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Tropical tree growth sensitivity to climate is driven by species intrinsic growth rate and leaf traits

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Abstract
A better understanding of how climate affects growth in tree species is essential for improved predictions of forest dynamics under climate change. Long-term climate averages (mean climate) drive spatial variations in species' baseline growth rates, whereas deviations from these averages over time (anomalies) can create growth variation around the local baseline. However, the rarity of long-term tree census data spanning climatic gradients has so far limited our understanding of their respective role, especially in tropical systems. Furthermore, tree growth sensitivity to climate is likely to vary widely among species, and the ecological strategies underlying these differences remain poorly understood. Here, we utilize an exceptional dataset of 49 years of growth data for 509 tree species across 23 tropical rainforest plots along a climatic gradient to examine how multiannual tree growth responds to both climate means and anomalies, and how species' functional traits mediate these growth
responses to climate. We show that anomalous increases in atmospheric evaporative demand and solar radiation consistently reduced tree growth. Drier forests and fast-growing species were more sensitive to water stress anomalies. In addition, species traits related to water use and photosynthesis partly explained differences in growth sensitivity to both climate means and anomalies. Our study demonstrates that both climate means and anomalies shape tree growth in tropical forests and that species traits can provide insights into understanding these demographic responses to climate change, offering a promising way forward to forecast tropical forest dynamics under different climate trajectories.

**KEYWORDS**
climatic anomalies, climate change, demography, fast-slow continuum, functional traits, photosynthesis, tree vital rates, tropical moist forest ecology, vapour pressure deficit (VPD), water use efficiency

## 1 | INTRODUCTION

Tropical forests are key contributors to global carbon sequestration (Needham et al., 2018; Pan et al., 2011), but climate change may reduce this important ecosystem service by suppressing tree growth or increasing mortality, particularly in tropical forests (Brodribb et al., 2020; Sullivan et al., 2020). Therefore, it is important to understand how climate influences tree growth, both through long-term local averages (hereafter, ‘mean climate’, calculated over a period of 30 years in climate science) and deviations from these averages in a given location over a given time period (hereafter, anomalies) (Harris et al., 2018; Jentsch et al., 2007; Malhi et al., 2009; Rifai et al., 2018, 2019). Long-term mean climate can constrain the ways species achieve different baseline growth rates (hereafter intrinsic growth rates) in different locations through their effect on tree physiological processes (Green et al., 2019; Rifai et al., 2018; Sullivan et al., 2020), while climate anomalies can also drive growth rate deviations from the local baseline at a range of temporal scales, including days (Zwiefel et al., 2021), months (Clark, 2003; Mendivelso et al., 2014; Rifai et al., 2018, 2019), and multiple years (Coomes et al., 2014; Mendivelso et al., 2014; Rohner et al., 2018; Uriarte et al., 2016; Yuan et al., 2019). The multiannual scale of growth variation is of particular interest in the context of climate change; growth changes over long periods that include sustained periods of constraining climatic anomalies (with less opportunity for recovery) likely reflect tree performance responses that impact long-term forest dynamics with climate change (Harris et al., 2018; Jentsch et al., 2007; Sanginés de Cárcer et al., 2018; Yuan et al., 2019). Evidence of such multiannual climate anomaly effects include growth reduction in tropical dry forests following high dry season atmospheric water demand (Mendivelso et al., 2014; Uriarte et al., 2016), or the negative effect of high water deficit on stem net primary productivity (Rifai et al., 2018).

Major difficulties to disentangle the effects of mean climate and anomalies on spatial and temporal growth variations across species in wet tropical forests include the high species diversity in this biome, the common lack of annual growth rings (Brienen et al., 2016) and the potentially wide range of species growth sensitivities to climate (e.g. DeSoto et al., 2020; Mendivelso et al., 2014; Sanginés de Cárcer et al., 2018; Uriarte et al., 2016). Addressing these challenges requires simultaneously studying multiple sites with different baseline mean climates that share common species. Consequently, we still have a limited understanding of the mechanisms responsible for species differences in their growth sensitivity to climate means and anomalies. Functional traits (sensu Violle et al., 2007) can capture species differences in ecological strategies and allocation tradeoffs to growth, survival and reproduction (McGill et al., 2006; Westoby et al., 2002), and can offer a path towards a more mechanistic insight into the ecological strategies underlying tree growth response to climate drivers (Brodribb et al., 2020; Laughlin et al., 2020; Uriarte et al., 2016; Wagner et al., 2014; Zambrano et al., 2017). Specifically, the ‘fast-slow’ plant economics spectrum links fast-growing and slow-growing species to acquisitive and conservative trait values, respectively (Reich, 2014). As high intrinsic growth rates may come with a cost of lower stress tolerance (Gilbert et al., 2016; Reich, 2014), acquisitive strategies could be associated with increased growth sensitivity to climate anomalies, while conservative strategies could attenuate it. Physiological traits directly related to photosynthesis and water use efficiency, in particular, are good candidates to reflect the effects of light- and water-related climate variables on tree growth and forest dynamics (Brodribb et al., 2020; Powers et al., 2020; Rowland et al., 2021; Wagner et al., 2014).

Further, where species perform better (e.g., grow faster) across a range of mean climatic conditions may also underlie different ecological strategies and help predict expected performance sensitivities to climatic anomalies (e.g., drier periods). However, the sign of this relation remains unclear, as species with long-term adaptations to a constraining climatic condition (e.g., drier mean climate) may be either better adapted or closer to a threshold with respect to further deviations (Aguirre-Gutiérrez et al., 2019, 2020; Bennett et al., 2021; Esquivel-Muelbert et al., 2020; Zuleta et al., 2017). A related uncertainty when scaling up from the species to the forest level is whether
growth sensitivity to climate anomalies varies overall depending on local mean climate. Recent works point to additive or interacting effects between mean climate or climate trends and climate anomalies of increasing magnitude and intensity on ecosystem functions and organisms responses to climate change (Aguirre-Gutiérrez et al., 2019, 2020; Harris et al., 2018; Sullivan et al., 2020), but whether the productivity of tropical forests is more at risk in relatively drier or warmer mean conditions remains little understood, in the face of a further increase in temperature and atmospheric water demand related to global warming (Grossiord et al., 2020; Harris et al., 2018; Sullivan et al., 2020; Yuan et al., 2019).

Here, we take advantage of a unique 49-year dataset of regularly censused tropical tree growth (two- to five-year intervals) spanning 509 species across 23 plots covering an elevation range of 1200 m and encompassing a broad range of climatic conditions, in North Queensland (Wet Tropics of Australia). We additionally use 15 morphological, chemical, and physiological traits related to leaf, wood, and maximum size collected within the plot network for 75 dominant species to test how these traits mediate species growth responses to climate drivers. We couple the multi-year census data with the detailed plant traits dataset in Bayesian hierarchical models to relate tree growth to species traits, forest plots, and climate (Figure 1). We examine the effects of both mean climate and climate anomalies on interannual tree growth variation, both within and across species, and evaluate the role of functional traits in capturing species differences in growth sensitivity. We also test whether the effects of climate anomalies on plot-level growth rate variation depend upon long-term mean climate. Specifically, we ask:

**FIGURE 1** Spatial and temporal dimensions of the tropical forest network. (a) Maps of North Queensland (Australia) and the 23 forest plots on a background of the long-term mean annual precipitation for woody vegetation areas. Circles: plots; Circle colours: Plot elevation (strongly negatively correlated to mean annual temperature, Table S3a). (b, c) Illustration of the temporal extent of the study and of the concepts of mean climate and anomalies for one plot (Mont Haig) presenting vapour pressure deficit (VPD) and solar radiation (SRAD) through time, respectively. (b, c) Show the mean climate (1981–2010) (horizontal black dashed line) and negative and positive anomalies (blue and red vertical segments and dots; monthly anomalies averaged per year). VPD and SRAD were modelled as a plot-specific function of year (see Section 2 and Table S4). The thin black line and shaded areas are the median and 95%-highest posterior density interval (HPDI) of the slope characterising the VPD increase over time. SRAD did not present any clear trend (slope not represented; i.e. the 95%-HPDI encompassed zero).
1. How do mean climate and climate anomalies determine interannual variation in tree growth rates, and what are the main climatic drivers?
2. Are species that grow faster in drier locations more resistant to water stress anomalies?
3. Can intrinsic growth rate and functional traits explain interspecific differences in growth sensitivities to climate?
4. Are drier and warmer forests more sensitive to positive anomalies in temperature and water stress?

2 | MATERIALS AND METHODS

2.1 | Study sites and demographic data

Individual tree annual absolute growth rates were calculated for 12,853 trees in 23 permanent forest plots of tropical rainforest located in northern Queensland, Australia, between 12°44′ S to 21°15′ S and 143°15′ E to 148°33′ E, and encompassing an elevation gradient between 15 and 1200 m a.s.l. and a period of 49 years (Figure 1a; Table S1) (20 CSIRO long-term plots (Bradford et al., 2014), and three more recent plots; see Supplementary Methods S1). Regular cyclonic disturbance contributes to the dynamics of the forests (Murphy et al., 2013). They cover a wide range of mean annual temperatures (19 to 26.1°C), precipitations (1213 to 3563 mm), solar radiation (17.8–19.4 MJ m⁻² day⁻¹) and vapour pressure deficit (VPD, 6.5–11.8 hPa) (Table S1). At plot establishment, all trees with stems ≥10 cm diameter at breast height (DBH) were mapped, identified to species level and measured for diameter. The 20 long-term plots were re-measured every 2 years for 10 years, and then at three- to four-year intervals, with diameter, recruits and deaths recorded, summing up to 11–17 censuses per plot. The remaining three plots were established between 2001 and 2012 and resampled one to three times (Table S1).

All available censuses were used to calculate individual annualized absolute growth rate (AGR) based on DBH at date 1 and 2 (t₁ and t₂), as:

\[
AGR = \frac{DBH_{t₂} - DBH_{t₁}}{(nb\ days)_{t₂-t₁}} \times 365 \tag{1}
\]

Abnormal AGR values were removed following Condit et al. (2004; see Supplementary Methods S1). Pteridophytes and palms species were excluded from the analyses due to their lack of secondary growth.

2.2 | Climate data

The effect of climate on growth was studied through four climate variables encompassing a wide range of variability across the plots and relevant for tree growth (see details in Supplementary Methods S1): mean temperature (T_{mean}), solar radiation (SRAD), VPD, and maximum climatological water deficit (MCWD; a proxy of the annual accumulated water stress over the drier season, estimated from climate data as the cumulative deficit between precipitation and evapotranspiration; hence, better capturing the seasonality of precipitation and potential soil water deficit than precipitation (Aragão et al., 2007; Malhi et al., 2009, 2015) (Tables S1 and S3a).

Climate data collection is detailed in the Supplementary Methods S1 and summarized here. Monthly climatic variables were obtained for the period 1970 to 2018 for each plot from ANUClimate v.2.0 (Hutchinson et al., 2014), a spatial model constructed from a new anomaly-based approach to the interpolation of Australia’s national point climate data to produce climate variables on a 0.01° longitude–latitude grid. The monthly climate variables were interpolated from the standard Bureau of Meteorology data network using elevation-dependent thin plate smoothing splines as calculated by the ANUSPLIN package (Hutchinson & Xu, 2013). The elevation dependence of these analyses yielded acceptably small predictive errors. Maximum and minimum temperatures were estimated with mean absolute predictive errors of around 0.5°C and solar radiation, VPD and rainfall were estimated with relative mean absolute predictive errors of 4%, 7% and 18%, respectively. The monthly actual evapotranspiration (a_{ave}) was derived from TerraClimate (Abatzoglou et al., 2018). The a_{ave} was used in combination with rainfall to calculate the monthly climatological water deficit (CWD). The CWD was reset to zero at the wettest month of the year and had an upper bound at 1000 mm. It was used to calculate monthly MCWD through a rolling maximum over the previous 12 months.

In each forest plot, a monthly 30-year historical mean and standard deviation were calculated over the 1981–2010 period for T_{mean}, SRAD, VPD, and MCWD (Table S1). On this basis, we calculated in each plot the monthly anomalies for each variable (i.e., monthly 30-year mean \( \mu \) subtracted from monthly value) and divided them by their location-specific 30-year monthly standard deviation \( \sigma \), yielding standardized anomalies (Aragão et al., 2007; Rifai et al., 2018):

\[
\text{anomaly}_{\text{std}}_{k,t} = \frac{(X_{k,t} - \mu_k)}{\sigma_k} \tag{2}
\]

where \( X_{k,t} \) is the climate variable value in plot \( k \) at time \( t \) (i.e., year and month), and \( \mu_k \) and \( \sigma_k \) are the monthly 30-year mean and standard deviation of the corresponding plot.

Standardized anomalies are expressed in units of standard deviation from monthly means over 1981–2010. This allows the comparison of plots differing not only in their historical means but also in the long-term variation range around them, that is, an important element to detect anomaly effects on tree growth across different climates (Rifai et al., 2018).

To build the climate covariates for the tree growth models, the monthly 30-year mean and standardized anomaly variables were averaged over the months between consecutive censuses (2–5 years). For MCWD, the maximum over the growth periods between two censuses was used instead of the weighted mean. The eight resulting interannual averaged variables were used as predictors to model
tree growth (see Section 2.5). Correlations among these variables, stand structure, and elevation are presented in Table S3a and the Supplementary Methods S1.

2.3 | Stand structure

As stand structure can vary between plots, we include its effect on tree growth through total plot basal area. Plot basal area (m² ha⁻¹) was calculated at each census, with expectations that increasing basal area would have a general negative effect on tree growth (Mulesdi et al., 2020; Sánchez-Salgueiro et al., 2015).

2.4 | Functional traits

Between July and September 2015, we measured 15 traits of 75 dominants, canopy tree species in eight plots along the gradient (Table 1; Tables S1 and S2) for plot and species details. Species were chosen to sample those that made up 80% of the standing biomass in these eight plots, but were also present in varying proportions in other plots; hence, covering the climatic range of the plot network. Trait data collection and measurement are detailed in Supplementary Methods S1. We measured leaf, wood, and maximum size traits for pairwise trait correlations, and Figure S1 for trait distribution.

Methods S1. We measured leaf, wood, and maximum size traits in these eight plots, but were also present in varying proportions in other plots; hence, covering the climatic range of the plot network. Trait data collection and measurement are detailed in Supplementary Methods S1. We measured leaf, wood, and maximum size traits for pairwise trait correlations, and Figure S1 for trait distribution along the elevation gradient. Traits were measured on three individuals per species and included photosynthesis and stomatal conductance at a reference CO₂ concentration of 400 µmol mol⁻¹ and irradiance of 1500 µmol photons m⁻² s⁻¹ (A_sat and gsat), dark respiration (R₀) at the same CO₂ concentration, the CO₂-saturated photosynthesis and stomatal conductance (A_max and gsmax), measured at 1200 µmol mol⁻¹ CO₂. The one-point method (De Kauwe et al., 2016) was used to estimate the maximum carbon transportation (V_max) for each individual from net photosynthesis measured at 400 µmol mol⁻¹ CO₂, and maximum light-driven electron flux (J_max) from net photosynthesis measured at 1200 µmol mol⁻¹ CO₂ (Bloomfield et al., 2018) (see Supplementary Methods S1 for correlation of one-point estimates and Aci curve-derived J_max and V_max of a subset of species). We also measured leaf stable carbon isotope ratio (δ¹³C), nutrient concentration, and leaf area, leaf mass per area (LMA), leaf thickness, and wood density (from branches, after bark removal). All traits were averaged at the species level for tree growth analyses.

2.5 | Data analysis

We addressed our four questions through three sets of Bayesian multilevel models (M1 to M3; details in Supplementary Methods S1).

2.5.1 | M1: Tree growth response to climate means and anomalies, and species differences in their sensitivities to climate

In the M1 models, we used 12,853 individuals from all 509 species to test the effects of climate on tree growth and to investigate trade-offs among species between intrinsic growth rate and growth sensitivity to climate covariates (ie, to address question 1). We built a two-level hierarchical Bayesian model of AGR, where the hierarchy included an upper level of response (hereafter grand coefficients or effects, affecting AGR across species) above a lower, species-level response. The higher level modelled AGR responses to covariates via hyperparameters (i.e., statistical distributions from which species-level intercepts and slope coefficients arose), while the lower level captured species-specific growth sensitivities to model covariates, and species-level intercepts (hereafter intrinsic AGR) captured unexplained growth variation across individuals, growth periods, and plots.

More specifically, we modelled individual log(AGR) as a species-specific function of (i) initial tree size (approximated by log(DBH) at the beginning of a growth period), (ii) the local 30-year mean of a climate variable, (iii) the anomalies of the same climate variable averaged over the studied growth period, and (iv) stand structure (approximated by plot basal area at the beginning of a growth period), using varying slopes (also known as random slopes) and a covariance matrix to estimate correlations among species-specific AGR sensitivities to the covariates, as:

\[
\log(\text{AGR}_{ijk,t}) \sim \text{Normal}(\mu_{ijk,t}, \sigma_R) \quad \text{[Likelihood]}, \quad (3.1)
\]

\[
\mu_{ijk,t} = a_0 + \beta_{i} \times \log(\text{DBH}_{i,t}) + \beta_{2} \times \text{meanClim}_{t} + \beta_{3} \times \text{climAnom}_{t} + \beta_{4} \times \text{BA}_{i,t} + \beta_{5} + \delta_{i} + \lambda_{i} \quad \text{[Linear model]}, \quad (3.2)
\]

\[
[\text{Adaptive priors for species - level param. }], \quad (3.3)
\]

\[
[S = \begin{pmatrix} 
\sigma_{\mu} & 0 & 0 & 0 \\
0 & \sigma_{\beta_{2}} & 0 & 0 \\
\vdots & \vdots & \ddots & \vdots \\
0 & 0 & \sigma_{\beta_{4}} & 0 \\
\end{pmatrix} R = \begin{pmatrix} 
\sigma_{\mu} & 0 & 0 & 0 \\
0 & \sigma_{\beta_{2}} & 0 & 0 \\
\vdots & \vdots & \ddots & \vdots \\
0 & 0 & \sigma_{\beta_{4}} & 0 \\
\end{pmatrix} \quad \text{[Construction of covariance matrix]}, \quad (3.4)
\]
### TABLE 1 Functional traits measured and their functions

<table>
<thead>
<tr>
<th>Organ</th>
<th>Trait type</th>
<th>Trait</th>
<th>Abbreviation</th>
<th>Units</th>
<th>Mean (min, max)</th>
<th>CV (%)</th>
<th>Functional role</th>
</tr>
</thead>
<tbody>
<tr>
<td>Leaf</td>
<td>Physiology</td>
<td>Net photosynthetic rate at saturating irradiance and ambient CO₂</td>
<td>$A_{sat}$</td>
<td>$\mu$mol CO₂ m⁻² s⁻¹</td>
<td>5.44 (0.98, 9.36)</td>
<td>28.6</td>
<td>Photosynthesis and growth</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Net photosynthetic rate at saturating irradiance and saturated CO₂</td>
<td>$A_{max}$</td>
<td>$\mu$mol CO₂ m⁻² s⁻¹</td>
<td>12.9 (7.7, 19.2)</td>
<td>19.1</td>
<td>Photosynthesis and growth</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Stomatal conductance at saturating irradiance and ambient CO₂</td>
<td>$g_{sat}$</td>
<td>mol H₂O m⁻² s⁻¹</td>
<td>0.071 (0.02, 0.146)</td>
<td>34.8</td>
<td>Control of carbon and water exchange between the leaf and the atmosphere; hence, influencing photosynthesis and water use efficiency (Liu et al., 2017)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Stomatal conductance at saturating irradiance and saturated CO₂</td>
<td>$g_{max}$</td>
<td>mol H₂O m⁻² s⁻¹</td>
<td>0.064 (0.018, 0.135)</td>
<td>33.1</td>
<td>Control of carbon and water exchange between the leaf and the atmosphere; hence, influencing photosynthesis and water use efficiency (Liu et al., 2017)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Maximum rate of electron transport</td>
<td>$J_{max}$</td>
<td>$\mu$mol e⁻ m⁻² s⁻¹</td>
<td>68.2 (39.3, 94.5)</td>
<td>18.0</td>
<td>Directly related to photosynthetic rate (Walker et al., 2014)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Maximum rate of Rubisco carboxylation</td>
<td>$V_{cmax}$</td>
<td>$\mu$mol CO₂ m⁻² s⁻¹</td>
<td>31.2 (11.2, 52.1)</td>
<td>25.3</td>
<td>Directly related to photosynthetic rates (Walker et al., 2014)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Ratio of maximum electron transport to maximum carboxylation rates</td>
<td>$J_{max}/V_{cmax}$</td>
<td>$\mu$mol e⁻ µmol⁻¹ CO₂</td>
<td>2.32 (1.67, 6.73)</td>
<td>26.9</td>
<td>Relative allocation to $J_{max}$ and $V_{cmax}$ (Smith et al., 2019)</td>
</tr>
<tr>
<td>Leaf</td>
<td>Metabolism</td>
<td>Maximum rate of dark respiration</td>
<td>$R_d$</td>
<td>$\mu$mol CO₂ m⁻² s⁻¹</td>
<td>0.826 (0.259, 1.74)</td>
<td>41.7</td>
<td>Metabolic rate; correlates with photosynthetic capacity (Atkin et al., 2015)</td>
</tr>
<tr>
<td>Leaf</td>
<td>Chemistry</td>
<td>Leaf carbon stable isotope ratio</td>
<td>leaf δ¹³C</td>
<td>‰</td>
<td>-30.4 (-32.9, -27.7)</td>
<td>4.6</td>
<td>Positively correlated with intrinsic water use efficiency and the ratio of intercellular to ambient CO₂ concentrations; hence, relying on stomatal conductance and photosynthetic capacity (Cernusak et al., 2013)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Leaf nitrogen per unit area</td>
<td>$N_{leaf}$</td>
<td>µg cm⁻²</td>
<td>177 (104, 268)</td>
<td>22.6</td>
<td>$N_{leaf}$ mainly supports the photosynthetic machinery, mostly the Rubisco carboxylation rate and hence photosynthesis (Quebbeman &amp; Ramirez, 2016; Walker et al., 2014; Wright et al., 2004)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Leaf phosphorus per unit area</td>
<td>$P_{leaf}$</td>
<td>µg cm⁻²</td>
<td>10.3 (4, 38.7)</td>
<td>46.5</td>
<td>Important determinant of photosynthetic rate (Walker et al., 2014)</td>
</tr>
<tr>
<td>Organ</td>
<td>Trait type</td>
<td>Trait</td>
<td>Abbreviation</td>
<td>Units</td>
<td>Mean (min, max)</td>
<td>CV (%)</td>
<td>Functional role</td>
</tr>
<tr>
<td>---------------</td>
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<td>--------</td>
<td>---------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Leaf</td>
<td>Structure</td>
<td>Leaf area</td>
<td>LA</td>
<td>cm²</td>
<td>540 (7, 32864)</td>
<td>695.9</td>
<td>Light capture efficiency and control of the boundary layer driving leaf heating-cooling dynamics (Wright et al., 2017)</td>
</tr>
<tr>
<td>Leaf thickness</td>
<td>Thickness</td>
<td>Thickness</td>
<td>mm</td>
<td></td>
<td>0.28 (0.17, 0.53)</td>
<td>25.6</td>
<td>Increases structural support and leaf lifespan; related to resource acquisition and use (Vile et al., 2005; Westoby et al., 2002)</td>
</tr>
<tr>
<td>Leaf mass per area</td>
<td>LMA</td>
<td>g m⁻²</td>
<td>126 (78, 216)</td>
<td></td>
<td>27.4</td>
<td></td>
<td>Relative allocation to biomass per leaf area; strongly correlated to leaf lifespan and thus nutrient use efficiency (Osnas et al., 2013)</td>
</tr>
<tr>
<td>Wood Structure</td>
<td>Wood density</td>
<td>WD</td>
<td>g cm⁻³</td>
<td></td>
<td>0.585 (0.312, 0.795)</td>
<td>17.1</td>
<td>Mechanical support, water transport and storage capacity (carbon and other nutrients, defence compounds) (Chave et al., 2009; Peters et al., 2021)</td>
</tr>
<tr>
<td>Whole plant</td>
<td>Maximum size</td>
<td>Maximum diameter (130 cm)</td>
<td>DBH_max</td>
<td>cm</td>
<td>54.6 (16.4, 113.2)</td>
<td>41.6</td>
<td>Proxy of maximum height, itself summarizing light-acquisition and growth strategies (Rüger et al., 2012; Westoby, 1998)</td>
</tr>
</tbody>
</table>

Note: These traits were measured on three adult individuals of 75 tree species and used to model intrinsic growth rate and growth response to mean climate, climatic anomalies and stand structure.

CV corresponds to coefficient of variation of a variable, as defined by the ratio of the standard deviation to the absolute value of the mean, expressed as a percentage.
$$R = \begin{pmatrix}
1 & \rho_{a_1,a_2} & \rho_{a_1,a_3} & \rho_{a_1,a_4} \\
\rho_{a_2,a_1} & 1 & \rho_{a_2,a_3} & \rho_{a_2,a_4} \\
\vdots & \vdots & \vdots & \vdots \\
\rho_{a_4,a_1} & \rho_{a_4,a_2} & \rho_{a_4,a_3} & 1
\end{pmatrix}$$

[Correlation matrix of species-level params.], (3.5)

$$\gamma_k \sim \text{Normal}(0, \sigma_j) \quad \text{[Adaptive priors for the k plots]}, \quad (3.6)$$

$$\delta_t \sim \text{Normal}(0, \sigma_j) \quad \text{[Adaptive priors for the t time periods]}, \quad (3.7)$$

$$\lambda_i \sim \text{Normal}(0, \sigma_j) \quad \text{[Adaptive priors for the i individuals]}, \quad (3.8)$$

$$a_{0, \beta_{1-4,0}} \sim \text{Normal}(0, 1) \quad \text{[Priors for grand intercept and slopes]}, \quad (3.9)$$

$$\sigma_{\alpha}, \sigma_{\beta_{1-4}}, \sigma_{\gamma}, \sigma_{\delta_1}, \sigma_{\lambda} \sim \text{HalfNormal}(0, 1)$$

[Adaptive priors for standard deviation params.], (3.10)

$$R \sim \text{LKJcorr}(2) \quad \text{[Prior for correlation matrix]}, \quad (3.11)$$

where $\alpha_j$ characterizes the intrinsic AGR of species $j$ and $\beta_{0, \beta_{1-4}}$ and $\beta_{0, \beta_{1-4}}$ characterize the AGR response of species $j$ to tree size, mean climate (1981–2010), standardized climatic anomalies, and plot basal area in plot $k$ for time interval $t$. The parameter $a_{0, \beta_{1-4,0}}$ represents the grand intercept, and the parameters $\beta_{1-4,0}$ are the grand slopes of model covariates whose posterior distributions represent the effect of covariates on AGR across all species.

The matrix of fitted correlation coefficients among all pairs of species-level intercepts and slopes $(\alpha_j, \beta_0, \beta_2, \beta_3, \beta_{1-4})$ allows evaluating correlations among species intrinsic growth rate (intercepts $\alpha_j$) and species AGR sensitivity to model covariates ($\beta_{1-4}$). For instance, if we consider VPD or MCWD, the $\rho_{\beta_{0, \beta_{1-4}}}$ parameter will be used to address question 2, that is, whether species AGR sensitivity to long-term conditions (eg, of VPD) is correlated to AGR sensitivity to climate anomalies (of VPD). Similarly, a model with a negative $\rho_{\beta_{0, \beta_{1-4}}}$ parameter and a negative $\beta_{1-4}$ slope would indicate that species with higher intrinsic growth rate ($\alpha_j$) tend to have higher sensitivity (ie, more negative slopes) to climate anomalies (ie, the intrinsic growth-related part of question 3). Using covariance matrix to pull information across species-level intercepts and slopes through the multinormal distribution also improves the accuracy of posterior likelihood estimates both across and within species (hierarchical levels 1 and 2, respectively) while limiting risks of overfitting through adaptive regularizing priors, or partial pooling (McElreath, 2020).

Parameters $\gamma_k, \delta_t, \lambda_i$ varying intercepts capturing the residual variation in expected individual AGR occurring among forest plots, time periods between consecutive censuses (characterized by the years beginning and ending a given census period) and individual stems, respectively. This model was run separately for each of the four climate variables ($T_{\text{mean}}, \text{SRAD, VPD and MCWD}$) to manage model complexity (representing a total of four M1 models).

### 2.5.2 M2: Trait-mediated species-level tree growth response to climate

Models M2 have the same hierarchical structure as M1, but additionally include the role of species traits in AGR response to climate and only consider 75 locally dominant species with trait data instead of all 509 species. Models M2 were used to address question 3, that is, to test whether traits mediate climate effects on tree growth. We thus used a subset of 5191 individuals from the 75 species with trait data in M2, the species-level intercept and slopes are modelled as depending on species mean trait value such that both species-specific intrinsic AGR and AGR sensitivity to covariates can be influenced (either accentuated or lessened) by species traits (Fortunel et al., 2018; Rüger et al., 2012; Uriarte et al., 2016) as:

$$\log(\text{AGR}_{ijk}) \sim \text{Normal}(\mu_{ijk}, \sigma_R) \quad \text{[Likelihood]}, \quad (4.1)$$

$$\mu_{ijk} = a_i + \beta_0 \times \text{DBH}_{ijk} + \beta_2 \times \text{meanClim}_{ijk} + \beta_3 \times \text{climAnom}_{ijk} + \beta_{1-4} \times \text{BA}_{ijk} + \gamma_k + \delta_t + \lambda_i$$

[Linear model - level 1], (4.2)

$$a_i = a_0 + \alpha_1 \times \log \text{(Trait)}$$

[Linear model - level 2], (4.3)

$$\beta_{2-4i} = \beta_{2-40} + \beta_{2-41} \times \log \text{(Trait)}$$

[Linear model - level 2], (4.4)

$$\begin{pmatrix}
\alpha_i \\
\beta_0 \\
\vdots \\
\beta_4
\end{pmatrix} \sim \text{MVNormal}$$

[Adaptive priors for species-level param.], (4.5)

$$S = \begin{pmatrix}
\sigma_{a} & 0 & 0 & 0 \\
0 & \sigma_{\beta_0} & 0 & 0 \\
\vdots & \vdots & \ddots & \vdots \\
0 & 0 & 0 & \sigma_{\beta_4}
\end{pmatrix}$$

[Construction of covariance matrix], (4.6)
where Equations (4.1), (4.2), (4.5)–(4.7) are the same as Equations (3.1)–(3.5) of M1, whilst species-level intercepts and slopes are mediated by species mean trait value (Equations 4.3–4.4; see priors in Supplementary Methods S1). Parameter $\alpha_i$ is the species-level departure from the grand intercept ($\alpha_{1,0}$) for an increase of one standard deviation in the log(Trait) value of species $j$ (direct effect of trait on AGR), whilst $\beta_{2,4,i}$ are the departures from the grand slope of the corresponding model covariates for an increase of one standard deviation in the log(Trait) value of species $j$ (trait mediation of AGR response to climate and stand structure). If the sign of a trait coefficient ($\beta_{2,4,i}$) is the same as that of the corresponding grand covariate effect ($\beta_{2,4,1}$), then increasing values of the Trait accentuate the effect of the covariate (meanClim, climAnom or BA) on tree growth (ie, push $\beta_{2,4,i}$ further away from 0). If the signs are different, increasing values of Trait, attenuate the effect of covariate c (ie, pull $\beta_{2,4,i}$ closer to 0). We did not include the role of species traits in AGR response to tree size because some traits can change through tree ontogeny (Fortunel et al., 2020) and our trait data does not encompass species tree size ranges. M2 models were run separately for each of the four climate variables and for each of the 15 functional traits to manage model complexity (representing a total of 60 M2 models).

In both M1 and M2 models, we standardized the response variable log(AGR) and all covariates—but climate anomalies—to mean zero and unit standard deviation, to allow relative importance comparisons between covariates through slope coefficients (Schielzeth, 2010), and to ease plausible weakly-informative prior assignment to the parameters (McElreath, 2020) (see Supplementary Methods S1). We did not standardize averaged monthly anomalies to maintain their interpretability as deviations from long-term means in terms of plot-specific units of standard deviation (see Equation 2; ie, mean anomaly covariate slope coefficients are not directly comparable to other covariate mean slopes). Individual trait measurements were averaged per species and log-transformed prior to standardization to mean zero and unit standard deviation, thus implying that parameter $\beta_{2,4,i}$ corresponds to $\beta_{2,4,0}$ at the mean trait value of the dataset.

### 2.5.3 | M3: Plot-level tree growth response to climate anomalies and interaction with mean climate

M3 models evaluate plot-level growth response to climate anomalies, and whether it varies depending on local mean climates (ie, question 4; e.g. whether plot-level AGR sensitivity to VPD anomalies is higher in drier sites). M3 models were run on the basis of all 509 species and did not include trait data, like M1 models. We focused on the tree growth at the plot level and modelled the expected log(AGR) as a linear function of mean climate and climate anomalies. We used a similar Bayesian hierarchical model as described for M2, where plotspecific average AGR depended on climate anomalies, whose effect on AGR itself depended on the plot mean climate, as:

$$\log(AGR_{ijk}) \sim \text{Normal}(\mu_{ijk}, \sigma_k)$$  \hspace{1cm} \text{(Likelihood),} \hspace{1cm} (5.1)

$$\mu_{ijk} = a_k + \beta_{1k} \times \text{climAnom}_{ik} + \tau_i + \delta_j$$  \hspace{1cm} \text{(Linear model - level 1),} \hspace{1cm} (5.2)

$$a_k = a_0 + \alpha_i \times \text{meanClim}_k$$  \hspace{1cm} \text{(Linear model - level 2),} \hspace{1cm} (5.3)

$$\beta_{1k} = \beta_{1,0} + \beta_{1,1} \times \text{meanClim}_k$$  \hspace{1cm} \text{(Linear model - level 2),} \hspace{1cm} (5.4)

where $a_k$ is the average growth rate in plot $k$, and $\beta_{1,1}$ characterizes the growth response of plot $k$ to standardized climate anomalies for time interval $t$. $\alpha_i$ is the mean intercept value (ie, mean absolute growth rate) across plots, and $a_k$ is the departure from the grand mean for one unit increase in mean climate (see d priors in Supplementary Methods S1). $\beta_{1,0}$ is the grand slope of climate anomalies, and $\beta_{1,1}$ is the departure from this grand mean for a one-unit increase in mean climate (medication of the effect of anomalies on growth by the plot mean climate). Parameters $\tau_i$, $\delta_j$, $\lambda_k$ are varying intercepts for species, census periods, and individual stems, respectively.

We run M3 models only for two climate variables (VPD and SRAD), as we found they were the most important climate variables for tree growth in M1 and M2 models (see Section 3). Standardization of variables was carried out as for M1.

### 2.5.4 | Trends in climate over time

To explore the implications of the effects of climate anomalies on tree growth, we built a separate set of hierarchical Bayesian models to test for linear temporal trends in mean annual climate variables between 1971 and 2019. We used varying year slopes per plots to...
allow plot-specific trends (model details in Supplementary Methods S1). We also run the models for the period 2000 to 2019 for comparison with recent analyses suggesting an increasing rate of VPD increase over time since the late 90s (Yuan et al., 2019). Annual mean temperature and VPD increased by 0.015°C and 0.02 hPa per year between 1971 and 2019 (R² = .97 and .84, respectively, Table S4; illustration in Figure 1b) and by 0.038°C and 0.045 hPa per year between 2000 and 2019 (R² = .98 and .81, respectively, Table S4). There was no general temporal trend for MCWD or SRAD (Figure 1c).

2.5.5 Analysis of model outcomes

All model parameter posteriors were summarized through their median and 95%-highest posterior density interval (HPDI) (i.e. the narrowest posterior interval encompassing 95% of the probability mass, corresponding to the coefficient values most consistent with the data; McElreath, 2020). Model covariates were considered important at two high levels of confidence when their coefficient had a posterior probability of over 95% or 90% of being either positive or negative (HPDI not encompassing zero).

The goodness-of-fit of the models was assessed through the squared Pearson correlation between the observed AGR and the AGR predicted by the fitted model (R²). M1 and M2 models had high explanatory power, with R² of .46 and .52 on average, respectively. M3 models, with VPD and SRAD as climate variables, had an R² of .67 and .63, respectively.

Bayesian updating of parameters was performed via the No-U-Turn Sampler (NUTS) in Stan (Carpenter et al., 2017), using three chains and 3000 steps (1500 warm-up). All models mixed well and converged (Rhat within <0.01 of 1). Models were run in the R environment (R Core Team, 2020) using the packages ‘brms’ (Bürkner, 2017), ‘tidybayes’ (Kay, 2020) and ‘tidyverse’ (Wickham et al., 2019).

3 RESULTS

3.1 Contribution of climate means and anomalies to tree growth

The M1 models of tree growth, based on all 509 tree species and not including trait data, indicated that the main climate drivers affecting tree growth across species were the climate means and anomalies in Tmean, SRAD and VPD (Figure 2; Figure S3, Table S5). Tree growth was higher in forests with higher mean Tmean, SRAD and VPD (βj: 0.17 [0.08, 0.26], 0.05 [0.02, 0.08], and 0.09 [0.02, 0.17], respectively; median and 95%-HPDI; unless otherwise stated, all intervals are 95%-HPDI). However, tree growth was reduced when forests experienced positive anomalies in temperature, SRAD and VPD (βj: −0.12 [−0.17, −0.07], −0.34 [−0.42, −0.26] and −0.13 [−0.19, −0.06], respectively). Contrary to our expectation, anomalies in MCWD had no clear effect on tree growth across species (Figure 2; Figure S2, Table S5). Tree growth sensitivity to climate, stand structure and tree size varied widely among species (illustration in Figure S3). Similar results were obtained from the M2 models (a subset of 75 species with trait data) (Figure S5a–d, Table S5), though we no longer detected the effects of temperature anomalies and VPD and solar radiation means in this reduced dataset.

3.2 Coordinated tree growth sensitivities to climate means and anomalies

Using the fitted matrix of correlations among species-level intercepts and slopes from the M1 models on all 509 species (matrix R, see Equation 3.5) allowed testing for different sensitivities to climate anomalies between fast- and slow-growing species, and between species growing better at opposite extremes of the range

![FIGURE 2 Grand effects of climate, stand structure and tree size on tree growth (based on all 509 species; four M1 models). Red and blue arrows indicate clear negative and positive effects (i.e., slope coefficient 95%-highest posterior density interval [HPDI] not encompassing zero). Arrow widths are proportional to the median of the covariate slope posteriors (grand slopes, values in rectangles; see βj in Equations 3.1-3.11) (details in Figure S2 and Table S5)
of mean climates. Fast-growing species (i.e., with high intrinsic AGR) were more sensitive than slow-growing species to the negative effects of both VPD anomalies and plot basal area on tree growth (Figure 3c; Figure S4; \( \rho = -0.36 [-0.48, -0.23] \) and \( \rho = -0.29 [-0.41, -0.17] \), respectively). Species that grew better in cloudier forests (i.e., lower SRAD) tended to show steeper growth decreases when experiencing positive anomalies in solar radiation (Figure 3b; \( \rho = 0.17 [0.01, 0.33] \)). Species that grew faster in drier forests (i.e., higher VPD) were more negatively affected by positive VPD anomalies (Figure 3a; \( \rho = -0.15 [-0.29, 0.00] \)). Finally, species most negatively affected by positive anomalies in VPD also experienced stronger growth decrease in denser forests (high basal area) (Figure 3d; \( \rho = 0.27 [0.14, 0.40] \)).

### 3.3 | Drier rainforests are more sensitive to VPD anomalies

M3 models highlighted clear interactions between the effects of climate anomalies and mean climate for VPD (\( \beta_{1,1} = -0.26 [-0.39, -0.13] \); see Equations (3.1)–(3.11)), and to a lesser extent for solar radiation (\( \beta_{1,1} = -0.09 [-0.18, -0.01] \), 90%-HPDI; Table S5). Drier tropical rainforests showed a steeper decrease in plot-level growth in response to increasing positive VPD anomalies (Figure 4a; Table S5). Cloudier forests exhibited a stronger decrease in plot-level growth with increasing positive SRAD anomalies (Figure 4b; Table S5).

### 3.4 | Functional traits influence species intrinsic tree growth and their response to climate drivers

Based on the M2 models of tree growth for the subset of 75 locally dominant species with trait data, species intrinsic growth increased with dark respiration rate \( R_d \), DBHmax, leaf P content, Asat, \( V_{cmax} \), leaf \( \delta^{13}C \) and LMA (Figure 5; Figures S5e; details in Table S5). Species traits also mediated the effects of climate and forest structure on tree growth, either by accentuating them (species with high values of the trait respond more strongly) or by attenuating them (species with low values of the trait are more sensitive) (Figure 5; details in Figure S6; Table S5). Leaf \( \delta^{13}C \) and P content exacerbated the negative effects of positive anomalies in SRAD on tree growth, whilst \( A_{max} \), \( g_{max} \), \( g_{sat} \) and \( J_{max} \) attenuated them (Figure 5; Figure S6f, Table S5). The negative effects of anomalies in VPD on tree growth were exacerbated in species with high leaf \( \delta^{13}C \), DBHmax, leaf P, and LMA, further confirming that VPD anomalies had the most negative effects on fast-growing species (Figure 3c), but also those with low \( g_{max} \) or leaf area (Figure 5; Figure S6g). Tree growth was less reduced by denser forest environments (high plot basal areas) in species with high wood density, low \( R_d \) and low leaf \( \delta^{13}C \) (Figure 5; Figure S6i–l).

**FIGURE 3** Correlations among species-level growth sensitivities highlighting joint responses to multiple drivers (M1 models; 509 species). Joint growth sensitivities to: (a) vapour pressure deficit (VPD) anomalies and mean VPD; (b) Solar radiation anomalies and mean solar radiation; (c) VPD anomalies and intrinsic growth rate; (d) VPD anomalies and plot basal area. Circles are species, placed at the median of their corresponding coefficient posteriors. Vertical and horizontal bars are 95%-highest posterior density interval (HPDI) for the corresponding coefficients. Species for which both plotted coefficients were important are plain blue; other species are shaded. Blue and red regression lines indicate positive and negative correlations (\( \rho \), see Equation 3.5 in Supplementary Methods S1), respectively. Values beyond and below zero indicate positive and negative effects on growth rates, respectively. Mean, lower and upper 95%-HPDI are in the upper right-hand corner of the figures.
FIGURE 4  Plot-level growth sensitivity to positive (a) vapour pressure deficit (VPD) anomalies and (b) solar radiation (SRAD) anomalies (b) across the full range of the corresponding mean climate variable (M3 models). Circles and vertical bars are the median and 95%-highest posterior density interval (HPDI) of the plot-level slope posteriors characterizing the growth rate responses to climate anomalies. The plot-level models including VPD (a) and SRAD (b) had a marked interaction between anomalies and long-term mean (Table S5), so that plot-level sensitivities to a given anomaly depend on plots’ long-term mean. (a, b) Illustrate those interactions through the differences of plot-level growth sensitivity to positive anomalies across the range of long-term means of the corresponding variable. The represented plot-level coefficients were calculated for a positive standardised anomaly equal to the 95th percentile of anomalies in the data, ie, a standardized anomaly of 0.8 (a) and 0.4 (b). The red and blue regression lines and shaded areas are decreasing and increasing slopes, respectively (median and 95%-HPDI, not encompassing zero), of the represented plot-level coefficients along the long-term means. Horizontal dashed line: limit between positive and negative slope coefficients indicating a growth rate increase and decrease, respectively, with the positive anomaly.

FIGURE 5  Mediation of intrinsic growth rate and climate anomaly effects by species functional traits (75 species). The figure only presents important trait effects (95%- or 90%-highest posterior density interval [HPDI] not encompassing zero; non-transparent and semi-transparent arrows, respectively) (see Figures S5 and S6 for all model coefficients and all traits). Red and blue plain arrows indicate negative and positive direct effects of traits on species’ intrinsic growth rate ($\alpha_i$, see Equation 4.3). Dashed arrows are slope coefficients indicating indirect trait effects on growth through the effects of environmental covariates, ie, accentuation (red) or attenuation (blue) of the negative effects of VPD or SRAD anomalies when trait values increase ($\beta_{3,1}$, see Equation 4.4). Arrow widths are proportional to the median of the covariate slope posterior across species (ie, grand slope; details in Figure S6 and Table S5).
4 | DISCUSSION

In this study, we aimed to disentangle the influences of mean climate and climate anomalies on interannual tree growth and defined how species functional traits mediated climate effects by combining 49 years of demographic data, functional traits and climatic data along a climatic gradient in 23 tropical rainforests of Australia.

4.1 | What are the important climatic drivers for tree growth?

Solar radiation (SRAD) and atmospheric water demand (VPD) anomalies were the two overarching climatic drivers of tree growth across pre-existing climatic conditions and species in our study. These two variables were also the main drivers of seasonal stand-level net primary productivity in aseasonal forests across the tropics (Rifai et al., 2018), and increasing VPD due to anthropogenic climate change has repeatedly been shown to impact tree growth, biomass and vegetation health (Eamus et al., 2013; Novick et al., 2016; Rifai et al., 2019; Sanginés de Cárcer et al., 2018; Yuan et al., 2019). The pervasive negative effect of VPD anomalies on tree growth in our study is consistent with expectations from stomatal conductance models (Grossiord et al., 2020), with stomatal closure and ensuing restriction of CO₂ assimilation rate triggered by VPD values exceeding the climate mean and usual variation range. This negative effect of VPD is expected to be amplified by SRAD anomalies, as VPD depends on leaf temperature, which itself increases with SRAD (Grossiord et al., 2020). The negative influence of VPD anomalies on tree growth may be additive to that of VPD anomalies, as previously shown (Krause & Winter, 2020; Rifai et al., 2018, 2019). Furthermore, positive SRAD anomalies did not enhance tree growth but reduced it, as would be expected from a VPD-related effect. However, the effect of SRAD anomalies on tree growth was probably more than a mere reflection of VPD, as anomalies in SRAD and VPD were only moderately correlated (r = .33, Table S3a). Excess or fluctuating light, and changes in light quantity and quality are other potential mechanisms underlying SRAD anomaly effects, as these can be direct physiological stressors (Krause & Winter, 2020; Roeber et al., 2020), or indirectly influence the response to other abiotic or biotic stresses (Roeber et al., 2020).

The strong effect of VPD anomalies compared to the undetectable effect of MCWD anomalies suggests that VPD may limit tree growth before soil water becomes limiting, further confirming previous results in temperate and tropical forests (Choat et al., 2012; Konings et al., 2017; Novick et al., 2016; Rifai et al., 2018; Sanginés de Cárcer et al., 2018). This is a key result, given the generalized tree growth decrease potentially driven by increasing VPD anomalies, as VPD has been strongly increasing in the tropics due to anthropogenic climate change (Rifai et al., 2019). Yuan et al. (2019) highlighted a particularly-strong increasing VPD trend at the global scale beginning in the late 1990’s (0.017 hPa/year). Modelling VPD anomalies through time from 2000 to 2019 in our dataset, we detected a 3.8-fold stronger VPD increase rate across all plots (0.045 hPa/year, 90%-HPDI: 0.019, 0.066; R² = .80; details in Table S4; eg Figure 1b). This trend itself was stronger than the 1971–2019 trend in our dataset (0.020 hPa/year; R² = .84; Table S4), indicating a sharper-than-previously-thought VPD increase in the past two decades. This rapid increase of VPD anomalies through time combined with the generalized ensuing decrease in tree growth and growth sensitivity variability to VPD among species (Figure S3, Table S5) suggests that tropical forest composition and functions may be strongly altered by ongoing climate change, especially by VPD. It is worth noting that soil water deficit also depends on evapotranspiration estimates accuracy and variables unaccounted for, here (eg, soil-water retention capacity, topography), so that the importance of soil-related water stresses should be interpreted with caution.

In spite of the suppressing effects of increasing anomalies in SRAD, VPD and Tmean, average growth rates were higher in warmer and sunnier forests (ie, higher long-term means), across species (Figure 2) and within many species (Table S5). While long-term Tmean was highly correlated with elevation (r = ~.95; Table S3a), mean solar radiation was not correlated with neither elevation nor the other climate variables (Table S3a). This suggests that these forests are in general energy-limited along the elevation gradient (faster growth in lowland forests), and light-limited across the gradient, supporting previous results along an Amazon-Andes elevation gradient (Fyllas et al., 2017). Our gradient of mean climates encompassed 7 to 51% of the global-scale climate space of tropical forests, but did not encompass their driest and warmest conditions (see Figure 57). Future studies will need to cover a broader range of climate values to test how generalizable the relationships that we detected are for tropical forests worldwide.

4.2 | Trade-offs in tree growth responses to climate

We showed that two aspects allowed understanding the broad range of species differences in growth response to VPD anomalies: the long-term mean VPD where species grew better, and the contrast between slow- and fast-growing species (Figure 3a,c). The models including plot-specific responses to climate anomalies additionally showed that forest growth sensitivity to VPD anomalies was stronger in drier forests, mostly at the higher end of the VPD range (Figure 4a). This result could be driven by higher levels of obligate or facultative deciduousness, as even the wettest rainforests have seasonal peaks in leaf fall (Edwards et al., 2018) and the drier the forest the earlier the leaf fall peak and the shorter the growing season. Our results support recent findings indicating that drier forests could be more sensitive to increasing VPD anomalies (Aguirre-Gutiérrez et al., 2019, 2020; Esquivel-Muñoz et al., 2020; Powers et al., 2020), which would here translate into drier rainforests already being under water stress and therefore closer to a threshold of further growth.
decrease than moist rainforests. This effect may not be linear and will need to be further tested with more plots encompassing diverse water-stress conditions.

Similarly, Sullivan et al. (2020) recently argued that warmer forests may be closer to a temperature threshold beyond which woody productivity would decrease. In our study, this would translate into expectations that forests and species adapted to warmer conditions would respond more negatively to further temperature increases. Our results are consistent with this expectation but suggest that the temperature effect manifests itself indirectly through VPD.

Species that grew faster in cloudier forests showed the strongest growth reduction due to positive SRAD anomalies (Figure 3b). This may reflect species differences in light-use strategies, with species that grow well under low direct-sunlight conditions not benefitting from brighter conditions. This was supported by the stronger negative effects of SRAD anomalies in species with lower maximum photosynthetic capacity, stomatal conductance and electron transport capacity (Figure 5), a trait syndrome consistent with shade-tolerance strategies (He et al., 2019). This interpretation was supported in the plot-level analyses by the steeper growth rate decreases in the cloudier forests in response to positive SRAD anomalies (Figure 4b), which may stem from a plot-wide relatively more marked adaptation to shade tolerance.

### 4.3 Functional traits mediate the effects of climate anomalies on tree growth

Traits directly influenced species intrinsic growth rate in locally dominant species in our plot network. As expected, the intrinsic growth rate increased with metabolism ($R_{ch}$, maximum size (DBH$_{max}$) and acquisitive chemical and physiological traits related to the photosynthetic machinery (leaf P content, $A_{sat}$ and $V_{max}$). However, it also increased with leaf $\delta^{13}$C and LMA, contrary to expectations as high values of these traits correspond to tough, long-lived leaves and high intrinsic water use efficiency (Cernusak et al., 2013; Osnas et al., 2013). In our study, leaf $\delta^{13}$C was positively correlated with leaf N and P contents (Table S3b), suggesting variation in $\delta^{13}$C among species may have been driven more by photosynthetic capacity than by stomatal conductance. The positive association of LMA and growth, also reported in previous studies (Gray et al., 2019; Poorter et al., 2008; Wills et al., 2018), could be explained by a change in the cost-benefit balance of acquisitive traits with plant size (Gibert et al., 2016; Gray et al., 2019).

An overarching finding is that species traits can enhance our understanding of differences in species growth response to the anomalies of SRAD and VPD, and to forest stand structure. Our results confirmed that resource-acquisitive species overall had a higher intrinsic growth rates and that their growth was more sensitive to positive anomalies in SRAD and VPD. This highlights a trade-off between fast growth (via high allocation to acquisitive tissues) and sensitivity to atmospheric water stress, consistent with expectations from the ‘fast-slow’ plant economics spectrum (Reich, 2014).

Most physiological traits directly related to photosynthesis (Table 1) successfully captured species differences in growth sensitivity to SRAD anomalies (Figure 5; Figure S6), confirming the importance of physiological traits to investigate potential mechanisms underlying differences in demographic responses to climate change among species (Brodribb et al., 2020; Powers et al., 2020; Rowland et al., 2021). Increasing values of these traits attenuated the tree growth reduction following increasing SRAD anomalies (Figure 5; Figure S7), suggesting that species investing in a more responsive and flexible photosynthetic machinery may cope better with unusually high direct exposure to sunlight. Whilst most traits that increased species intrinsic growth rate also exacerbated the negative effects of VPD anomalies on tree growth, the mediation of SRAD anomalies by species traits was mostly independent of the fast-slow spectrum (Figure 5; Figures S5 and S6). For example, whilst leaf P concentration, stable carbon isotope ratio and the maximum photosynthetic capacity tended to increase intrinsic growth rate, the two former accentuated while the latter attenuated the negative effects of SRAD anomalies on tree growth (Figure 5).

Here, the data at hand did not allow us to integrate intraspecific or temporal trait variability into the growth models, nor could we consider traits from rare species that may occupy different parts of the trait space (Leitão et al., 2016; Mouillot et al., 2013; Violle et al., 2017). These sources of variation may potentially change some effect sizes (Bloomfield et al., 2018; Rowland et al., 2021; Yang et al., 2018, 2021), although this would be unlikely to affect the direction of the effects shown here, as the structure of our multilevel models was centred on species ranking, which is usually conserved (Auger & Shipley, 2013; Oliveras et al., 2020). Exploring sources of uncertainty in species traits such as a potential climate change-driven trait acclimation over time for certain traits (Bloomfield et al., 2018; Way & Yamori, 2014), tissue-level traits not providing a consistent and generalizable upscaling to whole-tree energy allocations and strategies across species (Yang et al., 2018), or climate effects on growth being mediated by multivariate phenotypes (i.e., trait interaction effects) (Laughlin et al., 2018; Pistón et al., 2019), will be a challenge to address in future studies, mostly in terms of data collection.

### 4.4 Stand structure as a driver of tree growth variation

Plot basal area consistently strongly reduced tree growth across species and explained more growth variation than mean climate for all four climate variables. Although plot basal area was partly correlated with elevation, the 30-year average of $T_{\text{mean}}$ and VPD ($r = -0.63, -0.59$, and $-0.47$, respectively; Table S3a), the slope coefficient of the basal area remained virtually unchanged across models including $T_{\text{mean}}$, VPD, or the other less correlated covariates (and was much steeper than the slopes of long-term $T_{\text{mean}}$, VPD), so that the stand structure effect detected here is unlikely to indirectly reflect $T_{\text{mean}}$ or VPD, and is likely to approximate—though in an imperfect spatially implicit way—the mean neighbourhood crowding of the plot. Furthermore, the faster
growth in less dense environments across forest plots suggests a release from competition for light. This is supported by the general light-limitation suggested by the faster growth in sunnier sites. Slower growth in denser environments may also suggest an increase in competition for resources or attacks by natural enemies. Neighbourhood crowding has indeed been shown to strongly reduce tree growth in tropical and temperate forests (Clark et al., 2014; Fortunel et al., 2016; Uriarte et al., 2016; Zambrano et al., 2017). In line with these studies, we found that conservative species with high wood density suffered less growth reduction from increasing plot basal area, whilst acquisitive species with high dark respiration rate and leaf δ¹³C were more sensitive to increasing plot basal areas (Figures S5 and S6).

In summary, we have shown how long-term demographic data across multiple plots encompassing environmental gradients, combined with functional traits collection can yield insights into how climate affects the interannual variation of tree growth at different temporal scales, and give important clues into which species and forests may be particularly vulnerable to climate change, and why. Our findings emphasise the importance of functional traits—and notably those related to photosynthesis and water use efficiency—to understand species differences in demographic sensitivity to abiotic and biotic drivers. Future efforts to further characterise how climate and neighbourhood crowding affect tree growth, survival and population growth across environmental gradients, and how these effects are mediated by species traits will help improve predictions of forest response and future ecosystem functions to climate change under different trajectories.

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CONFLICT OF INTEREST
The authors declare there are no competing interests.

AUTHOR CONTRIBUTIONS
David Bauman, Claire Fortunel and Yadvinder Malhi designed the study. David Bauman tidied and vetted the demographic and trait data and performed the analyses. David Bauman and Claire Fortunel designed the statistical models of tree growth. Sami W. Rifai helped generating the climatic covariates and created Figure 1a. Imma Oliveras and Jesús Aguirre-Gutiérrez contributed ideas and constructive feedback to early versions of the work. Jesús Aguirre-Gutiérrez helped obtain climate data and provided feedback on an early version of the work. Lucas A. Cernusak and Lisa P. Bentley led the trait data collection, assisted by Raymond Dempsey, Brandon E. McNellis, Hugo R. Ninantay-Rivera, Jimmy R. Chambi Paucar and Paul E. Santos-Andrade. Michael F. Hutchinson provided the final raw climate data. Matt Bradford supplied demographic data for the 20 CSIRO plots and Robson Creek. Lucas A. Cernusak, Sami W. Rifai, Jesús Aguirre-Gutiérrez, Guillaume Delhaye provided feedback to part of the discussion. Susan G. W. Laurance contributed demographic data of Daintree Observatory. David Bauman led the writing with regular feedback from Yadvinder Malhi, Claire Fortunel and Sean M. McMahon on intermediate stages of the analyses and manuscript. All authors commented on the manuscript and gave their approval for publication.

DATA AVAILABILITY STATEMENT
The raw demographic data that supported the findings are available in Bradford et al. (2014; see References) and CSIRO Data Access Portal [https://doi.org/10.4225/08/59475c67be7a4]. The R code for the analyses of the study is in Supplementary Methods S2. The raw climate data and processed data used in the models are openly available in the Github repository at [https://github.com/davbauman/Bauman_et_al_2021_GCB], and raw trait data will be available upon reasonable request.

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