



Competition influences tree growth, but not mortality, across environmental gradients in Amazonia and tropical Africa

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Abstract. Competition among trees is an important driver of community structure and dynamics in tropical forests. Neighboring trees may impact an individual tree's growth rate and probability of mortality, but large-scale geographic and environmental variation in these competitive effects has yet to be evaluated across the tropical forest biome. We quantified effects of competition on tree-level basal area growth and mortality for trees ≥ 10 -cm diameter across 151 ~ 1 -ha plots in mature tropical forests in Amazonia and tropical Africa by developing nonlinear models that accounted for wood density, tree size, and neighborhood crowding. Using these models, we assessed how water availability (i.e., climatic water deficit) and soil fertility influenced the predicted plot-level strength of competition (i.e., the extent to which growth is reduced, or mortality is increased, by competition across all individual trees). On both continents, tree basal area growth decreased with wood density and increased with tree size. Growth decreased with neighborhood crowding, which suggests that competition is important. Tree mortality decreased with wood density and generally increased with tree size, but was apparently unaffected by neighborhood crowding. Across plots, variation in the plot-level strength of competition was most strongly related to plot basal area (i.e., the sum of the basal area of all trees in a plot), with greater reductions in growth occurring in forests with high basal area, but in Amazonia, the strength of competition also varied with plot-level wood density. In Amazonia, the strength of competition increased with water availability because of the greater basal area of wetter forests, but was only weakly related to soil fertility. In Africa, competition was weakly related to soil fertility and invariant across the shorter water availability gradient. Overall, our results suggest that competition influences the structure and dynamics of tropical forests primarily through effects on individual tree growth rather than mortality and that the strength of competition largely depends on environment-mediated variation in basal area.

Key words: climatic water deficit; competition; forest dynamics; mortality; neighborhood effects; soil fertility; trait-based models; tree growth; tropical forest; wood density.

INTRODUCTION

Competition is an important driver of community structure and dynamics in forests worldwide (Kunstler et al. 2016), particularly in closed-canopy forests such as mature, undisturbed tropical forests, where low light levels under the canopy typically limit tree growth. Generally, competition with neighboring trees is expected to decrease growth and increase the probability of mortality of individual tropical trees (Uriarte et al. 2004, Lasky et al. 2015). However, effects of competition on growth and mortality of individual trees have only been quantified within single tropical forest sites to date (e.g., Uriarte et al. 2004, Baribault et al. 2012). Whether strong effects of competition on demographic rates are pervasive, and whether they vary across environmental gradients in the tropics, remains unresolved.

Better knowledge of the effects of competition on tropical tree growth and mortality, and the geographic variation thereof, is essential for enhancing understanding of the global terrestrial carbon balance. Mature tropical forests have increased in biomass over recent decades (Lewis et al. 2009), and those in Amazonia have become more dynamic (McDowell et al. 2018). Mortality rates have a key role in controlling biomass in tropical forests (Johnson et al. 2016), as increases in mortality over time are influencing the carbon balance of Amazon forests (Brienen et al. 2015). Changes in the average strength of competition in forests might be one of the driving factors of such dynamic changes, because increased biomass (i.e., increased neighborhood crowding) leads to enhanced competition, with expected impacts in turn in decreased growth and increased mortality. More generally, the underlying causes of tree mortality in the tropics are still actively debated (e.g., McDowell et al. 2018), and quantifying their effects on the terrestrial carbon balance is a key challenge for ecologists and global change scientists. In addition to

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mortality that results from competition, trees may die from a range of other processes, including hydraulic failure in response to drought (large trees in particular; Phillips et al. 2010, Bennett et al. 2015, Rowland et al. 2015), from senescence (although effects are weak; Mencuccini et al. 2005), and from large-scale wind disturbance (Espírito-Santo et al. 2014), but which process(es) dominate(s) remains poorly understood.

Environmental conditions vary considerably across tropical forest sites, and this variation is known to influence forest structure and dynamics strongly. Across the Amazon basin, for example, water availability generally decreases from north to south, and soil fertility increases from east to west (ter Steege et al. 2006). Drier forests generally have a lower stature, lower aboveground biomass and basal area, and a more open canopy than wet forests (Quesada et al. 2012), with typically lower rates of tree growth (Toledo et al. 2011) and stem turnover (Quesada et al. 2012). Forests are more dynamic on the high-fertility soils of western Amazonia, with higher coarse woody productivity (Malhi et al. 2004, Baker et al. 2009), higher stem mortality (Johnson et al. 2016), lower basal area and aboveground biomass, and lower mean wood density (WD) than eastern Amazonia (Baker et al. 2004, Malhi et al. 2006, ter Steege et al. 2006, Quesada et al. 2012). Environmental gradients are also found across African tropical forests, where basal area decreases with both rainfall seasonality and soil fertility (sum of bases; Lewis et al. 2013).

Effects of competition on tree growth and mortality are expected to vary across continental environmental gradients in Amazonia and tropical Africa because water and soil nutrient availability influence forest structure and understory light availability. Competition has been hypothesized to intensify with resource availability because high resource levels lead to rapid growth and resource depletion, whereas plant growth is generally low in stressful habitats (Grime 1979). In tropical forests, competition is likely to be strongest at high resource (water and/or soil nutrient availability) levels, which support a higher basal area. Then, the resulting crowding leads to stronger competition because of reduced light availability to individual trees.

The response of any given focal tree to competition will likely depend not only on the degree of crowding in its local neighborhood, but also on its size and functional traits. Smaller trees are more strongly affected by competition (Uriarte et al. 2004) because they are more heavily shaded by taller neighbors, and likely suffer from greater belowground competition. Shade-intolerant tree species, which typically have low wood density (WD; van Gelder et al. 2006), respond more strongly to changes in light availability than shade-tolerant species (Bazzaz 1979), and thus are likely to be more strongly affected by competition. Indeed, shade-intolerant (Hubbell et al. 2001, Canham et al. 2006, Kunstler et al. 2011) and low WD tree species (Kunstler et al. 2016) often show greater growth decreases in response to

neighborhood crowding. Hence, variation in the plot-level strength of competition (i.e., the extent to which growth is reduced, or mortality is increased, by competition across all individual trees in a plot) across environmental gradients may not only depend on forest basal area, but also on tree size distributions and mean wood density. Nevertheless, forest basal area is expected to have the largest effect, because the basal area of neighbor trees directly influences resource availability to a focal tree.

In this study, we quantify the effects of neighborhood crowding on tree growth and mortality across gradients of moisture and soil nutrient availability in Amazonia and tropical Africa. Neighborhood crowding likely reflects competition for light (although competition for water and soil nutrients may also play a role), as light is typically the main factor limiting tree growth in closed-canopy forests. We use data from 151 ~1 ha-plots to fit nonlinear growth and mortality models based on tree WD, size, and neighborhood crowding. We use these models to estimate the predicted plot-level strength of competition, i.e., to what extent growth across all trees is reduced compared to a low level of neighborhood crowding, and assess how water availability and soil fertility influence the strength of competition through relationships with average tree size, plot basal area, and plot wood density. Specifically, we test the following predictions: (1) tree growth will decrease, and mortality increase, with neighborhood crowding; (2) low WD species will be most strongly affected by neighborhood crowding; (3) variation in the plot-level strength of competition will be more strongly related to plot basal area than to wood density or mean tree size; (4) the plot-level strength of competition will intensify with increasing climatic water availability through relationships with plot basal area on both continents; and (5) the predicted plot-level strength of competition will be negatively related to soil fertility in Africa because of decreasing basal area with increasing soil fertility (sum of bases; Lewis et al. 2013), but be largely independent of soil fertility in Amazonia because of weak correlations between soil fertility and basal area (Quesada et al. 2012).

METHODS

Plot data

We used data from 102 permanent plots in Amazonia from the RAINFOR network and 49 in tropical Africa from the AfriTRON network, curated at ForestPlots.net (Lopez-Gonzalez et al. 2009, 2011; Fig. 1), to span the environmental gradients in each tropical lowland forest region. Plots were all below 500 m above sea level (a.s.l.), in nonflooded, closed-canopy forests, with a five-fold range of mean annual precipitation in Amazonia (855–4273 mm) and twofold range in Africa (1,377–2,716 mm). Soil fertility, estimated by soil total exchange bases (in cmol(+)/kg), varied from 0.5 to 13.2 cmol(+)

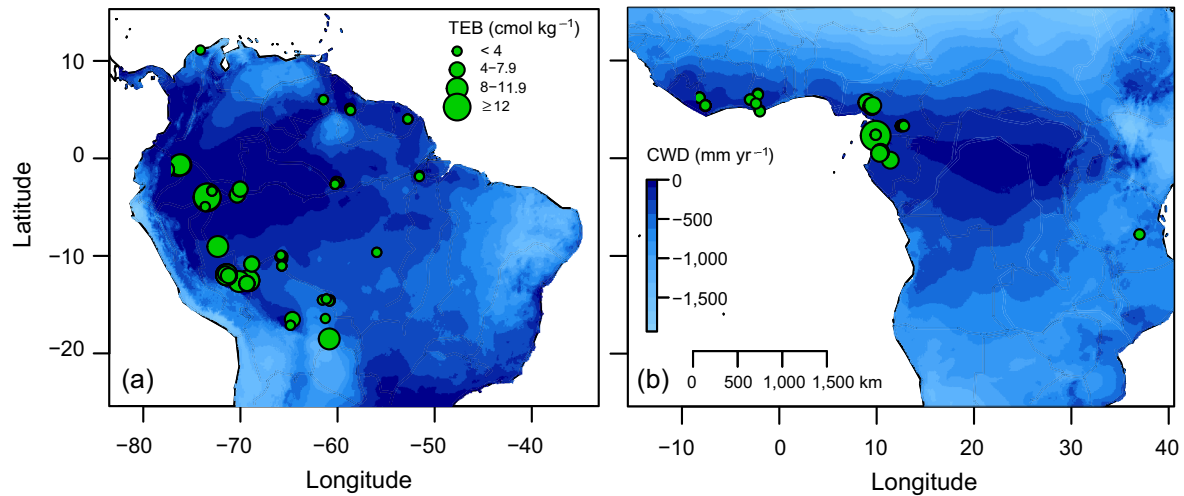


FIG. 1. Maps of the plot locations across gradients in climatic water deficit (CWD) and soil total exchange bases (TEB). (a) Amazonia (102 plots); (b) tropical Africa (49 plots).

kg in Amazonia, and from 2 to 13.5 cmol(+)/kg in Africa. Most plots were 1 ha in size, but plot size ranged from 0.25 to 9 ha (Appendix S1: Table S1). Trees ≥ 10 -cm diameter at breast height (dbh), or above buttresses, were measured for their diameter, identified to species, and either mapped or assigned to 0.04-ha subplots. Across all plots, 2,947 species and 73,100 trees were included in Amazonia, and 695 species and 20,705 trees in Africa. For each plot, we included data from two censuses with an average interval length of 6.3 yr (range: 3.0–12.7 yr; Appendix S1: Table S1) and an average starting year of 1994 (range: 1971–2008), and calculated annual basal area growth (in cm²/yr) for trees that were present in both censuses. We excluded monocotyledonous species (palms and Strelitziaceae) from the growth models, as they do not have secondary growth. Neighborhood crowding was expressed as the total basal area of neighbor trees within a 0.04-ha subplot (BA_{neigh}) in the first census. We defined neighborhoods based on subplots instead of on a fixed radius around each focal tree, to allow inclusion of plots for which individual trees were not mapped (Appendix S2). We found that BA_{neigh} accurately captured local effects of competition (Appendix S2). Neighborhood crowding likely reflects competition for light, although competition for water and soil nutrients may also occur. Other processes, for example pathogen accumulation at high densities of conspecific trees that increase mortality (negative density dependence; NDD), may also contribute, but effects of NDD are typically weak for large trees (Zhu et al. 2015).

Environmental conditions and wood density

Average annual rainfall (in mm/yr) for each of the plots was obtained from WorldClim 2 (Fick and Hijmans 2017). Climatic water deficit (CWD; in mm/yr; Chave et al. 2014) was included as a measure of seasonal

drought stress.¹ CWD is defined as the cumulative amount of water lost by the environment during months in which evapotranspiration exceeds rainfall. CWD is negative for sites that experience seasonal drought stress; a CWD of 0 indicates absence of seasonal drought stress. Topsoil total exchange bases (TEB; in cmol(+)/kg) was included as an indicator of soil fertility, and was obtained from the World Harmonized Soil Database (FAO/IIASA/ISRIC/ISS-CAS/JRC 2012). Wood density (WD) data were obtained from a global database (Chave et al. 2009, Zanne et al. 2009). In cases where a species-specific WD value was not available, we used genus- or family-level mean WD (Baker et al. 2004). Genus-level WD was used for 1,578 (out of 2,947) and for 233 (out of 695) species in Amazonia and Africa, respectively. Family-level WD was used for 235 and 186 species in Amazonia and Africa, respectively. For stems that remained unidentified, or for which family-level mean WD was unavailable (for 37 species in Amazonia and 31 in Africa), we used the mean WD across all stems in the plot.

Modeling approach

We used a combination of (modeling) approaches to evaluate whether the predicted strength of competition varied across environmental gradients in Amazonia and Africa. First, we used the plot data from both continents to construct nonlinear models of individual tree growth and mortality as functions of tree size (dbh), neighborhood crowding, and WD. Separate models were fitted for Amazonia and tropical Africa. Second, we used the estimated parameters of the fitted growth models to calculate the strength of competition (C_{plot}) at the plot level (mortality was excluded because competition effects on

¹ http://chave.ups-tlse.fr/pantropical_allometry.htm

mortality were very weak; see Results). As a last step, we assessed (1) whether C_{plot} varied with water availability and soil fertility, and (2) how C_{plot} was influenced by variation in plot basal area, plot-level WD, and average tree size. Variation in C_{plot} could arise from plot-to-plot differences in average neighborhood crowding (i.e., plot basal area), average WD, or average tree size, as each of these influenced the modeled effect of competition on individual tree growth. We describe each of these steps in greater detail below.

We modeled the annual basal area growth (G) and the annual probability of mortality (M) for individual trees on each continent as follows:

$$G = a_G \times p_G \times S_G \times C_G$$

$$M = (1 + a_M \times p_M \times S_M \times C_M)^{-1}$$

where a_G and a_M are constants, p_G and p_M are plot-level random effects, and S and C (each subscripted for growth and mortality) are nonlinear functions that capture effects of tree size and competition, respectively:

$$S = \text{dbh}^{s_1} \times \exp(-s_2 \times \text{dbh})$$

$$C = \exp(-c_1 \times \text{dbh}^{c_2} \times \text{BA}_{\text{neigh}})$$

where s_1 , s_2 , c_1 , and c_2 control the shape of the functions and have separate values for growth and mortality. S has a flexible form that can produce either an intermediate peak or a continuous increase in tree growth with tree size (dbh; Coomes et al. 2014). For mortality, S can produce a U-shaped response where mortality both decreases with size for small trees and increases with size for larger trees (Rüger et al. 2011, Iida et al. 2014). C is a decreasing function that can produce lower growth and higher mortality in trees with greater neighborhood crowding. The sensitivity of growth and mortality to competition may vary with tree size (as determined by c_2), as large trees may be less susceptible to competition than small trees.

We applied a trait-based approach to account for taxonomic variation in growth and mortality, as a species-level approach was not feasible given the huge diversity of tree species in the tropics (e.g., an estimated 15,000 tree species in the Amazon basin; ter Steege et al. 2015). WD is known to be a good predictor of tropical tree growth and mortality (e.g., Chao et al. 2008, Poorter et al. 2008, Wright et al. 2010, Rüger et al. 2012, Aleixo et al. 2019); therefore we defined model parameters a , s_1 , s_2 , c_1 , and c_2 as linear functions of WD. As such, WD could influence growth and mortality directly, as well as indirectly through effects on size relationships and responses to competition (e.g., Hérault et al. 2011, Iida et al. 2014, Kunstler et al. 2016). Models were fit using a hierarchical Bayesian approach (Appendix S2, Data S1: Model_script.R).

Using the fitted growth models, we calculated the strength of competition for each plot (C_{plot}) as the percent reduction in plot-level basal area growth due to competition compared to a low, baseline level of neighborhood crowding by assessing to what extent growth was reduced for each individual tree:

$$C_{\text{plot}} = \left(1 - \left(\frac{\sum_{i=1}^n G_i^{(c)}}{\sum_{i=1}^n G_i^{(lc)}} \right) \right) \times 100$$

where for tree i , $G_i^{(c)}$ represents predicted basal area growth with the observed level of competition, and $G_i^{(lc)}$ represents its potential growth at a low, baseline level of competition. Quantifying plot-level competition based on the growth reduction compared to potential growth in the absence of competition may be unrealistic, because a BA_{neigh} of zero is rarely found. Per continent, we calculated the 10th percentile of the plot-level 10th percentile values of BA_{neigh} (11.3 m²/ha for Amazonia; 9.8 m²/ha for Africa). We therefore calculated the strength of competition based on a general baseline level of $\text{BA}_{\text{neigh}} = 10$ m²/ha for both continents. Thus, C_{plot} was calculated by comparing predicted plot-level growth (based on all individual trees) with competition to growth at a BA_{neigh} of 10 m²/ha. Growth predictions were based on the posterior means of the model parameters.

For each continent, we examined whether C_{plot} was correlated with water availability (CWD) or soil fertility (TEB). To assess whether variation in C_{plot} was driven by variation in plot basal area, plot-level WD (basal area-weighted mean), or average tree size (the diameter of a tree with mean basal area; $\sqrt{(\sum \text{dbh}^2)/n}$), we modeled C_{plot} as a function of plot BA, plot-level WD, and average tree size using linear regression. In order to compare effect sizes among the three predictors, predictors were standardized by subtracting the mean and dividing the difference by the standard deviation. All analyses were performed in R 3.1.2 (R Development Core Team 2014).

RESULTS

Overall responses to competition

Individual tree growth was strongly affected by competition (Fig. 2; Appendix S1: Table S2), but competition effects were stronger in Amazonian than African tropical forests. For example, for a 20-cm diameter tree with a WD of 0.6 g cm⁻³, growth decreased by 34% in Amazonia (Fig. 2e, g) and 17% in Africa (Fig. 2f, h) as BA_{neigh} increased from 10 to 50 m²/ha. Further, even though plot-level basal area was on average slightly lower in Amazonia (25.9 ± 0.44 m²/ha; mean \pm SE) than in Africa (28.7 ± 0.64 m²/ha), the stronger response of trees to competition in Amazonia resulted in greater predicted decreases in plot-level wood production than in Africa. Competition reduced

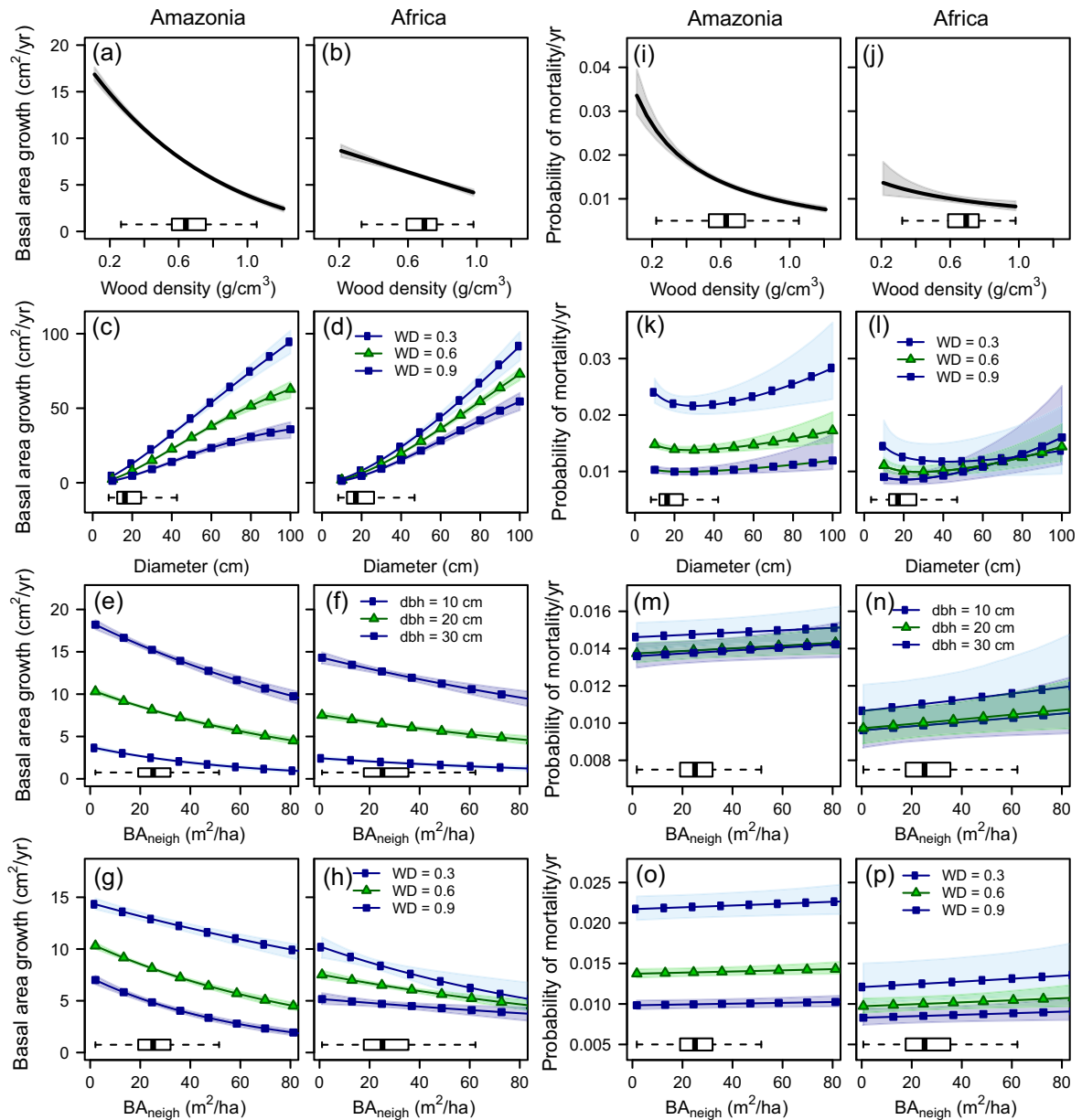


FIG. 2. Effects of wood density (WD), tree size, and competition (subplot neighbor basal area; BA_{neigh}) on predicted annual basal area growth and mortality across Amazonia ($n = 102$ plots) and tropical Africa ($n = 49$). Solid lines and symbols indicate predicted effects based on the posterior means; shaded areas indicate the 95% credible interval. Boxplots indicate the distribution of the variable on the x-axis. BA_{neigh} was kept constant at the mean for quantifying effects of WD and tree size on growth and mortality; tree size was kept constant at 20-cm diameter for quantifying effects of WD and BA_{neigh}. dbh = diameter at breast height.

plot-level basal area growth (compared to a baseline, low-BA_{neigh} value of 10 m²/ha) by, on average, 31.1% (range: 4.5–25.2%; Fig. 3a, c) in Amazonia, and by 7.4% in Africa (range: 5.3–11.7%; Fig. 3b, d).

In contrast to effects on growth, competition with neighboring trees had little or no effect on the probability of mortality. Nevertheless, the mortality model that

included competition performed better than the no-competition model for Amazonia (Appendix S1: Table S2). The predicted probability of mortality for a 20-cm dbh tree with a WD of 0.6 gr cm⁻³ remained constant at 1.4% (Fig. 2m, o) and 1.0% (Fig. 2n, p) per year as BA_{neigh} increased from 10 to 50 m² ha⁻¹ in Amazonia and Africa, respectively.

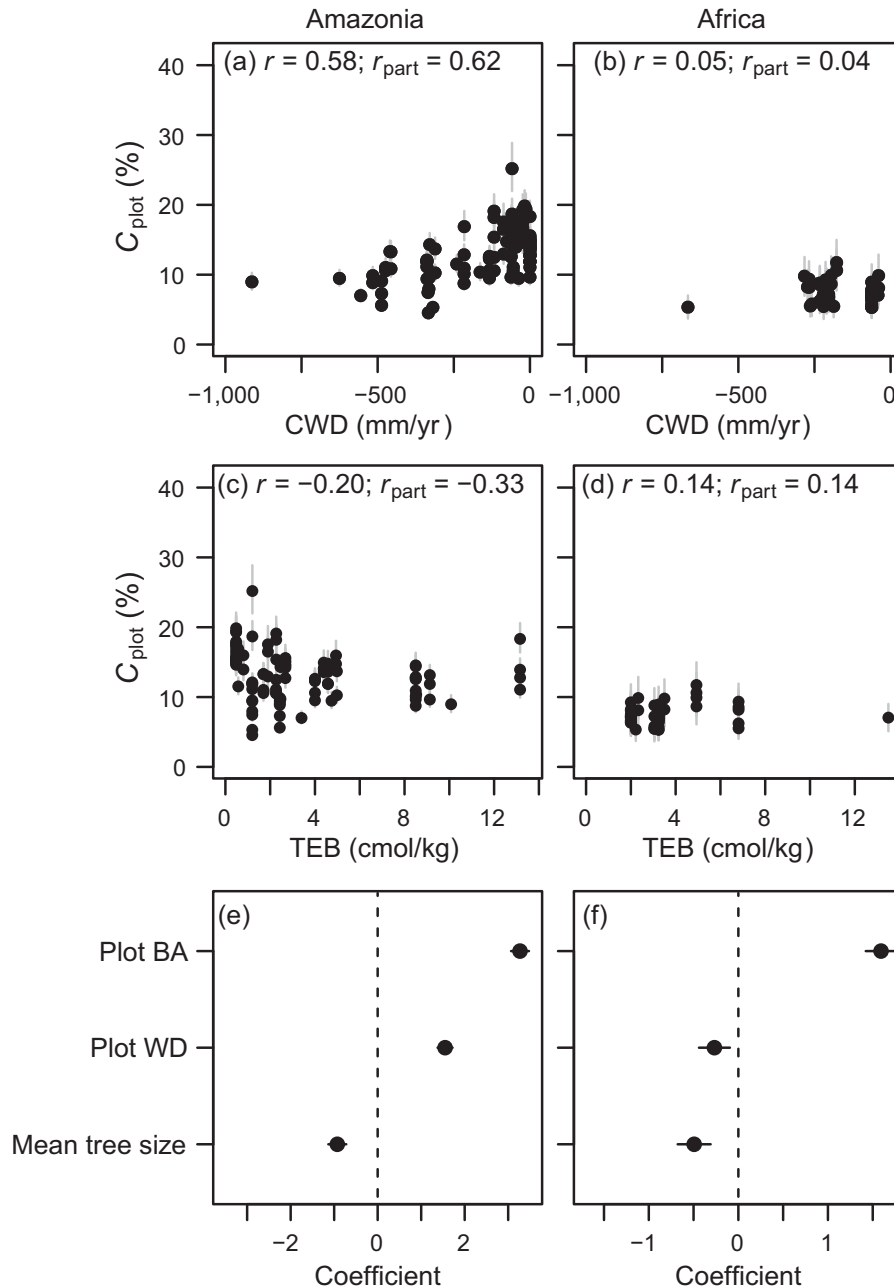


FIG. 3. Relationships between the strength of competition on basal area growth (C_{plot} : reduction in plot-level basal area growth by competition based on a reference value of 10 m²/ha) and climatic water deficit (CWD), soil total exchange bases (TEB), plot basal area (BA), plot wood density (WD), and mean tree size in Amazonia ($n = 102$ plots) and tropical Africa ($n = 49$ plots). (a–d) Gray bars represent 95% credible intervals; Pearson’s correlation (r) and partial (r_{part}) correlation coefficients are indicated; (e, f) standardized regression coefficients with 95% confidence intervals are indicated.

Effects of wood density and tree size

Tree basal area growth decreased with increasing WD on both continents (Fig. 2a, b). In Amazonia, a 20-cm tree with low WD (0.3 g/cm³) grew more than twice as fast as a high-WD (0.9 g/cm³) tree of the same size (Fig. 2a). In

Africa, the growth decrease with increasing WD was less pronounced (Fig. 2b). Growth increased with tree size on both continents (Fig. 2c, d), with low WD species exhibiting stronger size-related increases in growth.

On both continents, small trees were more strongly affected by competition than large trees (Fig. 2e, f). In

Amazonia, growth of a 10-cm tree and a 30-cm tree decreased by 49% and 27%, respectively, as neighbor basal area increased from 10 to 50 m²/ha (Fig. 2e). Similar growth decreases were found in Africa, with a 28% and 18% growth decrease for a 10-cm and a 30-cm tree, respectively (Fig. 2f). Amazonian trees with different WD showed similar absolute decreases in growth resulting from competition, but on a proportional basis high WD species expressed greater decreases than low WD species (48% and 17%, respectively) as BA_{neigh} increased from 10 to 50 m²/ha (Fig. 2g). Conversely, the growth of high WD species in Africa was less affected by competition than that of low WD species (decreases of 14% and 28%, respectively; Fig. 2h).

The probability of mortality decreased with WD on both continents (Fig. 2i, j), but the decline was more pronounced and more consistent in Amazonia than in Africa. Mortality generally increased with tree size (Fig. 2k, l), particularly for trees >50-cm dbh, although low abundances increased uncertainty for large trees. Small trees with low WD had higher mortality than mid-sized trees (7% and 23% higher mortality at 10-cm dbh than at 50-cm dbh in Amazonia and tropical Africa, respectively), leading to a U-shaped size–mortality relationship. Effects of competition on mortality were very weak on both continents, regardless of WD or tree size (Fig. 2m, n, o, p).

Variation in the strength of competition

In Amazonia, the plot-level strength of competition (C_{plot}) was strongly and positively correlated with CWD, but negatively correlated with TEB, particularly after accounting for variation in CWD (Fig. 3a, b). Plot basal area had the largest effect on C_{plot} , followed by a positive effect of plot WD, and a small negative effect of mean tree size (Fig. 3e). In Africa, C_{plot} was not correlated with CWD, and just weakly, positively correlated with TEB (Fig. 3c, d). Like in Amazonia, C_{plot} was largely driven by a positive effect of plot basal area. Unlike Amazonia, plot-level WD had little influence on C_{plot} in tropical Africa (Fig. 3f).

DISCUSSION

Large variation in the strength of competition on tree growth across environmental gradients

Across two continents, we found that competition is an important driver of tropical tree growth, but unexpectedly not of mortality. Variation in the plot-level strength of competition across tropical forests was large for both continents. As expected, individual tree growth was most strongly affected by competition in forests with high basal area, although in Amazonia competition was also strong in high WD forests. In Amazonia, as expected, the strength of competition on tree growth increased with water availability (CWD), likely because

of higher plot basal area in wetter forests (Appendix S1: Fig. S1). However, the strength of competition declined slightly with soil fertility (TEB), likely because of lower plot-level WD at high soil fertility (Appendix S1: Table S3), and because low WD species in Amazonia appeared to be less susceptible to competition. Unexpectedly, the strength of competition did not vary with water availability, nor with soil fertility, in Africa. This may have been due to the shorter water availability and soil fertility gradients compared to Amazonia in our study, which likely partly explains the lack of relationships with environmental conditions in tropical Africa. Given these differences, we must be careful in drawing general conclusions across continents. Across the same range in environmental conditions (based on Africa, excluding two outliers; Fig. 3b, d), the relationship between the strength of competition and CWD was stronger in Amazonia (Pearson's $r = 0.40$, $n = 38$ plots) than in Africa ($r = -0.12$). The relationship between the strength of competition and TEB was somewhat stronger for Africa because of outlier exclusion ($r = 0.23$) than for Amazonia ($r = 0.10$, $n = 41$ plots). Overall, our results are partly consistent with Grime's (1979) hypothesis that competition is strongest in resource-rich environments because of the increased strength of competition under high water availability in Amazonia.

Effects of WD and tree size on growth and mortality

In contrast, effects of WD and tree size on individual tree growth and mortality were largely consistent between Amazonia and tropical Africa. In general, our results confirmed findings of previous studies that were based on a single, or a few, tropical forest sites, and indicated that these attributes control growth and mortality across most of the tropical forest biome. Tree growth and mortality both decreased with WD, as reported by smaller-scale Neotropical studies (e.g., Chao et al. 2008, Keeling et al. 2008, Poorter et al. 2008, Wright et al. 2010, Rüger et al. 2012). Low WD is associated with an acquisitive strategy that confers rapid growth, but that comes at the cost of high mortality because of lower tolerance to stress and damage compared to high WD species (Wright et al. 2010). Basal area growth increased with tree size, presumably because larger trees have more resources and/or leaf area available to support assimilation of carbon (Stephenson et al. 2014). The ontogenetic increase in growth was strongest for low WD species (Fig. 2c, d), probably because of the low construction cost of low-density wood. These findings are consistent with single-site studies that found that low-WD tropical tree species had the strongest increase in diameter growth at intermediate tree size (King et al. 2006, Hérault et al. 2011, but see Rüger et al. 2012).

Our study is one of the first to show a clearly U-shaped size–mortality relationship (cf. Rüger et al. 2011, Iida et al. 2014, Pillet et al. 2018), which we found for low-WD species. For trees ≥ 30 cm dbh, and for high

WD trees in general, the risk of death increased nearly monotonically with size. Small trees, particularly those with low WD, may be most susceptible to physical damage in the understory (Clark and Clark 1991). The higher mortality risk for large trees may be a result of the stronger risk of hydraulic failure for large trees (Rowland et al. 2015) rather than senescence (Mencuccini et al. 2005).

Competition decreased tree growth but did not influence mortality

Our results show that growth decreases with increased neighborhood crowding across tropical forests on two continents, particularly for small trees. This provides large-scale confirmation that results reported to date for single Neotropical forest sites in Costa Rica, Ecuador, Panama, and Puerto Rico (Uriarte et al. 2004, Baribault et al. 2012, Grote et al. 2013, Lasky et al. 2015, Fortunel et al. 2016) are typical of the biome. We also expected that low-WD species would be most strongly affected by competition. Low-WD species were indeed most affected by competition in Africa, consistent with earlier findings of strong growth responses of low-WD species to competition (Kunstler et al. 2016) and light availability (Rüger et al. 2012), which supports the notion that shade-intolerant tree species respond more strongly to changes in resource levels. However, it remains unclear why high WD species in Amazonia were more susceptible to competition. The mean and range of neighborhood crowding levels did not vary across WD classes ($<0.35 \text{ g/cm}^3$; $0.35\text{--}0.75 \text{ g/cm}^3$; $>0.75 \text{ g/cm}^3$; results not shown), thus effects of competition were not weaker because low WD species were confined to areas with low neighborhood crowding.

Our results suggest that competition does not strongly influence tree mortality in either Amazonia or tropical Africa. The lack of evidence for impacts of competition on mortality could be partly due to only including trees $\geq 10 \text{ cm dbh}$ in our study. Generally, mortality rates are highest for seedlings and saplings (trees $< 10 \text{ cm dbh}$; Clark and Clark 1992, Condit et al. 1995) because of the low-light conditions in the understory, and mortality resulting from negative density-dependent effects (Zhu et al. 2015). Those studies that have found clear effects of competition on tropical tree mortality included trees $< 10 \text{ cm dbh}$, and likely included a larger range of resource levels by focusing on forests in recovery from disturbances such as agricultural use (Lasky et al. 2014) and hurricanes (Uriarte et al. 2004). Our findings suggest that competition is not a widespread and important driver of mortality for trees $\geq 10 \text{ cm dbh}$ in mature tropical forests. Instead, it appears that processes such as hydraulic failure (e.g., Rowland et al. 2015) and stochastic wind disturbances (Espírito-Santo et al. 2014, Aleixo et al. 2019) may be the dominant causes of mortality, although accelerated growth may eventually increase mortality by ensuring that trees reach larger sizes more

quickly (cf. Brien et al. 2015, McDowell et al. 2018). Nevertheless, the effects of competition on growth may still indirectly lead to an increased risk of mortality, as suppressed trees will be less likely to escape from suppression because of their slow growth, and thus accumulate mortality risk over a longer period of time.

Implications for projecting the tropical forest carbon sink

Our results provide some insights into how competition may influence ongoing and future changes in the tropical forest carbon sink. First, we found that the decrease in basal area growth due to competition increased strongly with forest basal area. Hence, when forests gain basal area over time, greater competition between trees is likely to reduce tree growth, which might explain why long-term increases in productivity in Amazonia have leveled off since 2000 (Brien et al. 2015). Secondly, we found that, particularly in Amazonia, effects of competition are also influenced by stand-level WD. Changes in WD over time (e.g., van der Sande et al. 2016) may not only influence standing biomass (Baker et al. 2004), but also alter the strength of competition.

Competition effects should be appropriately incorporated into models that are used for projecting future dynamics of tropical forests. In individual-based forest dynamics models, effects of competition are typically included (Fyllas et al. 2014), but models could be further improved by also including effects of WD, and tree size, on the strength of competition. These changes are relatively easy to implement, as direct effects of tree size are already included, and WD data are available for many species (Chave et al. 2009). In dynamic global vegetation models that are applied over broad geographical scales, inclusion of forest basal area as a measure of neighborhood crowding will mostly account for geographical variation in the strength of competition. Such models could be improved further by including average plot WD.

In conclusion, our study revealed that in 151 forest plots distributed across Amazonia and tropical Africa competition is an important driver of individual tree growth rates, but not of the probability of tree mortality. This is, to our knowledge, the first study to evaluate the effects of competition on tropical tree growth and mortality at such a broad geographical scale. Given that geographic variation in the strength of competition is mainly driven by forest basal area (i.e., neighborhood crowding), we anticipate that wood production might decrease as tropical forests accrue higher basal area.

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LITERATURE CITED

- Aleixo, I., D. Norris, L. Hemerik, A. Barbosa, E. Prata, F. Costa, and L. Poorter. 2019. Amazonian rainforest tree mortality driven by climate and functional traits. *Nature Climate Change* 9:384–388.
- Baker, T. R. et al. 2004. Variation in wood density determines spatial patterns in Amazonian forest biomass. *Global Change Biology* 10:545–562.
- Baker, T. R. et al. 2009. Do species traits determine patterns of wood production in Amazonian forests? *Biogeosciences* 6:297–307.
- Baribault, T. W., R. K. Kobe, and A. O. Finley. 2012. Tropical tree growth is correlated with soil phosphorus, potassium, and calcium, though not for legumes. *Ecological Monographs* 82:189–203.
- Bazzaz, F. A. 1979. Physiological ecology of plant succession. *Annual Review of Ecology and Systematics* 10:351–371.
- Bennett, A. C., N. G. McDowell, C. D. Allen, and K. J. Anderson-Teixeira. 2015. Larger trees suffer most during drought in forests worldwide. *Nature Plants* 1:15139.
- Brienen, R. J. W. et al. 2015. Long-term decline of the Amazon carbon sink. *Nature* 519:344–348.
- Canham, C. D., M. J. Papaik, M. Uriarte, W. H. McWilliams, J. C. Jenkins, and M. J. Twery. 2006. Neighborhood analyses of canopy tree competition along environmental gradients in New England forests. *Ecological Applications* 16:540–554.
- Chao, K. J., O. L. Phillips, E. Gloor, A. Monteagudo, A. Torres-Lezama, and R. Vásquez-Martínez. 2008. Growth and wood density predict tree mortality in Amazon forests. *Journal of Ecology* 96:281–292.
- Chave, J., D. Coomes, S. Jansen, S. L. Lewis, N. G. Swenson, and A. E. Zanne. 2009. Towards a worldwide wood economics spectrum. *Ecology Letters* 12:351–366.
- Chave, J. et al. 2014. Improved allometric models to estimate the aboveground biomass of tropical trees. *Global Change Biology* 20:3177–3190.
- Clark, D. A., and D. B. Clark. 1992. Life-history diversity of canopy and emergent trees in a neotropical rain-forest. *Ecological Monographs* 62:315–344.
- Clark, D. B., and D. A. Clark. 1991. The impact of physical damage on canopy tree regeneration in tropical rain-forest. *Journal of Ecology* 79:447–457.
- Condit, R., S. P. Hubbell, and R. B. Foster. 1995. Mortality rates of 205 Neotropical tree and shrub species and the impact of a severe drought. *Ecological Monographs* 65:419–439.
- Coomes, D. A., O. Flores, R. Holdaway, T. Jucker, E. R. Lines, and M. C. Vanderwel. 2014. Wood production response to climate change will depend critically on forest composition and structure. *Global Change Biology* 20:3632–3645.
- Espírito-Santo, F. D. B. et al. 2014. Size and frequency of natural forest disturbances and the Amazon forest carbon balance. *Nature Communications* 5:3434.
- FAO/IIASA/ISRIC/ISS-CAS/JRC. 2012. Harmonized World Soil Database (version 1.2). FAO, Rome, Italy and IIASA, Laxenburg, Austria.
- Fick, S. E., and R. J. Hijmans. 2017. WorldClim 2: new 1-km spatial resolution climate surfaces for global land areas. *International Journal of Climatology* 37:4302–4315.
- Fortunel, C., R. Valencia, S. J. Wright, N. C. Garwood, and N. J. B. Kraft. 2016. Functional trait differences influence neighbourhood interactions in a hyperdiverse Amazonian forest. *Ecology Letters* 19:1062–1070.
- Fyllas, N. M. et al. 2014. Analysing Amazonian forest productivity using a new individual and trait-based model (TFS vol. 1). *Geoscientific Model Development* 7:1251–1269.
- Grime, J. P. 1979. Plant strategies and vegetation processes. John Wiley and Sons, Chichester, UK.
- Grote, S., R. Condit, S. Hubbell, C. Wirth, and N. Rüger. 2013. Response of demographic rates of tropical trees to light availability: Can position-based competition indices replace information from canopy census data? *PLoS ONE* 8:e81787.
- Héroult, B., B. Bachelot, L. Poorter, V. Rossi, F. Bongers, J. Chave, C. E. T. Paine, F. Wagner, and C. Baraloto. 2011. Functional traits shape ontogenetic growth trajectories of rain forest tree species. *Journal of Ecology* 99:1431–1440.
- Hubbell, S. P., J. A. Ahumada, R. Condit, and R. B. Foster. 2001. Local neighborhood effects on long-term survival of individual trees in a neotropical forest. *Ecological Research* 16:859–875.
- Iida, Y., L. Poorter, F. Sterck, A. R. Kassim, M. D. Potts, T. Kubo, and T. S. Kohyama. 2014. Linking size-dependent growth and mortality with architectural traits across 145 co-occurring tropical tree species. *Ecology* 95:353–363.
- Johnson, M. O. et al. 2016. Variation in stem mortality rates determines patterns of above-ground biomass in Amazonian forests: implications for dynamic global vegetation models. *Global Change Biology* 22:3996–4013.
- Keeling, H. C., T. Baker, R. Martinez, A. Monteagudo, and O. Phillips. 2008. Contrasting patterns of diameter and biomass increment across tree functional groups in Amazonian forests. *Oecologia* 158:521–534.
- King, D. A., S. J. Davies, and N. S. M. Noor. 2006. Growth and mortality are related to adult tree size in a Malaysian mixed dipterocarp forest. *Forest Ecology and Management* 223:152–158.
- Kunstler, G., C. H. Albert, B. Courbaud, S. Lavergne, W. Thuiller, G. Vieilledent, N. E. Zimmermann, and D. A. Coomes. 2011. Effects of competition on tree radial-growth vary in importance but not in intensity along climatic gradients. *Journal of Ecology* 99:300–312.
- Kunstler, G. et al. 2016. Plant functional traits have globally consistent effects on competition. *Nature* 529:204–U174.
- Lasky, J. R. et al. 2015. Ontogenetic shifts in trait-mediated mechanisms of plant community assembly. *Ecology* 96:2157–2169.
- Lasky, J. R., M. Uriarte, V. K. Boukili, and R. L. Chazdon. 2014. Trait-mediated assembly processes predict successional changes in community diversity of tropical forests. *Proceedings of the National Academy of Sciences of the United States of America* 111:5616–5621.
- Lewis, S. L. et al. 2009. Increasing carbon storage in intact African tropical forests. *Nature* 457:1003–U1003.

- Lewis, S. L. et al. 2013. Above-ground biomass and structure of 260 African tropical forests. *Philosophical Transactions of the Royal Society B* <http://dx.doi.org/10.1098/rstb.2012.0295>
- Lopez-Gonzalez, G., S. L. Lewis, M. Burkitt, and O. L. Phillips. 2009. ForestPlots.net Database. www.forestplots.net
- Lopez-Gonzalez, G., S. L. Lewis, M. Burkitt, and O. L. Phillips. 2011. ForestPlots.net: a web application and research tool to manage and analyse tropical forest plot data. *Journal of Vegetation Science* 22:610–613.
- Malhi, Y. et al. 2004. The above-ground coarse wood productivity of 104 Neotropical forest plots. *Global Change Biology* 10:563–591.
- Malhi, Y. et al. 2006. The regional variation of aboveground live biomass in old-growth Amazonian forests. *Global Change Biology* 12:1107–1138.
- McDowell, N. et al. 2018. Drivers and mechanisms of tree mortality in moist tropical forests. *New Phytologist* 219:851–869.
- Mencuccini, M., J. Martinez-Vilalta, D. Vanderklein, H. A. Hamid, E. Korakaki, S. Lee, and B. Michiels. 2005. Size-mediated ageing reduces vigour in trees. *Ecology Letters* 8:1183–1190.
- Phillips, O. L. et al. 2010. Drought–mortality relationships for tropical forests. *New Phytologist* 187:631–646.
- Pillet, M. et al. 2018. Disentangling competitive vs. climatic drivers of tropical forest mortality. *Journal of Ecology* 106:1165–1179.
- Poorter, L. et al. 2008. Are functional traits good predictors of demographic rates? Evidence from five Neotropical forests. *Ecology* 89:1908–1920.
- Quesada, C. A. et al. 2012. Basin-wide variations in Amazon forest structure and function are mediated by both soils and climate. *Biogeosciences* 9:2203–2246.
- R Development Core Team. 2014. R: A language and environment for statistical computing. R Foundation for Statistical Computing, Vienna, Austria. www.r-project.org
- Rowland, L. et al. 2015. Death from drought in tropical forests is triggered by hydraulics not carbon starvation. *Nature* 528:119–122.
- Rüger, N., A. Huth, S. P. Hubbell, and R. Condit. 2011. Determinants of mortality across a tropical lowland rainforest community. *Oikos* 120:1047–1056.
- Rüger, N., C. Wirth, S. J. Wright, and R. Condit. 2012. Functional traits explain light and size response of growth rates in tropical tree species. *Ecology* 93:2626–2636.
- Stephenson, N. L. et al. 2014. Rate of tree carbon accumulation increases continuously with tree size. *Nature* 507:90–93.
- ter Steege, H. et al. 2006. Continental-scale patterns of canopy tree composition and function across Amazonia. *Nature* 443:444–447.
- ter Steege, H. et al. 2015. Estimating the global conservation status of more than 15,000 Amazonian tree species. *Science Advances* 1:e1500936.
- Toledo, M. et al. 2011. Climate is a stronger driver of tree and forest growth rates than soil and disturbance. *Journal of Ecology* 99:254–264.
- Uriarte, M., C. D. Canham, J. Thompson, and J. K. Zimmerman. 2004. A neighborhood analysis of tree growth and survival in a hurricane-driven tropical forest. *Ecological Monographs* 74:591–614.
- van der Sande, M. T. et al. 2016. Old-growth Neotropical forests are shifting in species and trait composition. *Ecological Monographs* 86:228–243.
- van Gelder, H. A., L. Poorter, and F. J. Sterck. 2006. Wood mechanics, allometry, and life-history variation in a tropical rain forest tree community. *New Phytologist* 171:367–378.
- Wright, S. J. et al. 2010. Functional traits and the growth–mortality trade-off in tropical trees. *Ecology* 91:3664–3674.
- Zanne, A. E. et al. 2009. Data from: Towards a worldwide wood economics spectrum. Dryad Digital Repository. <https://doi.org/10.5061/dryad.234>
- Zhu, Y., L. S. Comita, S. P. Hubbell, and K. P. Ma. 2015. Conspecific and phylogenetic density-dependent survival differs across life stages in a tropical forest. *Journal of Ecology* 103:957–966.

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