

## RESEARCH ARTICLE

# Demographic composition, not demographic diversity, predicts biomass and turnover across temperate and tropical forests

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#### Abstract

The growth and survival of individual trees determine the physical structure of a forest with important consequences for forest function. However, given the diversity of tree species and forest biomes, quantifying the multitude of demographic strategies within and across forests and the way that they translate into forest structure and function remains a significant challenge. Here, we quantify the demographic rates of 1961 tree species from temperate and tropical forests and evaluate how demographic diversity (DD) and demographic composition (DC) differ across forests, and how these differences in demography relate to species richness, aboveground biomass (AGB), and carbon residence time. We find wide variation in DD and DC across forest plots, patterns that are not explained by species richness or climate variables alone. There is no evidence that DD has an effect on either AGB or carbon residence time. Rather, the DC of forests, specifically the relative abundance of large statured species, predicted both biomass and carbon residence time. Our results demonstrate the distinct DCs of globally distributed forests, reflecting biogeography, recent history, and current plot conditions. Linking the DC of forests to resilience or vulnerability to climate change, will improve the precision and accuracy of predictions of future forest composition, structure, and function.

#### KEYWORDS

aboveground biomass, carbon residence time, forest dynamics, ForestGEO, size-dependent survival, species richness, tree demography

## 1 | INTRODUCTION

Forests store approximately 50% of the world's carbon (Keenan & Williams, 2018) and play a critical role in regulating the world's

biogeochemical and hydrological cycles (Gloor et al., 2013). They cover 30% of the Earth's land surface and span vastly different environments. There are 50,000–80,000 tree species (Beech et al., 2017), and individual forest plots can vary in species richness by

over three orders of magnitude (Anderson-Teixeira et al., 2015; Davies et al., 2021).

Tree species vary in their life histories, demographic rates, and environmental specialization, yet within a forest, a diversity of life history strategies is not necessarily correlated with a diversity of species. Life-histories range from short-lived small understory species (Condit et al., 1994) to canopy species that reach 100 m in height (Koch et al., 2004; Sillett et al., 2010) or survive for thousands of years (Brokaw, 1987; Chambers et al., 1998). Life histories can be specialized for specific environmental conditions, for example, fast-growing pioneer species that occur in gaps (Brokaw, 1987), and can have very different functional roles within the ecosystem in terms of carbon storage (Hubau et al., 2019; Lutz et al., 2018) and nutrient cycling (Ordoñez et al., 2009). Life history types may reflect strategies for dealing with forest perturbations, and are expected to differ in their response to climate change and other global change factors (Phillips et al., 2002). Demographic rates form an important component of a species' overall life history strategy, reflecting the influence of both the environment and evolved strategies on the acquisition and allocation of resources. Greater species richness in the tropics, however, is not correlated with a greater range of growth and survival rates (Condit et al., 2006). Furthermore, the biomass and turnover of forests do not show the same magnitude of variation as species richness (Lutz et al., 2018), suggesting a large degree of redundancy in the way that demographic rates of species scale to forest structure and function.

Demographic rates have typically been quantified by arranging species on the slow-fast continuum; an axis from slow-growing, long-lived species to fast-growing, short-lived species (Wright et al., 2010). However, multiple species show demographic rates that do not align well with this single axis of demographic variation (Rüger et al., 2018, 2020; Russo et al., 2021). Furthermore, the strength of the slow-fast continuum varies among forests, due to differences in environmental conditions (Russo et al., 2021). In highly disturbed forests, the trade-off between growth and survival among species may be absent, as slow-growing, high survival species are never able to become established (Russo et al., 2021). Quantifying demographic diversity (DD), the range of demographic strategies, and demographic composition (DC), the relative abundance of different demographic strategies in a forest, therefore requires multiple axes of demographic variation (Rüger et al., 2018).

Here, we define the DD of a forest as the area occupied by species in demographic space—defined by multiple axes of demographic variation. Forests with high DD will have species with a wide range of growth and survival rates, whereas the species in forests with low DD will have a more limited range of growth and survival rates. We group species by demographic strategy through the use of clustering algorithms on species' positions in demographic space and define the DC of a forest as the relative abundance of demographic strategies. Due to the ubiquity of gap-phase dynamics in closed-canopy forests (Brokaw, 1987), we might expect to see some similarities in DC across forests because all forests will contain demographic types specialized for different light conditions. However, it is not known whether the same strategies are

comparable in absolute terms across forests, nor whether the proportions of demographic types vary across forests, or if particular demographic types are absent or overrepresented in some forests.

There are at least two hypotheses for how species richness might relate to forest biomass and carbon residence time via DD and DC, Figure 1. First, higher species richness could lead to higher DD and aboveground biomass (AGB) because of niche-partitioning, where a more efficient use of resources would emerge from finer partitioning of resource space (Tilman et al., 1997). Second, the mass ratio hypothesis, first proposed by (Grime, 1998), suggests that observed patterns of AGB relate not to the number of species, but rather to the demographic rates of dominant species in the forest community. In this case, species richness may be unrelated to DD and AGB, as it is the presence or absence of species with particular demographic strategies (i.e., forest DC), which define the structure and functioning of the forest. A number of features unique to a location, such as seasonality, recent land use history, geophysical features, biogeography, and disturbance regimes could all shape DC. The relationship between carbon residence time and species richness, DD, and DC is more complex as it depends on both AGB and AGB loss from mortality. Under the niche-complementarity hypothesis higher species richness would lead to higher DD and AGB but either lower or higher mortality rates, depending on the increase in DD. As a result, carbon residence times would likely increase, but in some cases could decrease. Under the mass ratio hypothesis, AGB and AGB losses from mortality have no relationship with species richness and DD; therefore, carbon residence time would be defined by the dominant species, rather than species diversity.

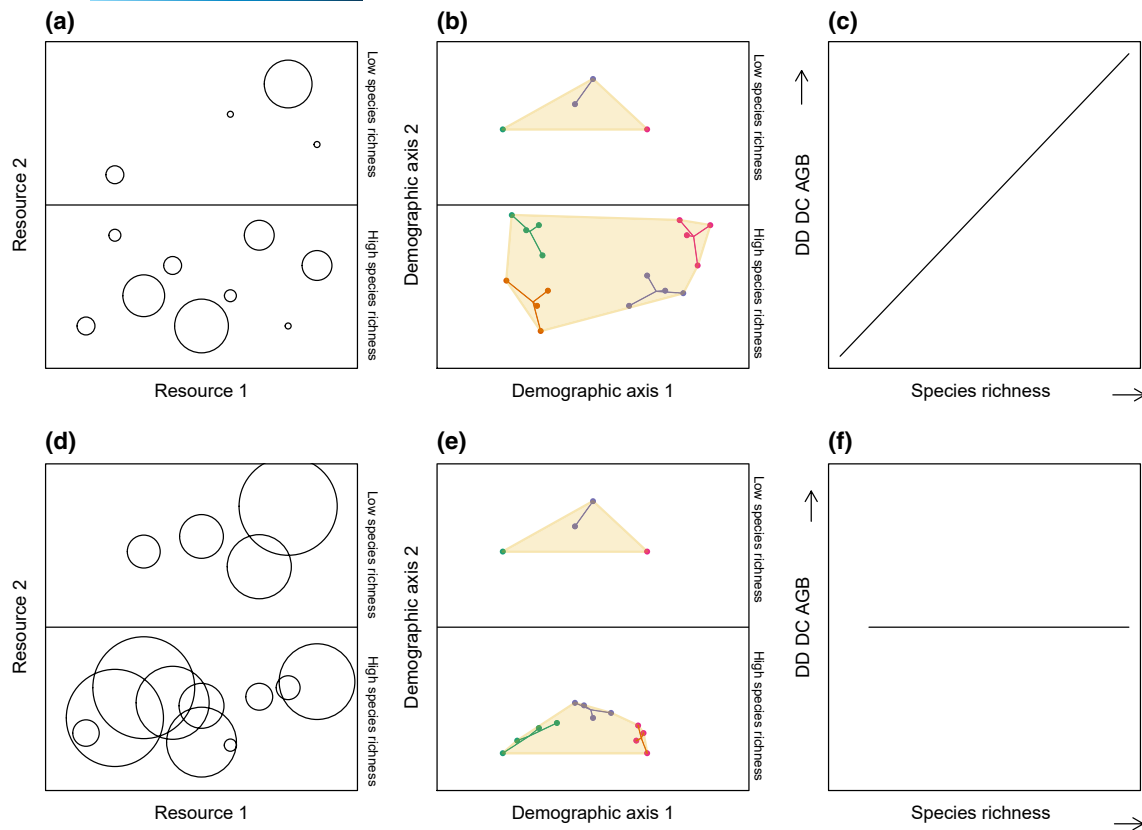
Although a number of studies have addressed aspects of demography and diversity either directly (e.g., Condit et al., 2006; Rüger et al., 2020), or indirectly through the effects of mechanisms on demographic rates (e.g., Comita et al., 2010), the majority of studies have focused on distinct biomes, and have not incorporated forests across orders of diversity, with distinct floras, and unique climate and disturbance regimes.

Here, we quantify the diversity of demographic rates across temperate and tropical forests in an effort to understand the extent to which demographic strategies explain how forests that exhibit conspicuously different species richness vary in terms of AGB and turnover, and how this relates to climatic conditions. We ask (1) How much does DD vary among forests around the world and how does species richness correlate with DD? (2) How do forests differ in DC, that is, how do the proportions of demographic strategies differ across forests? (3) How are DD and DC related to environmental conditions? and (4) How are DD and DC related to AGB and carbon residence time?

## 2 | METHODS

### 2.1 | Plots and data

We used demographic data from 20 plots across the ForestGEO network of permanent forest plots ([Davies et al., 2021], <https://forestgeo.si.edu>, [Figure S1]). Plots span a latitudinal gradient



**FIGURE 1** Conceptual model of the relationship between species richness, demographic diversity (DD), and aboveground biomass (AGB). In the niche complementarity hypothesis (top row) species have different resource requirements resulting in different demographic rates (a). (b) Shows the position of species in demographic space, defined by demographic rates, with DD depicted as the yellow convex hull around species, and demographic composition (DC) shown by clusters of species in different colors. As species richness increases more resources are used resulting in higher DD, DC, and AGB (c). Alternatively, in the mass ratio hypothesis (bottom row), species may overlap in their resource usage (d). Increasing species richness, therefore, does not necessarily result in greater DD or DC (e). Under this hypothesis, AGB is defined by the presence or absence of large statured species, that is, by demographic composition. DD, DC and AGB may have no relationship with species richness

from the most equatorial at 0°69'S (Yasuní Ecuador) to the most northerly at 51°77'N (Wytham, UK). Data collection at each plot follows a standardized protocol described in Condit (1998), where every stem  $\geq 1$  cm DBH (diameter at breast height, 1.3 m above the ground) is measured, mapped, and identified to species. Censuses are typically carried out every five years. For this analysis, we used one census interval from every plot (see Table S1). To have sufficient sample sizes, we limited our analysis to species with at least 200 individuals. In highly diverse or relatively small plots, this threshold means that many rare species were excluded from the analysis. However, due to highly skewed species abundance distributions, the percentage of stems included in the analysis ranged from 45% to 98% across plots, with a mean of 84% of stems included. Tree ferns, palms, lianas, and bamboos were also excluded due to non-standard growth forms. In total, the data set includes 2,195,135 stems of 1961 species. For further details of each plot see Table S1. Although some of the plots experience disturbance, there were no major cyclones or droughts at any of the plots between the two censuses that we used in this analysis.

## 2.2 | Statistical analysis

### 2.2.1 | Survival

We simultaneously fitted separate size-dependent survival curves to small and large stems, using a common asymptote, so that survival probability was a smooth function of stem size.

We modelled size dependent survival probability across a census interval, as:

$$s = \left( \frac{K}{1 + (e^{-r_1(z-p_1)})} \right)^t \text{ for all } z < c$$

$$s = \left( \frac{K}{1 + (e^{-r_2(z-p_2)})} \right)^t \text{ for all } z \geq c$$
(1)

where  $z$  is DBH in mm and  $K$  is the upper asymptote of the curves and represents maximum survival (MS), often constant across much of the DBH range,  $r$  is a rate parameter that determines the rate of survival increases ( $r_1$ ) or decreases ( $r_2$ ) with size, and  $p$  is the inflection point, that is, the size at which survival probability

is .5. Subscripts 1 and 2 denote the parameters for the survival curves at sizes below and above the DBH threshold where the curves meet,  $c$ . We set  $c$  to  $0.2 \times \max(\text{DBH})$  to ensure sufficient individuals above and below the DBH threshold. Raising to the power  $t$ , the time in years between censuses, makes parameters describe the annual probability of survival. For all further analysis, we used MS (usually, but not always  $K$ ), and juvenile survival (JS)—the survival probability at 10 mm DBH. Although we refer to this as JS, in some species individuals may already be several decades old and reproductive at 10 mm DBH, the smallest size in the inventory data.

### 2.2.2 | Growth

To account for negative growth in the census data, which may result from measurement error, stem damage or differences in water content, from one census to the next, we adapted the method described by (Rüger et al., 2011). We resampled all stem measurements 1000 times according to the distribution  $sda + sdb \times dbh$ , where  $sda = 0.927$  mm and  $sdb = 0.0038$  mm. If any increments were still negative, we assigned a growth increment by sampling from the positive growth increments in the plot.

We divided individuals of each species at a given plot into the slowest growing 95% of the population and the fastest 5% of the population and fit Gamma distributions to the DBH increment data for each group separately. The Gamma distribution has the flexibility to capture both the highly skewed distribution of slow growth and the more symmetric distribution of faster growth:

$$\Delta z \sim \begin{cases} \text{Gamma}(\alpha_1, \beta_1) & \text{for } \Delta z < q \\ \text{Gamma}(\alpha_2, \beta_2) & \text{for } \Delta z \geq q \end{cases} \quad (2)$$

Delta  $z$  is the absolute annual change in DBH in mm, alpha and beta are the shape and rate parameters of the Gamma distribution, respectively, and  $q$  is the percentile determining the proportion of individuals in the fast growth distribution, here the 95% percentile of DBH increment. Subscripts 1 and 2 denote growth distributions fit to individuals with growth below and above  $q$ , respectively. In further analyses, we used the expectation of growth (in  $\text{mm yr}^{-1}$ ), given by  $\alpha/\beta$  for the slow and fast growth distribution for each species, hereafter 95G and 5G.

We estimated growth and survival parameters using Bayesian Markov Chain Monte Carlo with the *rstan* package (Stan Development Team, 2020) in R (R Core Team, 2021). For each species, we ran three chains with 2500 iterations and checked diagnostic plots for convergence. For further details of the growth and survival models see Needham et al. (2018).

We did not include reproduction in these analyses due to a lack of data at the global scale. In several studies, (e.g., Rüger et al., 2020; Salguero-Gómez et al., 2016), recruitment into a census is used as

a proxy for reproductive effort. However, in our study, which includes all species with a DBH of 1 cm, recruitment into a census does not have a consistent interpretation across species. For example, a 1 cm DBH stem could be a reproductive stem of an adult shrub species, or a non-reproductive juvenile sapling of a canopy species. We, therefore, chose to focus our analysis on growth and survival, which better capture forest biomass and turnover across life-history strategies.

### 2.3 | Principal component analysis

To identify orthogonal axes of demographic variation, we performed a principal component analysis (PCA) on the growth and survival parameters for each species at each plot. We used logistic transformations of MS and JS (to improve normality), and non-transformed 5G and 95G. We also included the maximum DBH for each species, denoted ST. We used the PCA function from the *FactoMineR* R package (Lê et al., 2008) as it allowed us to weight species by the inverse of the number of plots they occur in, preventing species with wider geographic distributions having disproportionate influence on the results (115 species [5.9%] occurred in more than one plot). We did not attempt a phylogenetically controlled PCA due to the lack of well resolved phylogeny for the majority of species in our analysis. Incorrect tree topologies, along with incomplete taxon sampling, would introduce biases into any estimation of a phylogenetic signal in demographic rates (Ackerly, 2000; Symonds, 2002). All PCA axes are shown in Figure S2.

### 2.4 | Demographic diversity across plots

To address question one, we calculated the DD of each plot as the convex hull of species from that plot along the first two dimensions of PCA space, which together explain 76.2% of variation in growth and survival among plots (Cornwell et al., 2006) (Figure 3). We scaled the coordinates of each species along each axis by the percentage variation explained by that axis. We then plotted DD and species richness across plots and fitted a spline through the data using the *ss* function of the *npreg* package in R (Helwig, 2021) (Figure 4). As a measure of the degree of demographic similarity among plots, we calculated the pairwise overlap of convex hulls (Figure S3). To account for different plot sizes, we randomly sampled 500 bootstrap samples of 400 20 m  $\times$  20 m quadrats (equivalent to the size of the smallest plot, 16 ha) from each plot with replacement. For each unique bootstrapped sample, we calculated the species richness and then calculated the convex hull for those species present in the sampled 16 ha. We present here the mean number of species with  $\geq 200$  individuals across samples as species richness, the mean convex hull volume as DD, the median area of overlap from pairwise overlaps of 500 convex hulls at each plot for similarity in DD. In all but one plot (BCI), all species with more than 200 individuals in the full data set were found in all 500 bootstrap

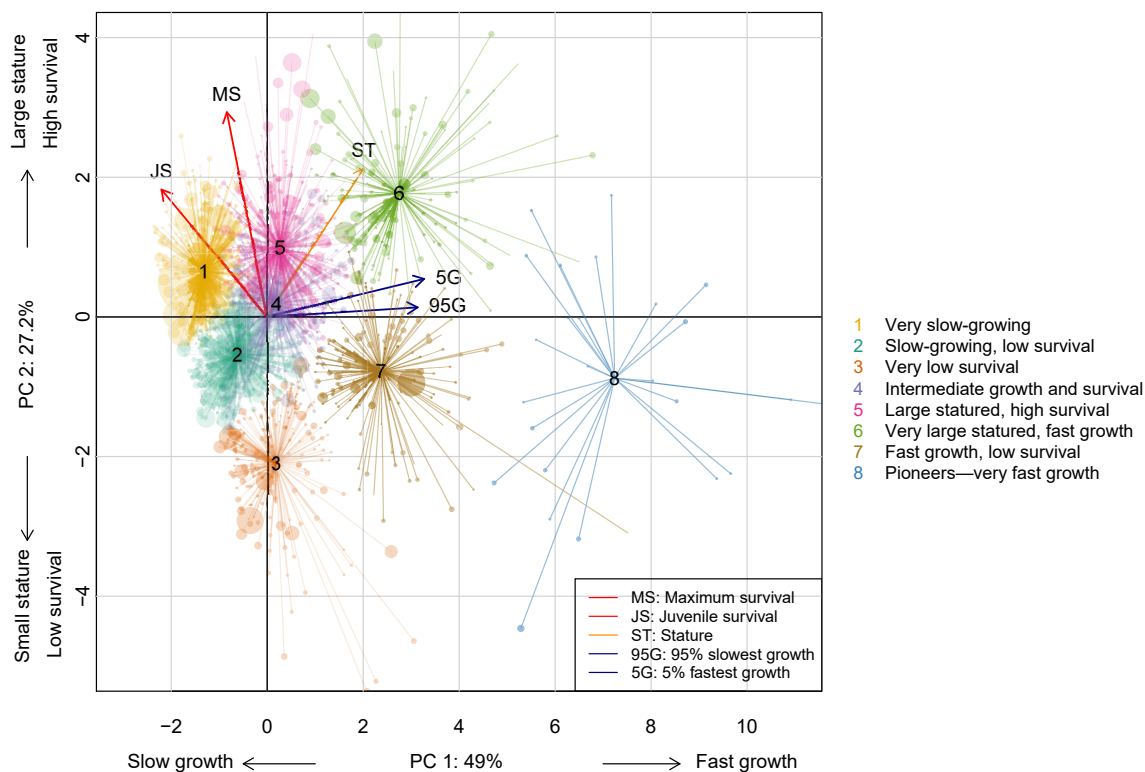
samples. Convex hulls and overlaps were calculated using the geometry package in R (Habel et al., 2019).

## 2.5 | Demographic composition across plots

To address question two, we first clustered species into demographic types, or specifically, growth-survival-stature modes (GSSMs), based on similarities in observed demographic rates. To determine the similarities in demographic rates, we used the position of each species in multidimensional PCA space using the clustering kmeans function in base R (R Core Team, 2021), with 200 starting points. The *kmeans* function minimizes the sum of squares from each species to the center of its assigned GSSM. Using 200 starting points allows the algorithm to explore multiple centers for each GSSM and, thus, optimizes clustering. We tested different combinations of clustering algorithms and cluster numbers (Figures S4 and S5) and identified eight GSSMs that were biologically interpretable (Figure 2) and that met statistical standards in cluster analysis: high Dunn index, low number of negative silhouettes, no cluster with too few species, and high explained variance (Figures S4 and S5). For each plot, we calculated the relative abundance of each GSSM as a measure of DC (Figure S6). We then tested whether DC was correlated with species richness across plots (Figure S7). Our third question was how

are DD and DC related to environmental conditions? We therefore tested whether DD and DC at each plot are predicted by mean annual temperature (MAT) and mean annual precipitation (MAP), taken from Anderson-Teixeira et al. (2015) (Figure 5 and Figures S8 and S9). We did not correct for multiple hypotheses related to the relative abundance of GSSMs and MAT, MAP, and species richness, as we are more interested in the relationship between each GSSM and each variable (see [Cabin & Mitchell, 2000] for discussions of the problems with Type II errors in ecological studies). We present  $R^2$  and p-values but avoid significance testing and discuss inference using the language of “evidence” as discussed in Muff et al. (2021).

To explore differences in life histories across GSSMs, we calculated passage times and life expectancies for each GSSM using a simple individual-based model. For each GSSM we found the mean growth and survival parameters from all species in that GSSM. We simulated 25,000 individuals until all individuals had died. Five percent of individuals grew according to the fast growth distribution, and 95% according to the slow growth distribution (Equation (2)). Survival was a draw from a binomial distribution each year with probabilities given by the size-dependent function described in Equation (1). We calculated life expectancies as the mean time to death for a 10 mm DBH stem, and passage times as the mean time taken to grow from 10 mm DBH to 100 mm DBH (a size that all GSSMs can reach), conditional on survival.



**FIGURE 2** Growth-survival state space. Principal component analysis and clustering analysis on growth and survival parameters for 1961 species across 20 tropical and temperate forests. Parameters are described in detail in the main text. PC 1 corresponds to a trade-off between growth and juvenile survival and accounts for 49.0% of the variation among species. PC 2 accounts for 27.2% of variation and is a stature and survival axis, characterized by high maximum survival and large stature at one extreme and low maximum survival and small stature at the other. We used clustering algorithms to group species into eight “growth-survival-stature modes,” depicted with different colors and given a number from 1 to 8



## 2.6 | Aboveground biomass across plots

Our final question was how do DD and DC vary with AGB and carbon residence across plots? To test this, we estimated AGB and carbon residence time at each plot. We estimated AGB using equations and parameters from Chave et al. (2014) for tropical plots and from Jenkins et al. (2003) and Chojnacky et al. (2014) for temperate plots. We calculated AGB for all stems in each plot using the following equations.

For tropical plots, we calculated AGB as:

$$\text{AGB} = \exp \left[ -1.803 - 0.976E + 0.976 \ln(\text{wsg}) + 2.673 \ln(D) - 0.0299 [\ln(D)]^2 \right] \quad (3)$$

where  $D$  is DBH in cm,  $\text{wsg}$  is wood-specific gravity and  $E$  is a local bioclimatic variable (Chave et al., 2014).  $\text{wsg}$  estimates come from Zanne et al. (2009), see Chave et al. (2009). We use species-level estimates of  $\text{wsg}$  where available, and genus, family or global means depending on availability. Values of  $E$  for each plot come from [http://chave.ups-tlse.fr/pantropical\\_allometry.htm](http://chave.ups-tlse.fr/pantropical_allometry.htm).  $E$  is defined as

$$E = 1. e - 3 \times (0.178 \times TS - 0.938 \times CWD - 6.61 \times PS) \quad (4)$$

where  $TS$  is temperate seasonality as defined in the Worldclim data set (bioclimatic variable 4).  $CWD$  is climatic water deficit and  $PS$  is precipitation seasonality as defined in the Worldclim data set (bioclimatic variable 15). Higher values of  $E$  result in lower estimates of AGB. For more details, see [http://chave.ups-tlse.fr/pantropical\\_allometry.htm](http://chave.ups-tlse.fr/pantropical_allometry.htm).

For temperate plots, we calculated AGB as:

$$\text{AGB} = \exp \left[ \beta_0 + \beta_1 \ln(D) \right] \quad (5)$$

We used parameter values from Chojnacky et al. (2014), which depend on wood specific gravity. Again, these come from Zanne et al. (2009), see Chave et al. (2009).

## 2.7 | Aboveground carbon residence time

We calculated aboveground carbon residence time at each plot as  $\text{AGB}/\text{AGB}_{\text{mort}}$ , where  $\text{AGB}_{\text{mort}}$  is AGB loss from mortality, following Koven et al. (2015). For  $\text{AGB}_{\text{mort}}$  we followed Sullivan et al. (2020), to take into account unobserved growth of trees present at the first census that had died by the second census, and recruits that entered during a census interval but died before being recorded in the second census. The unobserved growth of trees that died, and unobserved recruits were estimated from per area annual recruitment and per capita annual mortality using equations 11 and 5 from Kohyama et al. (2017). For more information, see SI Methods S1. We tested the correlations between AGB and carbon residence time with DD. We further tested for correlations between AGB and carbon residence time with the relative abundance of each GSSM (Figure 6 and Figures S10 and S11). Again, we present tests independently, but follow Moffit et al. and discuss our findings in terms of "evidence."

## 3 | RESULTS

### 3.1 | Axes of demographic variation

The first principal component (PC) accounts for 49% of the variation among species (Table 1) and corresponds to a trade-off between growth and JS (Figure 2, growth rates [both 95G and 5G], as growth and JS rates [JS] have opposite loadings on the first axis). MS is largely orthogonal to growth rates. The positive loadings of both 95G and 5G on PC 1 suggests that species are generally consistently fast-growing or slow-growing, that is, if the fastest 5% of the population (5G) are fast-growing then the slowest 95% of the population (95G) are also relatively fast-growing. PC 2 explains 27.2% of variation and describes an axis associated with low survival and small stature at one extreme and high survival and large stature at the other. Other axes are shown in Figure S2. Species are not evenly distributed across the growth-survival space, but are clustered in regions of high survival and slow growth. GSSM 2, a slow-growing, small-statured and low survival mode, had the most species, 29%, followed by GSSM 1, 23%, a very slow-growing, high-survival mode.

### 3.2 | Demographic diversity shows a complex association with species richness

Our first question asked how DD varies among forests and whether species richness correlates with DD. We estimated DD at each plot by calculating the area of the convex hull around species' positions in PCA space (Figure 3). We find that DD varies widely across plots and initially shows a steep increase with increasing species richness (Figure 4). DD increases from low to intermediate levels of species richness and is roughly level at high-species-richness plots.

### 3.3 | Demographic composition varies across forests

Our second question asked how DC varies across plots. We compared where species from each plot are located in PCA space (Figure 3), the pairwise overlap of plots in PCA space (Figure S3), as well as the relative abundance of GSSMs at each plot (Figure S6). Communities of species from each plot occupy different regions of PCA space (Figure 3) with some clustered in regions of high survival, and others more spread out to include fast growing species. Tropical plots have a higher degree of overlap with other plots, due to higher species richness and larger convex hulls. BCI (Panama), La Planada (Colombia), and other plots with fast growing species have the greatest degree of overlap with other plots (Figure S3). We found moderate evidence for an effect of species richness on the relative abundance of GSSM 6 ( $R_{sq} = 0.29$ ,  $p = .018$ ), but weak or no evidence of species richness having an effect on the relative abundance of any other GSSM (Figure S7). There is little evidence for an effect of species richness on the relative abundance of GSSM 6 among

TABLE 1 Variable loadings, eigenvalues, and percent variance explained by each principal component analysis (PCA) axis

	PC 1	PC 2	PC 3	PC 4	PC 5
MS	-0.24	0.84	0.27	-0.41	-0.01
JS	-0.63	0.52	0.13	0.57	0.01
ST	0.57	0.61	-0.54	0.04	0.07
95G	0.89	0.04	0.36	0.13	0.22
5G	0.93	0.15	0.14	0.12	-0.27
Eigenvalue	2.45	1.36	0.54	0.52	0.13
% variance	48.97	27.22	10.79	10.45	2.56
Cumulative % variance	48.97	76.19	86.98	97.44	100

tropical plots ( $R_{sq} = 0.11$ ,  $p = .24$ ). The relationship between species richness and the relative abundance of GSSM 6 is driven by the inclusion of temperate plots.

### 3.4 | Climate variables predict demographic diversity and composition to some extent

Our third question asked how DD and DC are related to environmental conditions. We find strong evidence that DD increases with increasing MAT ( $R_{sq} = 0.43$ ,  $p = .0022$ ), but this correlation is mostly driven by a clear trend in temperate plots (Figure 5), and, therefore, would be unlikely to confirm a causal relationship from temperature

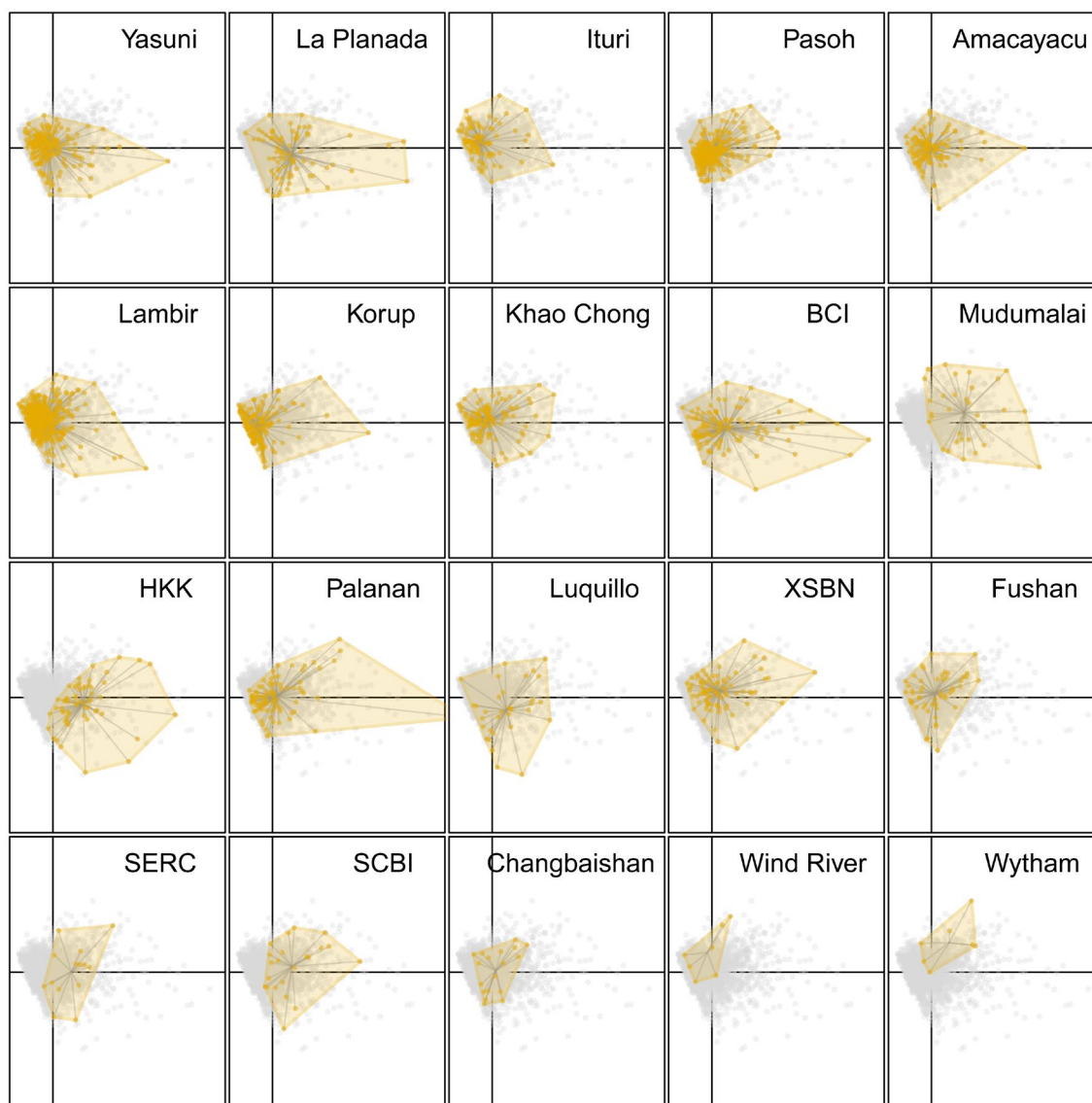
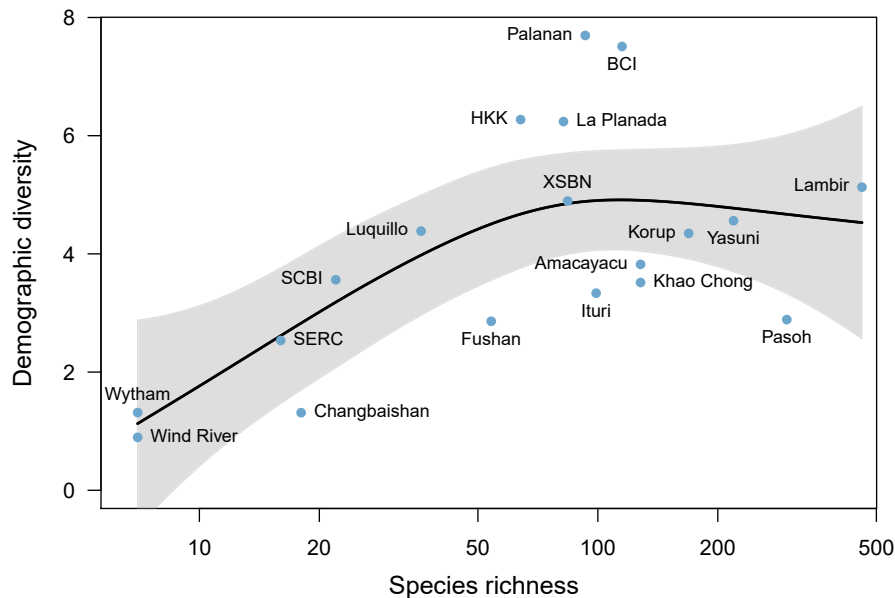


FIGURE 3 Position of plots in growth-survival space. Position of species in the first two dimensions of demographic space for each plot sorted from left to right, top to bottom by latitude (plots on the bottom row are temperate plots). Gray dots show species from all plots; colored dots indicate the species from the named plot, with lines connecting each species to the community centroid. Shading shows the demographic diversity of each plot, calculated as the convex hull area for species from each plot





**FIGURE 4** Relationship between species richness and demographic diversity. Species richness here is the mean number of species with over 200 individuals found in 500 bootstrap samples of 16 ha from each plot. To estimate demographic diversity, we calculated the volume of the convex hull of species along the first two axes of the growth-survival PCA for each bootstrap sample of species richness. Units of demographic diversity thus refer to the PCA axes. We fit a spline through the data using the *ss* function from the *npreg* package in R. Demographic diversity increases from low to moderate levels of species richness and is roughly level at high species richness. Note that the x-axis is logged for visualization purposes

to DD globally. Among tropical plots, there is no evidence for an effect of MAT on DD ( $R_{sq} = 0.002$ ,  $p = .88$ ). We also find no evidence of an effect of MAP on DD ( $R_{sq} = 0.14$ ,  $p = .11$ ). There is strong evidence that the relative abundance of the large statured GSSM 6 decreases with increasing MAT ( $R_{sq} = 0.48$ ,  $p = .00094$ ). Again, this correlation is mostly driven by the inclusion of temperate plots that contain a greater proportion of large statured species, given their lower species diversity. Small-statured, high-survival GSSM 2 has a positive correlation with MAT ( $R_{sq} = 0.32$ ,  $p = .012$ ), as does large statured, fast growing GSSM 7 ( $R_{sq} = 0.2$ ,  $p = .058$ ). There is moderate evidence that GSSM 3 is positively correlated with MAP ( $R_{sq} = 0.28$ ,  $p = .02$ ) and GSSM 6 negatively correlated with MAP ( $R_{sq} = 0.23$ ,  $p = .039$ ). Otherwise, there is weak or no evidence for an effect of MAT or MAP on any other GSSM (Figures S8 and S9).

### 3.5 | Demographic composition, not diversity, predicts forest structure

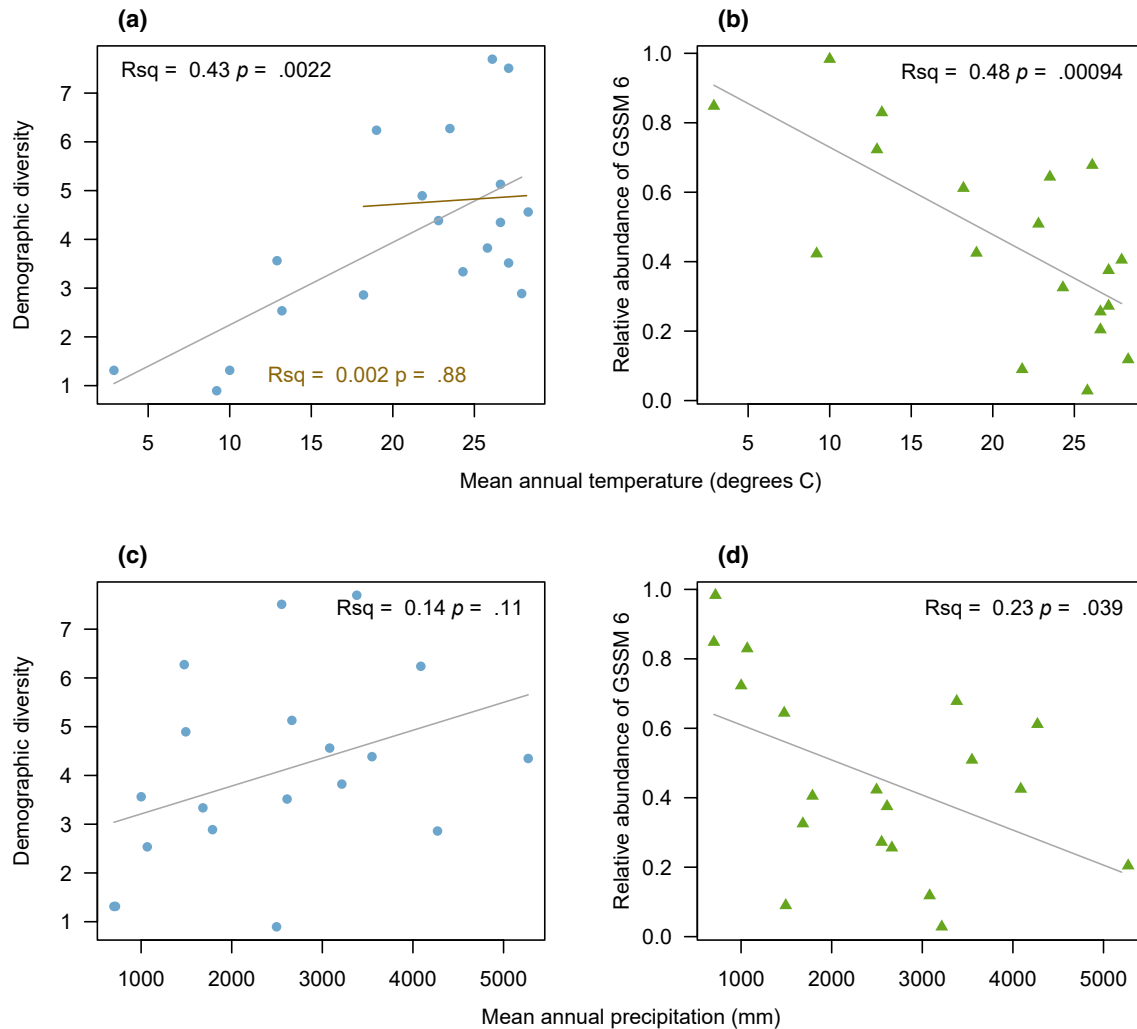
Our final question asked how DD and DC are related to AGB and carbon residence time. We find no evidence that DD is related to either AGB or carbon residence time across plots ( $R_{sq} = 0.014$ ,  $p = .63$ ;  $R_{sq} = 0.15$ ,  $p = .12$ ) (Figure 6). We find strong evidence that the relative abundance of GSSM 5 is positively correlated with AGB ( $R_{sq} = 0.33$ ,  $p = .0095$ ), and moderate evidence that the relative abundance of GSSM 6 is correlated with carbon residence time ( $R_{sq} = 0.23$ ,  $p = .043$ ), even when we remove Wytham Wood, which had a carbon residence time more than double that of any other plot.

## 4 | DISCUSSION

We fitted growth and survival models to 1961 species from 20 temperate and tropical forests and assessed global patterns of DD and DC. While DD initially increases with species richness it levels off in high diversity plots and there is no evidence of a relationship with either AGB or carbon residence time. In contrast, we find strong evidence for a relationship between DC and AGB, specifically for an increase in AGB with an increase in the relative abundance of large statured demographic types, and moderate evidence that DC has an effect on carbon residence time across forests.

### 4.1 | Demographic diversity has a complex association with species richness

Our first question was how does DD vary among forests around the world and are species-rich forests more demographically diverse? We find wide variation in DD across forests, but a complex association between species richness and DD. DD initially increases with increasing species richness but peaks at intermediate levels of species richness and reaches a plateau across highly species rich plots (Figure 4). Condit et al. (2006) similarly found that across ten tropical forests the most species rich forests had the lowest DD. This might be due to the fact that, in species rich tropical forests an increase in species richness does not necessarily translate to an increase in richness at higher taxonomic levels, rather, particular genera are highly species-rich (Davies et al., 2005; Richardson et al., 2001). If so,

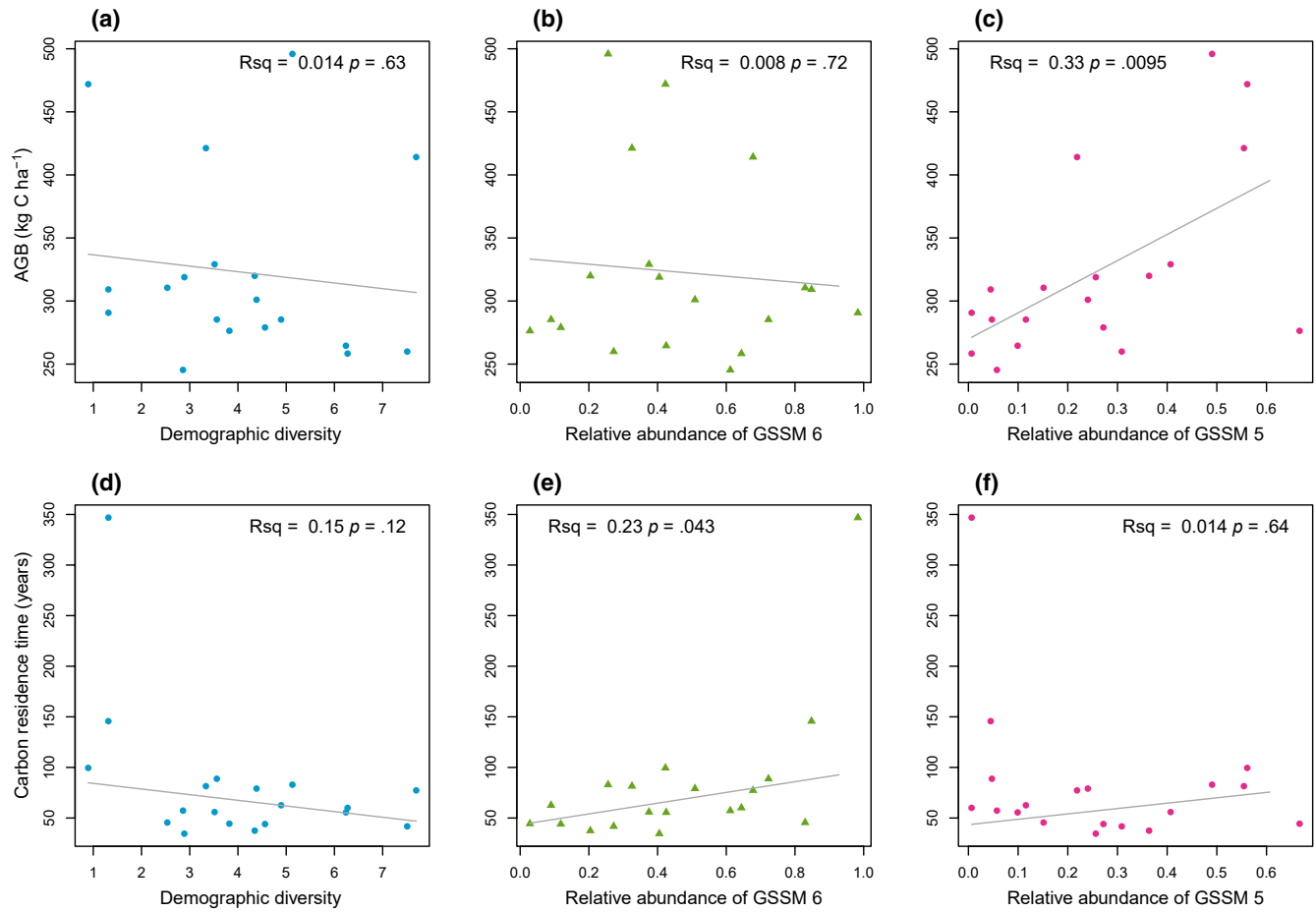


**FIGURE 5** Correlation between climate variables and both demographic diversity and the relative abundance of large statured species. Mean annual temperature (MAT) and mean annual precipitation (MAP) are taken from Anderson-Teixeira et al. (2015). Growth survival stature mode (GSSM) 6 is the demographic type characterized by fast growing large statured species. Demographic diversity (DD) is the volume of the convex hull around species from each plot in the first two dimensions of PCA space. There is strong evidence for DD increasing with increasing MAT ( $Rsq = 0.43$ ,  $p = .002$ ). However, within the tropics, there is no evidence that MAT has an effect on DD ( $Rsq = 0.002$ ,  $p = .88$ ). There is very strong evidence that the relative abundance of GSSM 6 decreases with MAT ( $Rsq = 0.48$ ,  $p = .00095$ ), but weak evidence for an effect of MAP on the relative abundance of GSSM 6 ( $Rsq = 0.23$ ,  $p = .039$ ). For the effect of climate variables on other GSSMs see Figures S8 and S9

this accumulation of species within genera could introduce demographic redundancy; explaining the plateau reached after intermediate levels of species richness. Because demographic rates within many species-rich tropical tree genera tend to be phylogenetically conserved, additional species do not always increase the DD of the forest (Davies et al., 2005). More information on the similarity of demographic rates among closely related species could provide insights into the mechanisms that determine DD across forests.

The relationship between DD and species richness also likely relates to disturbance regimes. Highly disturbed plots in our analysis (e.g., HKK [Thailand], Palanan [Philippines], Fushan [Taiwan], and Luquillo [Puerto Rico]) tend to have lower species richness than less disturbed sites. Yet despite having fewer species, disturbed sites have high DD due to greater numbers of fast-growing species, which can capitalize on increased light availability and nutrients generated

by disturbance (Brokaw, 1987). Since variation in growth rates is larger than variation in survival rates, plots with high disturbance rates have higher DD by our measure because they contain species located on the edge of PCA space towards very fast growth. Examples include HKK (Thailand), a seasonally dry forest with frequent ground fires, and Palanan (Philippines), a cyclone-prone forest. In contrast, plots that have relatively low DD but high species richness, such as Lambir (Malaysia), are dominated by slow-growing, high survival species. This is likely due to exceptionally low soil fertility, which also limits growth (Russo et al., 2005). Our results are in line with findings from Russo et al. (2021) who found that in highly disturbed plots the strength of the growth-survival trade-off is weaker than in relatively stable undisturbed plots because slow-growing, high-survival species are not able to become established (Russo et al., 2021). Quantifying disturbance rates is challenging due



**FIGURE 6** Demographic diversity (DD) versus aboveground biomass (AGB) and carbon residence time. DD is the volume of the convex hull around species from each plot in the first two dimensions of principal component analysis space. In panels (d, e and f) Wytham Woods is excluded from the analysis (but shown in the figure) due to its very high carbon residence time of 347 years. There is no evidence that DD has an effect on either AGB or carbon residence time ( $Rsq = 0.014$ ,  $p = .63$ ;  $Rsq = 0.15$ ,  $p = .12$ ). There is strong evidence that the relative abundance of growth-survival-stature mode (GSSM) 5 has an effect on AGB ( $Rsq = 0.33$ ,  $p = .0095$ ). There is moderate evidence that the relative abundance of GSSM 6 has an effect on carbon residence time ( $Rsq = 0.23$ ,  $p = .043$ )

to the variability in disturbance regimes and the difficulty in quantifying disturbance across many forests. For instance, disturbances affecting forests in this study include elephants, wildfires, cyclones, and windthrows. Estimating disturbance intensity at a given site is challenging, as intensity can vary greatly over small spatial scales due to topographical heterogeneity, wind exposure (Negrón-Juárez et al., 2014) and soil saturation (Margrove et al., 2015). In many cases the best indicator of the severity of a disturbance is mortality rates. Our results suggest that disturbances may be a key driver of demographic differences across sites, making this an important area of future research.

## 4.2 | Demographic composition varies across forests

Because we are interested in the role that demography plays in shaping the AGB and turnover of forests, we also explored differences in DC across plots. We find that despite the diversity of forest

types included in our study, many of the same demographic strategies are present across forests (Figure S6). This may be partially due to physiological constraints on tree life-history. Although tree life-history strategies can be very varied, growth and survival of trees is relatively limited compared with the differences in demographic rates across the whole plant kingdom (Salguero-Gómez et al., 2016). In most forests, there are canopy and understory species with a range of growth and survival rates. Yet, differences in biogeography and environmental conditions alter the relative abundance of demographic types, with consequences for vertical forest structural complexity and the strength of demographic trade-offs (Russo et al., 2021).

One important difference among forests in terms of DC is the relative abundance of large statured species, which is highest in forests with low species richness (Figure S7). This finding is in agreement with previous work across temperate and tropical plots (Lutz et al., 2018) and can be explained by the greater structural complexity of tropical forests. Due to limited space in the canopy, high stem density in tropical forests is normally associated with an

increase in canopy layers and higher numbers of small-statured understory individuals (Bohlman & Pacala, 2012; Farrior et al., 2016). Increased subcanopy layers lead to a decrease in the relative abundance of large canopy trees, even if the absolute abundance remains the same. Why there is such a diversity of subcanopy species in some tropical forests, as opposed to a high stem density of fewer species, remains unanswered, although one hypothesis is that greater niche complementarity in the tropics allows for a higher number of species to coexist (Chesson et al., 2013; Loreau, 1998).

### 4.3 | There is moderate evidence for a relationship between climate variables and demographic diversity and demographic composition

To better understand the mechanisms driving differences in demography across forests, we tested for correlations between climate variables, and both DD and DC. Demographic rates have been shown to correlate with climate variables at the regional and pan-tropical scale (Esquivel-Muelbert et al., 2020; Johnson et al., 2018). Most notably, annual growth rates are slower at higher latitudes with lower temperatures. Although we also find evidence of a correlation between temperature and DD, there is no evidence for such a relationship within the tropics (Figure 5). Survival rates within the tropics decrease with increasing aridity, e.g. (Locosselli et al., 2020), yet we find no evidence of an effect of MAP on DD, possibly because DD is also defined by growth rates and stature in our analysis, which may show little correlation with precipitation. DC is mostly unrelated to temperature and precipitation, with the exception that the relative abundance of large-statured species decreases with increasing MAT. These correlations are likely driven by the strong increase in species richness with MAT, which as noted above, leads to greater structural complexity and a decrease in the relative abundance of large canopy trees. Our results suggest that, although DD and DC show some correlation with climate, the dynamics at a particular plot cannot be predicted by these metrics alone.

### 4.4 | Demographic composition, not diversity, predicts forest structure

In answer to our fourth question, we find evidence that DC, but not DD, is related to AGB and carbon residence time (Figure 6). Despite species richness varying by nearly 60 times, the two highest biomass plots in our study are Lambir, a dipterocarp forest in Malaysian Borneo, and Wind River, a temperate forest in the Pacific Northwest of the United States that is dominated by *Tsuga heterophylla* (western hemlock), and *Pseudotsuga menziesii* (Douglas-fir). Our results highlight that there are multiple ways to be a high biomass forest, from species-rich, tropical forests, to temperate forests dominated by just a handful of species. Both Lambir and Wind River are dominated by species in GSSMs 5 and 6, demographic types characterized by large stature. In Lambir just 26% of individuals, but 75% of AGB is in

GSSMs 5 and 6, whereas in Wind River 78% of individuals and 98% of AGB is in GSSMs 5 and 6. Our results support previous findings that the maximum height of dominant species, rather than a diversity of functional traits, determines biomass across forests (Finegan et al., 2015; Van Pelt et al., 2016). Our results suggest that species identity, rather than species richness or DD, determines carbon dynamics.

Aboveground carbon residence time, here defined as the length of time that carbon remains in living, aboveground tissue of trees, is strongly influenced by the growth and mortality of individual trees. Together growth and survival rates determine how quickly carbon is accumulated and how long that carbon is stored. Differences in growth and survival rates among GSSMs results in passage times that range from means of 15 to 151 years, and life expectancies that range from mean of 17 to 87 years (Figure S13). Despite small stature, GSSM 1 had the longest mean life expectancy (87 years) due to the slow growth and high survival of species in this mode. This finding is in line with previous studies that have shown that understory trees make substantial contributions to carbon storage in African forests (Hubau et al., 2019). Disturbances lead to higher mortality and thus decrease carbon residence time (e.g. Gora et al., 2020). Increases in disturbance frequency could directly decrease carbon residence time through short-term increases in mortality but also indirectly through shifts in composition towards demographic types with faster growth rates and higher mortality.

## 5 | CONCLUSIONS AND FUTURE DIRECTIONS

Here we have presented a quantitative analysis of the relationship between species richness, demography, and forest structure and turnover. Although we find little evidence for an effect of DD on AGB or carbon residence time, an important question that remains unanswered is whether forests that are more demographically diverse will show greater resilience to future perturbations. Many combinations of functional traits can give rise to the same demographic rates. As a result, species with similar demographic rates may respond differently to disturbance, and a forest with a high degree of demographic redundancy could be buffered from disturbance if some of the species are themselves resilient, or the mixture of species confers resilience. As disturbances are predicted to increase in the future (McDowell et al., 2020), it is important to explore the extent to which DD and DC determine forest response to changes in disturbance frequency and intensity. This may be especially important with regards to large trees. Our results highlight the importance of large-statured trees for forest ecosystem function across temperate and tropical regions. Shifts in DC following changes to disturbance regimes could have significant consequences for AGB and carbon residence time if they result in a decrease in the abundance of large statured demographic types. An improved understanding of the determinants of DC across forests will be essential to predict forest response to future climates and disturbance regimes.

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## CONFLICT OF INTEREST

The authors declare no conflict of interest.

## DATA AVAILABILITY STATEMENT

Growth and survival parameters for each species, excluding species names, along with R scripts for analyzing the data and producing the figures in the paper are available from Jessica Needham's GitHub at [https://github.com/JessicaNeedham/ForestGEO\\_global\\_demography](https://github.com/JessicaNeedham/ForestGEO_global_demography) or from the ForestGEO GitHub at [https://github.com/forestgeo/Needham\\_et\\_al\\_2022\\_GCB](https://github.com/forestgeo/Needham_et_al_2022_GCB). The data for individual sites in this study are available from the relevant authors or through the ForestGEO website <https://www.forestgeo.si.edu> upon reasonable request.

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## REFERENCES

- Ackerly, D. D. (2000). Taxon sampling, correlated evolution, and independent contrasts. *Evolution*, 54(5), 1480–1492. <https://doi.org/10.1111/j.0014-3820.2000.tb00694.x>
- Anderson-Teixeira, K. J., Davies, S. J., Bennett, A. C., Gonzalez-Akre, E. B., Muller-Landau, H. C., Joseph Wright, S., Abu Salim, K., Almeyda Zambrano, A. M., Alonso, A., Baltzer, J. L., Basset, Y., Bourg, N. A., Broadbent, E. N., Brockelman, W. Y., Bunyavejchewin, S., Burslem, D. F. R. P., Butt, N., Cao, M., Cardenas, D., ... Zimmerman, J. (2015). CTFS-ForestGEO: A worldwide network monitoring forests in an era of global change. *Global Change Biology*, 21(2), 528–549. <https://doi.org/10.1111/gcb.12712>
- Beech, E., Rivers, M., Oldfield, S., & Smith, P. P. (2017). GlobalTreeSearch: The first complete global database of tree species and country distributions. *Journal of Sustainable Forestry*, 36(5), 454–489. <https://doi.org/10.1080/10549811.2017.1310049>
- Bohman, S., & Pacala, S. (2012). A forest structure model that determines crown layers and partitions growth and mortality rates for landscape-scale applications of tropical forests. *Journal of Ecology*, 100(2), 508–518.
- Brokaw, N. V. L. (1987). Gap-phase regeneration of three pioneer tree species in a tropical forest. *Journal of Ecology*, 75(1), 9. <https://doi.org/10.2307/2260533>
- Cabin, R. J., & Mitchell, R. (2000). To Bonferroni or not to Bonferroni: When and how are the questions. *Bulletin of the Ecological Society of America*, 81(3), 246–248.
- Chambers, J. Q., Higuchi, N., & Schimel, J. P. (1998). Ancient trees in Amazonia. *Nature*, 391(6663), 135–136.
- Chave, J., Coomes, D., Jansen, S., Lewis, S. L., Swenson, N. G., & Zanne, A. E. (2009). Towards a worldwide wood economics spectrum. *Ecology Letters*, 12(4), 351–366.
- Chave, J., Réjou-Méchain, M., Búrquez, A., Chidumayo, E., Colgan, M. S., Delitti, W. B. C., Duque, A., Eid, T., Fearnside, P. M., Goodman, R. C., Henry, M., Martínez-Yrizar, A., Mugasha, W. A., Muller-Landau, H. C., Mencuccini, M., Nelson, B. W., Ngomanda, A., Nogueira, E. M., Ortiz-Malavassi, E., ... Vieilledent, G. (2014). Improved allometric models to estimate the aboveground biomass of tropical trees. *Global Change Biology*, 20(10), 3177–3190. <https://doi.org/10.1111/gcb.12629>
- Chesson, P., Pacala, S., & Neuhauser, C. (2013). *Environmental niches and ecosystem functioning*, Vol. 10. Princeton University Press.
- Chojnacky, D. C., Heath, L. S., & Jenkins, J. C. (2014). Updated generalized biomass equations for North American tree species. *Forestry*, 87(1), 129–151. <https://doi.org/10.1093/forestry/cpt053>
- Comita, L. S., Muller-Landau, H. C., Aguilar, S., & Hubbell, S. P. (2010). Asymmetric density dependence shapes species abundances in a tropical tree community. *Science*, 329(5989), 330–332.
- Condit, R. (1998). The CTFS and the standardization of methodology. *Tropical forest census plots* (pp. 3–7). Springer Berlin Heidelberg.
- Condit, R., Ashton, P., Bunyavejchewin, S., Dattaraja, H. S., Davies, S., Esufali, S., Ewango, C., Robin, F., Gunatilleke, I. A. U. N., Gunatilleke, C. V. S., Hall, P., Harms, K. E., Hart, T., Hernandez, C., Hubbell, S., Itoh, A., Kiratiprayoon, S., LaFrankie, J., deLao, S. L., ... Zillio, T. (2006). The importance of demographic niches to tree diversity. *Science*, 313(5783), 98–101.
- Condit, R., Hubbell, S. P., & Foster, R. B. (1994). Density dependence in two understory tree species in a neotropical forest. *Ecology*, 75(3), 671–680. <https://doi.org/10.2307/1941725>
- Cornwell, W. K., Schwilk, L. D. W., & Ackerly, D. D. (2006). A trait-based test for habitat filtering: Convex hull volume. *Ecology*, 87(6), 1465–1471.
- Davies, S. J., Abiem, I., Abu Salim, K., Aguilar, S., Allen, D., Alfonso, A., Kristina, A.-T., Ana, A., Gabriel, A., Ashton, P. S., Baker, P. J., Baker, M. E., Baltzer, J. L., Yves, B., Pulchérie, B., Stephanie,



- B., Bourg, N. A., Brockelman, W. Y., Sarayudh, B., ... Daniel, Z. (2021). ForestGEO: Understanding forest diversity and dynamics through a global observatory network. *Biological Conservation*, 253, 108907. <https://doi.org/10.1016/j.biocon.2020.108907>
- Davies, S. J., Tan, S., LaFrankie, J. V., & Potts, M. D. (2005). Soil-related floristic variation in a hyperdiverse dipterocarp forest. In D. W. Roubik, S. Sakai, & A. A. Hamid Karim (Eds.), *Pollination ecology and the rain forest* (pp. 22–34). Springer-Verlag.
- Esquivel-Muelbert, A., Phillips, O. L., Brienen, R. J. W., Fauset, S., Sullivan, M. J. P., Baker, T. R., Kuo-Jung, C., Feldpausch, T. R., Emanuel, G., Niro, H., Jeanne, H.-D., Jon, L., Haiyan, L., Yadvinder, M., Beatriz, M., Hur, M. J. B., Abel, M.-M., Lourens, P., Marcos, S., ... David, G. (2020). Tree mode of death and mortality risk factors across Amazon forests. *Nature Communications*, 11(1). <https://doi.org/10.1038/s41467-020-18996-3>
- Farrior, C. E., Bohlman, S. A., Hubbell, S., & Pacala, S. W. (2016). Dominance of the suppressed: Power-law size structure in tropical forests. *Science*, 351(6269), 155–157. <https://doi.org/10.1126/science.aad0592>
- Finegan, B., Peña-Claros, M., de Oliveira, A., Ascarrunz, N., Bret-Harte, M. S., Geovana, C.-R., Fernando, C., Sandra, D., Paul, E. V., Fernando, F., Carlos, L. J., Leda, L., Beatriz, S. N., Marcel, V., & Lourens, P. (2015). Does functional trait diversity predict above-ground biomass and productivity of tropical forests? Testing three alternative hypotheses. *Journal of Ecology*, 103(1), 191–201. <https://doi.org/10.1111/1365-2745.12346>
- Gloor, M., Brienen, R. J. W., Galbraith, D., Feldpausch, T. R., Schöngart, J., Guyot, J.-L., Espinoza, J. C., Lloyd, J., & Phillips, O. L. (2013). Intensification of the Amazon hydrological cycle over the last two decades. *Geophysical Research Letters*, 40(9), 1729–1733. <https://doi.org/10.1002/grl.50377>
- Gora, E. M., Burchfield, J. C., Muller-Landau, H. C., Bitzer, P. M., & Yanoviak, S. P. (2020). Pantropical geography of lightning-caused disturbance and its implications for tropical forests. *Global Change Biology*, 26(9), 5017–5026.
- Grime, J. P. (1998). Benefits of plant diversity to ecosystems: Immediate, filter and founder effects. *Journal of Ecology*, 86(6), 902–910. <https://doi.org/10.1046/j.1365-2745.1998.00306.x>
- Habel, K., Grasman, R., Gramacy, R. B., Mozharovskiy, P., & Sterratt, D. C. (2019). Geometry: Mesh generation and surface tessellation. R package version 0.4.5.
- Helwig, N. E. (2021). npreg: Nonparametric regression via smoothing splines. R package version 1.0-6. CRAN R project.
- Hubau, W., De Mil, T., Van den Bulcke, J., Phillips, O. L., Angoboy Ilondea, B., Joris, V. A., Sullivan Martin, J. P., Laurent, N., Benjamin, T., Camille, C., Banin, L. F., Begne, S. K., Baker, T. R., Nils, B., Eric, C., Clark, C. J., Murray, C., Comiskey, J. A., Aida, C.-S., ... Hans, B. (2019). The persistence of carbon in the African forest understory. *Nature Plants*, 5(2), 133–140. <https://doi.org/10.1038/s41477-018-0316-5>
- Jenkins, J. C., Chojnacky, D. C., Heath, L. S., & Birdsey, R. A. (2003). National scale biomass estimators for United States tree species. *Forest Science*, 49, 12–35.
- Johnson, D. J., Needham, J., Xu, C., Massoud, E. C., Davies, S. J., Anderson-Teixeira, K. J., Sarayudh, B., Chambers, J. Q., Chia-Hao, C.-Y., Jyh-Min, C., Chuyong, G. B., Richard, C., Susan, C., Christine, F., Giardina, C. P., Giambelluca, T. W., Nimal, G., Savitri, G., Chang-Fu, H., ... McMahon, S. M. (2018). Climate sensitive size-dependent survival in tropical trees. *Nature Ecology & Evolution*, 2(9), 1436–1442. <https://doi.org/10.1038/s41559-018-0626-z>
- Keenan, T. F., & Williams, C. A. (2018). The terrestrial carbon sink. *Annual Review of Environment and Resources*, 43(1), 219–243.
- Koch, G. W., Sillett, S. C., Jennings, G. M., & Davis, S. D. (2004). The limits to tree height. *Nature*, 428(6985), 851–854.
- Kohyama, T. S., Kohyama, T. I., & Sheil, D. (2017). Definition and estimation of vital rates from repeated censuses: Choices, comparisons and bias corrections focusing on trees. *Methods in Ecology and Evolution*, 9(4), 809–821.
- Koven, C. D., Chambers, J. Q., Georgiou, K., Knox, R., Negron-Juarez, R., Riley, W. J., Arora, V. K., Brovkin, V., Friedlingstein, P., & Jones, C. D. (2015). Controls on terrestrial carbon feedbacks by productivity versus turnover in the CMIP5 Earth System Models. *Biogeosciences*, 12(17), 5211–5228. <https://doi.org/10.5194/bg-12-5211-2015>
- Lê, S., Josse, J., & Husson, F. (2008). FactoMineR: An R package for multivariate analysis. *Journal of Statistical Software*, 25(1).
- Locosselli, G. M., Brienen, R. J. W., De Souza Leite, M., Gloor, M., Krotenthaler, S., de Oliveira, A. A., Jonathan, B., Dieter, A., Gregorio, C., & Jochen, S. (2020). Global tree-ring analysis reveals rapid decrease in tropical tree longevity with temperature. *Proceedings of the National Academy of Sciences of the United States of America*, 117(52), 33358–33364. <https://doi.org/10.1073/pnas.2003873117>
- Loreau, M. (1998). Biodiversity and ecosystem functioning: A mechanistic model. *Proceedings of the National Academy of Sciences of the United States of America*, 95(10), 5632–5636. <https://doi.org/10.1073/pnas.95.10.5632>
- Lutz, J. A., Furniss, T. J., Johnson, D. J., Davies, S. J., Allen, D., Alfonso, A., Anderson-Teixeira, K. J., Ana, A., Jennifer, B., Becker Kendall, M. L., Blomdahl, E. M., Bourg, N. A., Sarayudh, B., Burslem David, F. R. P., Alina, C. C., Ke, C., Min, C., Dairon, C., Li-Wan, C., ... Andrew, K. (2018). Global importance of large-diameter trees. *Global Ecology and Biogeography*, 27(7), 849–864. <https://doi.org/10.1111/geb.12747>
- Margrove, J. A., Burslem, D. F. R. P., Ghazoul, J., Khoo, E., Kettle, C. J., & Maycock, C. R. (2015). Impacts of an extreme precipitation event on dipterocarp mortality and habitat filtering in a bornean tropical rain forest. *Biotropica*, 47(1), 66–76. <https://doi.org/10.1111/btp.12189>
- McDowell, N. G., Allen, C. D., Anderson-Teixeira, K., Aukema, B. H., Bond-Lamberty, B., Chini, L., Clark, J. S., Dietze, M., Grossiord, C., Hanbury-Brown, A., Hurr, G. C., Jackson, R. B., Johnson, D. J., Kueppers, L., Lichstein, J. W., Ogle, K., Poulter, B., Pugh, T. A. M., Seidl, R., ... Xu, C. (2020). Pervasive shifts in forest dynamics in a changing world. *Science*, 368(6494). <https://doi.org/10.1126/science.aaz9463>
- Muff, S., Nilsen, E. B., O'Hara, R. B., & Nater, C. R. (2021). Rewriting results sections in the language of evidence. *Trends in Ecology & Evolution*.
- Needham, J., Merow, C., Chang-Yang, C.-H., Caswell, H., & McMahon, S. M. (2018). Inferring forest fate from demographic data: From vital rates to population dynamic models. *Proceedings of the Royal Society B: Biological Sciences*, 285(1874). <https://doi.org/10.1098/rspb.2017.2050>
- Negrón-Juárez, R., Chambers, J., Hurr, G., Annane, B., Cocke, S., Powell, M., Stott, M., Goosem, S., Metcalfe, D., & Saatchi, S. (2014). Remote sensing assessment of forest disturbance across complex mountainous terrain: The pattern and severity of impacts of tropical cyclone Yasi on Australian rainforests. *Remote Sens (Basel)*, 6(6), 5633–5649. <https://doi.org/10.3390/rs6065633>
- Ordoñez, J. C., van Bodegom, P. M., Witte, J.-P. M., Wright, I. J., Reich, P. B., & Aerts, R. (2009). A global study of relationships between leaf traits, climate and soil measures of nutrient fertility. *Global Ecology and Biogeography*, 18(2), 137–149. <https://doi.org/10.1111/j.1466-8238.2008.00441.x>
- Phillips, O. L., Vásquez, M. R., Arroyo, L., Baker, T. R., Killeen, T., Lewis, S. L., Malhi, Y., Monteagudo, M. A., Neill, D., Núñez, V. P., Alexiades, M., Cerón, C., Di Fiore, A., Erwin, T., Jardim, A., Palacios, W., Saldias, M., & Vinceti, B. (2002). Increasing dominance of large lianas in Amazonian forests. *Nature*, 418(6899), 770–774. <https://doi.org/10.1038/nature00926>
- R Core Team. (2021). R: A language and environment for statistical computing. R Foundation for Statistical Computing. <https://www.R-project.org/>
- Richardson, J. E., Pennington, R. T., Pennington, T. D., & Hollingsworth, P. M. (2001). Rapid diversification of a species-rich genus of neotropical rain forest trees. *Science*, 293(5538), 2242–2245.
- Rüger, N., Berger, U., Hubbell, S. P., Vieilledent, G., & Condit, R. (2011). Growth strategies of tropical tree species: Disentangling light and



- size effects. *PLoS One*, 6(9), e25330. <https://doi.org/10.1371/journal.pone.0025330>
- Rüger, N., Comita, L. S., Condit, R., Purves, D., Rosenbaum, B., Visser, M. D., Wright, S. J., & Wirth, C. (2018). Beyond the fast-slow continuum: Demographic dimensions structuring a tropical tree community. *Ecology Letters*, 21(7), 1075–1084. <https://doi.org/10.1111/ele.12974>
- Rüger, N., Condit, R., Dent, D. H., DeWalt, S. J., Hubbell, S. P., Lichstein, J. W., Lopez, O. R., Christian, W., & Farrior Caroline, E. (2020). Demographic trade-offs predict tropical forest dynamics. *Science*, 368(6487), 165–168. <https://doi.org/10.1126/science.aaz4797>
- Russo, S. E., Davies, S. J., King, D. A., & Tan, S. (2005). Soil-related performance variation and distributions of tree species in a Bornean rain forest. *Journal of Ecology*, 93(5), 879–889. <https://doi.org/10.1111/j.1365-2745.2005.01030.x>
- Russo, S. E., McMahon, S. M., Detto, M., Ledder, G., Wright, S. J., Condit, R. S., Davies, S. J., Ashton, P. S., Sarayudh, B., Chia-Hao, C.-Y., Sisira, E., Ewango Corneille, E. N., Christine, F., Foster, R. B., Savi, G. C. V., Nimal, G. I. A. U., Terese, H., Chang-Fu, H., Hubbell, S. P., ... Zimmerman Jess, K. (2021). The interspecific growth-mortality trade-off is not a general framework for tropical forest community structure. *Nature Ecology & Evolution*, 5(2), 174–183. <https://doi.org/10.1038/s41559-020-01340-9>
- Salguero-Gómez, R., Jones, O. R., Jongejans, E., Blomberg, S. P., Hodgson, D. J., Cyril, M.-A., Zuidema, P. A., de Kroon, H., & Buckley Yvonne, M. (2016). Fast-slow continuum and reproductive strategies structure plant life-history variation worldwide. *Proceedings of the National Academy of Sciences of the United States of America*, 113(1), 230–235. <https://doi.org/10.1073/pnas.1506215112>
- Sillett, S. C., Van Pelt, R., Koch, G. W., Ambrose, A. R., Carroll, A. L., Antoine, M. E., & Mifsud, B. M. (2010). Increasing wood production through old age in tall trees. *Forest Ecology and Management*, 259(5), 976–994. <https://doi.org/10.1016/j.foreco.2009.12.003>
- Stan Development Team. (2020). RStan: The R interface to Stan. R package version 2.19.3. <http://mc-stan.org/>
- Sullivan, M. J. P., Lewis, S. L., Affum-Baffoe, K., Castilho, C., Costa, F., Cuni, S. A., Ewango Corneille, E. N., Wannes, H., Beatriz, M., Abel, M.-M., Lan, Q., Bonaventure, S., Vasquez, M. R., Baker, T. R., Brienen Roel, J. W., Feldpausch, T. R., David, G., Manuel, G., Yadvinder, M., ... Phillips Oliver, L. (2020). Long-term thermal sensitivity of earth's tropical forests. *Science*, 368(6493), 869–874. <https://doi.org/10.1126/science.aaw7578>
- Symonds, M. R. E. (2002). The effects of topological inaccuracy in evolutionary trees on the phylogenetic comparative method of independent contrasts. *Systematic Biology*, 51(4), 541–553.
- Tilman, D., Lehman, C. L., & Thomson, K. T. (1997). Plant diversity and ecosystem productivity: Theoretical considerations. *Proceedings of the National Academy of Sciences of the United States of America*, 94(5), 1857–1861. <https://doi.org/10.1073/pnas.94.5.1857>
- Van Pelt, R., Sillett, S. C., Kruse, W. A., Freund, J. A., & Kramer, R. D. (2016). Emergent crowns and light-use complementarity lead to global maximum biomass and leaf area in *Sequoia sempervirens* forests. *Forest Ecology and Management*, 375, 279–308. <https://doi.org/10.1016/j.foreco.2016.05.018>
- Wright, S. J., Kitajima, K., Kraft, N. J. B., Reich, P. B., Wright, I. J., Bunker, D. E., Condit, R., Dalling, J. W., Davies, S. J., Díaz, S., Engelbrecht, B. M. J., Harms, K. E., Hubbell, S. P., Marks, C. O., Ruiz-Jaen, M. C., Salvador, C. M., & Zanne, A. E. (2010). Functional traits and the growth-mortality trade-off in tropical trees. *Ecology*, 91(12), 3664–3674. <https://doi.org/10.1890/09-2335.1>
- Zanne, A. E., Lopez-Gonzalez, G., Coomes, D. A., Ilic, J., Jansen, S., Lewis, S. L., Miler, R. B., Swenson, N. G., Wiemann, C., & Chave, J. (2009). Data from: Towards a worldwide wood economics spectrum. *Dryad Digital Repository*.

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