 454 Parthasarathy N, Pérez-Salicrup DR, Putz FE, Romero-Saltos H, Thomas D. 2010. Annual

455	rainfall and seasonality predict pan-tropical patterns of liana density and basal area. Biotropica
456	42: 309-317.
457	Fu R, Yin L, Li W, Arias PA, Dickinson RE, Huang L, et al. 2013. Increased dry-season length
458	over southern Amazonia in recent decades and its implication for future climate projection.
459	Proceedings of the National. Academy of Sciences 110: 18110–18115.
460	Garcia-Leon M, Martinez-Izquierdo L, Powers JS, Schnitzer SA. 2018. Lianas reduce community-
461	level canopy tree reproduction in a Panamanian forest. Journal of Ecology 106: 737 - 745.
462	Gentry AH. 1991. The distribution and evolution of climbing plants. Pages 3-49 in F.E. Putz and
463	H.A. Mooney, eds. The Biology of Vines. Cambridge University Press, Cambridge.
464	Gianoli E. 2015. Evolutionary implications of the climbing habit in plants. In S. A. Schnitzer, F.
465	Bongers, R. J. Burnham and F. E. Putz (Eds.). Ecology of Lianas, pp. 239-250. John Wiley &
466	Sons, Oxford.
467	Graham EA., Mulkey SS, Kitajima K, Phillips NG, Wright SJ. 2003. Cloud cover limits net CO2
468	uptake and growth of a rainforest tree during tropical rainy seasons. Proceedings of the National
469	Academy of Sciences 100:572-576.
470	Grauel WT and Putz FE. 2004. Effects of lianas on growth and regeneration of Prioria copaifera
471	in Darien, Panama. Forest Ecology and Management 190: 99-108.
472	Hutchinson GE. 1961. The paradox of the plankton. The American Naturalist 95: 137-145.

- 473 Kainer KA, Wadt LHO, Staudhammer CL. 2014. Testing a silvicultural recommendation: Brazil
- 474 nut responses 10 years after liana cutting. *Journal of Applied Ecology* 51: 655-663.
- 475 Körner C. 2015. Paradigm shift in plant growth control. *Current Opinion in Plant Biology* 25:
 476 107–114.

- 477 Krebs CJ. 1972. Ecology: the experimental analysis of distribution and abundance. Harper & Row,
 478 New York.
- 479 Ledo A, Schnitzer SA. 2014. Disturbance and clonal reproduction determine liana distribution and
 480 maintain liana diversity in a tropical forest. *Ecology* 95: 2169-2178.
- 481 Leigh EG. 1999. Tropical Forest Ecology: A View from Barro Colorado Island. Oxford University
 482 Press, Oxford.
- Lewis SL, Brando PM, Phillips OL, van der Heijden GMF, Nepstad D. 2011. The 2010 Amazon
 drought. *Science* 331: 554.
- 485 Manzané-Pinzón E, Goldstein G, Schnitzer SA. 2018. Does soil moisture availability explain liana
- 486 seedling distribution across a tropical rainfall gradient. *Biotropica* 50: 215-224.
- 487 Maréchaux I, Bartlett MK, Iribar A, Sack L, Chave J. 2017. Stronger seasonal adjustment in leaf
- turgor loss point in lianas than trees in an Amazonian forest. *Biology Letters* 13: 20160819.
- 489 Martinez-Vilalta J, Sala A, Asensio D, Galiano, L, Hoch G, Palacio S, Piper FI, Lloret F. 2016.
- 490 Dynamics of non-structural carbohydrates in terrestrial plants: a global synthesis. *Ecological*
- 491 *Monographs* 86: 495-516.
- 492 Martinez-Izquierdo L, Garcia-Leon MM, Powers JS, Schnitzer SA. 2016. Lianas suppress
- 493 seedling growth and survival of 14 tree species in a Panamanian tropical forest. *Ecology* 97:
- 494 215-224.
- 495 Muller-Landau HC, Pacala SW. 2018. What determines the abundance of lianas? In Unsolved
- 496 Problems in Ecology, edited by A. Dobson, R. Holt, and D. Tilman. In press.
- 497 Pasquini SC, Wright SJ, Santiago LS. 2015. Lianas always outperform tree seedlings regardless of
- 498 soil nutrients: results from a long-term fertilization experiment. *Ecology* 96: 1866-1876.

499	Phillips OL, Vasquez Martinez R, Arroyo L, Baker TR, Killeen T, Lewis SL, et al. 2002.
500	Increasing dominance of large lianas in Amazonian forests. Nature 418: 770–774.
501	Reid JP, Schnitzer SA, Powers JS. 2015. Soil moisture variation after liana removal in a
502	seasonally moist, lowland tropical forest. PLoS One DOI 10.1371/journal.pone.0141891
503	Rodriguez-Ronderos ME, Bohrer G, Sanchez-Azofeifa A, Powers JS, Schnitzer SA. 2016.
504	Contribution of lianas to plant area index and structure in a Panamanian forest. <i>Ecology</i> 97:
505	3271-3277.
506	Santiago LS, Wright SJ. 2007. Leaf functional traits of tropical forest plants in relation to growth
507	form. <i>Functional Ecology</i> 21: 19–27.
508	Schnitzer SA. 2005. A mechanistic explanation for global patterns of liana abundance and
509	distribution. The American Naturalist 166: 262-276.
510	Schnitzer SA. 2015a. The ecology of lianas in forest ecosystems. Pages 185-197 in: Handbook of
511	Ecology, Peh, K., R. Corlett, Y. Bergeron, editors. Routledge Publishing, New York, NY.
512	Schnitzer SA. 2015b. Increasing liana abundance and biomass in neotropical forests: causes and
513	consequences. Pages: 451-464 in: Ecology of Lianas, Schnitzer, S.A., F. Bongers, R.J.
514	Burnham, F.E. Putz, editors. Wiley-Blackwell Publishing, Oxford.
515	Schnitzer SA 2018. Testing ecological theory with lianas. New Phytologist 220: 366-380.
516	Schnitzer SA, Bongers F. 2002. The ecology of lianas and their role in forests. Trends in Ecology
517	& Evolution 17: 223–230.
518	Schnitzer SA, Bongers F. 2011. Increasing liana abundance and biomass in tropical forests:
519	emerging patterns and putative mechanisms. Ecology Letters 14: 397-406.
520	Schnitzer SA, Carson WP. 2010. Lianas suppress tree regeneration and diversity in treefall gaps.
521	Ecology Letters 13: 849–857.

- 522 Schnitzer SA, Dalling JW and Carson WP. 2000. The impact of lianas on tree regeneration in
- 523 tropical forest canopy gaps: Evidence for an alternative pathway of gap-phase regeneration.
- 524 *Journal of Ecology* 88: 655-666.
- 525 Schnitzer SA, Mangan SA, Hubbell SP. 2015. The lianas of Barro Colorado Island, Panama.
- 526 Pages: 76-90 in: Ecology of Lianas, Schnitzer, S.A., F. Bongers, R.J. Burnham, F.E. Putz,
- 527 *editors*. Wiley-Blackwell Publishing, Oxford.
- Schnitzer SA, Rutishauser S, Aguilar S. 2008a. Supplemental protocol for liana censuses. *Forest Ecology and Management* 255: 1044-1049.
- 530 Smith-Martin, CM, Bastos CL, Lopez OR, Powers JS, Schnitzer SA. In review. Effects of dry-
- season irrigation on leaf physiology and biomass allocation in tropical lianas and trees. *Ecology*.
- 532 Stahl C, Burban B, Bompy F, Jolin ZB, Sermage J, Bonal D. 2010. Seasonal variation in
- 533 atmospheric relative humidity contributes to explaining seasonal variation in trunk
- 534 circumference of tropical rain-forest trees in French Guiana. *Journal of Tropical Ecology* 26:

535 **393-405**.

- 536 Stewart TE, Schnitzer SA. 2017. Blurred lines between competition and parasitism. *Biotropica* 49:
 537 433-438.
- Swaine MD, Grace J. 2007. Lianas may be favoured by low rainfall: evidence from Ghana. *Plant Ecology* 192: 271-276.
- 540 Toledo-Aceves, T. 2015. Above- and belowground competition between lianas and trees. Pages
- 541 147–163 in Ecology of Lianas, Schnitzer SA, Bongers F, Burnham RJ, Putz FE, editors. Wiley-
- 542 Blackwell Publishing, Oxford.
- van der Heijden GMF, Powers SJ, Schnitzer SA. 2015. Lianas reduce carbon accumulation in
- tropical forests. *Proceedings of the National Academy of Sciences* 112: 13267-13271.

545	van der Heijden, GMF, Powers JS, Schnitzer SA. In review. No seasonal differences in liana effect
546	on forest-level tree biomass growth in a liana removal experiment in Panama. Journal of
547	Ecology.
548	van der Sande MT, Poorter L, Schnitzer SA, Engelbrecht BMJ, Markesteijn L. In review. The
549	hydraulic efficiency-safety trade-off differs between lianas and trees. Ecology.
550	Visser MD, Schnitzer SA, Wright SJ, Muller-Landau HC, Jongejans E, Comita LS de Kroon H,
551	Condit R, Hubbell SP. 2018a. Tree species vary widely in their tolerance for liana infestation: A
552	case study of differential host response to generalist parasites. Journal of Ecology 106: 784-794.
553	Visser MD, Muller-Landau HC, Schnitzer SA, de Kroon H, Jongejans E, Wright SJ. 2018b. A
554	host-parasite model explains variation in liana infestation among co-occurring tree species.
555	Journal of Ecology in press.
556	Wright SJ, van Schaik CP. 1994. Light and the phenology of tropical trees. The American
557	Naturalist 143: 192-199.
558	Wright SJ, Kitajima K, Kraft NJB, Reich PB, Wright IJ, Bunker DE, Condit R, Dalling JW,
559	Davies SJ, Diaz S, Engelbrecht BMJ, Harms KE, Hubbell SP, Marks, Ruiz-Jaen MC, Salvador
560	CM, Zanne AE. 1999. Functional Traits and the growth-mortality trade-off in tropical trees.
561	<i>Ecology</i> 80: 1632-1647.
562	Würth, M. K. R., S. Peláez-Riedl, S. J. Wright, and C. Körner. 2005. Non-structural carbohydrate
563	pools in a tropical forest. Oecologia 143: 11–24.
564	Wyka TP, Oleksyn J, Karolewsk P, Schnitzer SA. 2013. Phenotypic correlates of the lianescent
565	growth form - a review. Annals of Botany. 112: 1667-1681.

566 Figures

567 Figure 1. Mean bootstrapped relative growth (based on initial size) and annualized relative growth 568 for lianas (first column, N=648 individuals and 54 species) and trees (second column, N=1117 569 individuals and 128 species) over a five-year period (2011-2016) on Gigante Peninsula in central 570 Panama. Error bars represent 95% confidence intervals based on 100,000 bootstrap iterations; ** 571 indicates P < 0.05, * indicates 0.05 >= P <= 0.10. 572 573 Figure 2. Mean bootstrapped relative growth (based on initial size) and annualized relative growth 574 for common and rare liana species over a five-year period (2011-2016) on Gigante Peninsula in 575 central Panama. Common species were those with more than 30 replicate individuals among the 8 576 plots. Rare species were those with fewer than 5 individuals, and each individual was used as a 577 replicate to calculate a mean response. There were 4 common and 30 rare liana species. Error bars

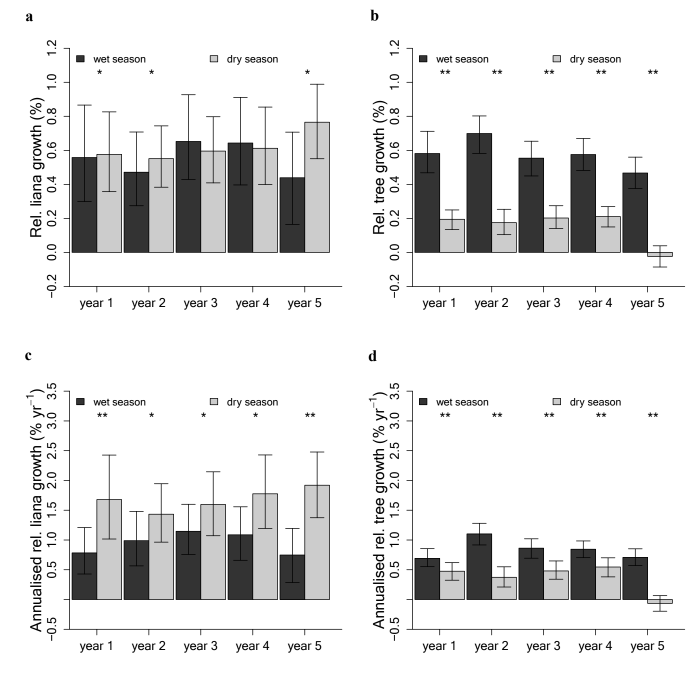
578 indicate 95% confidence intervals based on 100,000 bootstrap iterations; ** indicates P<0.05, *

579 indicates $0.05 \ge P \le 0.10$.

580

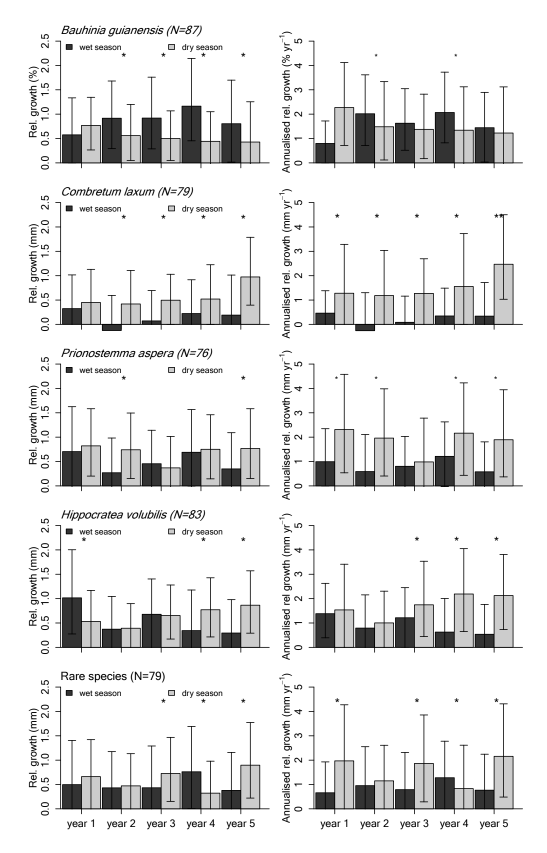
Figure 3. Mean bootstrapped relative growth (based on initial size) and annualized relative growth for common and rare tree species over a five-year period (2011-2016) on Gigante Peninsula in central Panama. Common species were those with more than 30 replicate individuals among the 8 plots. Rare species were those with fewer than 5 individuals, and each individual was used as a replicate to calculate a mean response. There were 7 common and 96 rare tree species. Error bars indicate 95% confidence intervals based on 100,000 bootstrap iterations; ****** indicates P<0.05, ***** indicates 0.05>= P <= 0.10.

588 Figure 1



589

591 Figure 2



593 Figure 3

