Evaluating the potential of full-waveform lidar for mapping pan-tropical tree species richness


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Abstract Aim: Mapping tree species richness across the tropics is of great interest for effective...
**1 | INTRODUCTION**

Tropical forests are known for their high tree species diversity. Current estimates suggest in the order of 15,000 tree species in Amazonia alone, in contrast to 124 tree species in temperate forests in Europe, and more than 40,000 different tree species across the tropical region (Slik et al., 2015; Ter Steege et al., 2015). High levels of tree species richness may contribute to maximizing the provision of essential ecosystem services (Liang et al., 2016). Unfortunately, 35% of pre-agricultural global forest cover has been lost over the past 300 years, largely due to increasing human pressures on the environment. Eighty-two percent of the remaining forest is estimated to have experienced some degree of human impact (Watson et al., 2018). The Convention of Biological Diversity (CBD) and Group on Earth Observations Biodiversity Observation Network (GEO BON) have developed a list of important variables aiming to provide quantitative information on biodiversity to reach the Aichi biodiversity targets 2020 (Pereira et al., 2013; Skidmore et al., 2015). Among the identified needs is the mapping of taxonomic diversity at high spatial resolution over large scales (Pereira et al., 2010). Here, we focus on tree species richness. The collection of tree species diversity data is traditionally done in the field, and such information has previously been used to create predictive maps of tree species richness across the globe at low spatial resolution (Kier et al., 2005; Mutke & Barthlott, 2005). More recently, passive remote sensing data, such as optical imagery from various airborne and spaceborne platforms, have been used in combination with field reference data to predict tree species diversity in different regions (Bongalov et al., 2019; Carlson, Asner, Hughes, Ostertag, & Martin, 2007; Féret & Asner, 2014; Foody & Cutler, 2006; Rocchini et al., 2016; Schäfer, Heiskanen, Heikinheimo, & Pellikka, 2016). Even though such methods have conservation and biodiversity management. In this study, we evaluated the potential of full-waveform lidar data for mapping tree species richness across the tropics by relating measurements of vertical canopy structure, as a proxy for the occupation of vertical niche space, to tree species richness.

**Location:** Tropics.

**Time period:** Present.

**Major taxa studied:** Trees.

**Methods:** First, we evaluated the characteristics of vertical canopy structure across 15 study sites using (simulated) large-footprint full-waveform lidar data (22 m diameter) and related these findings to in-situ tree species information. Then, we developed structure–richness models at the local (within 25–50 ha plots), regional (biogeographical regions) and pan-tropical scale at three spatial resolutions (1.0, 0.25 and 0.0625 ha) using Poisson regression.

**Results:** The results showed a weak structure–richness relationship at the local scale. At the regional scale (within a biogeographical region) a stronger relationship between canopy structure and tree species richness across different tropical forest types was found, for example across Central Africa and in South America [$R^2$ ranging from .44–.56, root mean squared difference as a percentage of the mean (RMSD%) ranging between 23–61%]. Modelling the relationship pan-tropically, across four continents, 39% of the variation in tree species richness could be explained with canopy structure alone ($R^2 = .39$ and RMSD% = 43%, 0.25-ha resolution).

**Main conclusions:** Our results may serve as a basis for the future development of a set of structure–richness models to map high resolution tree species richness using vertical canopy structure information from the Global Ecosystem Dynamics Investigation (GEDI). The value of this effort would be enhanced by access to a larger set of field reference data for all tropical regions. Future research could also support the use of GEDI data in frameworks using environmental and spectral information for modelling tree species richness across the tropics.

**KEYWORDS**

biodiversity, canopy structure, GEDI, lidar, plant area index, tropical forests
been developing progressively over the last decade, they are not yet operational for mapping tree species richness across the tropics due to, among others, a lack of consistent remote sensing and training data over such scales, insufficient model accuracy and/or low spatial resolution.

The scientific community has called for bolder science in conservation strategies to enable effective management of the Earth’s forests and allow for better conservation of our natural ecosystems (Lewis, Edwards, & Galbraith, 2015; Watson et al., 2016). In this study, we focus on the use of active remote sensing, specifically lidar, for mapping taxonomic tree species richness in the tropics. While local tropical forest diversity is largely independent of biomass in intact forests (Sullivan et al., 2017), it remains unclear if substantial amounts of variation in species diversity are associated with other features of forest structure. Here, we explore for the first time whether small-scale vertical canopy structure variation is significantly associated with the spatial variation in tropical tree species richness. On a global scale it has previously been shown that canopy height explains a limited portion of the variation in tree species diversity, as such data provide information on the available niche space (Gatti, Di Paola, Bombelli, Noce, & Valentini, 2017). It has since been hypothesized that including information on the vertical canopy structure must explain more of the variation in tree species diversity than canopy height alone, as such data provide information on the occupation of the vertical niche space. Marselis et al. (2019) demonstrated that information on canopy height and vertical canopy structure, expressed as the plant area index (PAI) profile from full-waveform airborne lidar data, could be used to map tree species diversity in Gabon, Africa. However, it is not clear whether this relationship is of a similar nature and strength across different regions, or even the entire tropics. If existent, then, the use of such a structure–diversity relationship(s) could be applied at a pan-tropical scale with the rapidly increasing availability of spaceborne canopy structure information derived from the Global Ecosystem Dynamics Investigation (GEDI), a full-waveform spaceborne lidar system (Dubayah, Blair, et al., 2020). GEDI is expected to provide over 10 billion measurements of vertical canopy structure across the temperate and tropical forests between 2019 and 2021.

Factors influencing tree species diversity on a global scale differ from those affecting spatial patterns at regional or local scales. In general, tropical tree species diversity increases with increasing precipitation, forest stature, soil fertility, time since catastrophic disturbance, and rate of canopy turnover; and decreases with seasonality, latitude and altitude (Givnish, 1999). At large-grain scales historical biogeographical processes are more important, whereas at the plot scale environmental variables strongly influence diversity (Keil & Chase, 2019).

Similar to species diversity, forest structure at the global scale is influenced by interacting historic, environmental and human-related variables, precipitation in the wettest month being the most important single predictor of plant height (Moles et al., 2009). Forest structure measured in the field is mainly comprised of four variables: canopy height, biomass, basal area and tree density (Palace et al., 2015). However, active remote sensing techniques have revolutionized the study of canopy structure (Newnham et al., 2015). With lidar remote sensing, for example, it is now possible to obtain information on canopy height, as well as the position and amount of plant material along the vertical axis of the canopy (Tang et al., 2012). Palace et al. (2015) stressed that high resolution lidar data possess vertical structure information that is inherently linked to ecological processes.

We hypothesize that structure–diversity relationships will vary across different biogeographical and phylogenetic regions (Corlett & Primack, 2011; Slik et al., 2018) and that it may be more fruitful to develop multiple relationships rather than one pan-tropical relationship for operationalizing tree species diversity mapping with spaceborne active remote sensing data. Additionally, the strength of the relationship between a variable and tree species diversity often changes with resolution (plot size) as tree species diversity is not linearly related with area (species–area curve) (MacArthur & Wilson, 1967). This complicates the development of predictive models at specific resolutions, and also limits the extrapolation of estimates at one resolution to a larger area, which impedes the mapping of pan-tropical tree species diversity at high spatial resolution.

In sum, we know that both species diversity and canopy structure vary greatly within and across continents. Hence, our objective is to assess whether canopy structure information can explain tree species richness at the local, regional and/or pan-tropical scale with the ultimate goal to evaluate the efficacy of spaceborne full-waveform lidar for mapping tree species richness across the tropics. First, we compare characteristics of the vertical canopy structure, measured with full-waveform lidar data, for tropical forests across the world. Second, we evaluate the differences in species richness and species–area curves across the different study sites using field measurements. Third, we evaluate the potential for developing local (within 25–50 ha field plots), regional (within biogeographical regions) and pan-tropical structure–richness relationships, relating canopy structure metrics from lidar to tree species richness measurements from the field at three spatial resolutions (0.0625, 0.25 and 1.0 ha). Lastly, we discuss the potential of full-waveform lidar data from GEDI for mapping tree species richness across the tropics using structure–richness relationships.

2 | MATERIALS AND METHODS

We address the relationship between canopy structure and tree species richness in terra firme forest in the tropical region between 23.5° N and S. We compiled a field and lidar data set covering colonizing forest, old-growth tropical forest and forests under different degrees of degradation and savanna. We included such a wide variety of forest stages as most of the Earth’s tropical forests have been degraded or otherwise affected by natural and human influences (Lewis et al., 2015). Hence, when developing a method that allows for estimating pan-tropical tree species richness it is important to include data covering this range of possibilities.
Species diversity can be expressed with a variety of indicators. Generally, three levels of diversity are recognized: α-, β- and γ-diversity. α-diversity refers to the local diversity of a community, habitat or field plot. β-diversity refers to the differences in diversity between habitats and γ-diversity to the total diversity of a region (Colwell, 2009). In this study, we focus on α-diversity. α-diversity can be expressed with many different metrics. In this study, we focus on one dimension of species diversity: species richness (S) expressed as the total number of species in a plot of a given size. From here on forward, we only refer to tree species richness, used to express the local tree species diversity. We chose species richness as it is easy to interpret, and it can probably be used most directly by ecosystem managers. This measure of species diversity is sometimes referred to as species density as it does not consider the number of trees sampled in each plot.

2.1 | Field data sets

Field data were used to calculate the reference values of tree species richness. We used 15 data sets: one from Australia, two from Southeast Asia, six from Africa, three from South America and three from Central America (Figure 1). All field data sets used in this study have been previously collected and published and have coincident airborne lidar data available. Each field data set is labelled with a three-letter code and contains information on tree location, species, and diameter at breast height (DBH). All data sets were collected by different organizations and research teams resulting in different data characteristics (Table 1, Supporting Information Appendix S1). Four data sets consisted of one large plot of 25 ha (rob, Australia and rab, Gabon) or 50 ha (dan, Malaysia and bci, Panama). The other 11 data sets consisted of multiple (3–21) smaller plots with sizes ranging from 0.16 to 4.0 ha.

In this study, we assessed the structure–richness relationship at three spatial resolutions (1.0, 0.25, 0.0625 ha) because of the non-linear relationship between the number of tree species (S) and sampled area. We selected squares of 1.0 ha (100 m × 100 m) because they are often used in ecology and it has been shown that the spatial mismatch of plot location and remote sensing products is minimized at this resolution (Réjou-Méchain et al., 2014). We used squares of 0.25 ha (50 m × 50 m) because these yielded the best results describing the structure–diversity relationship in Gabon (Marselis et al., 2019), and squares of 0.0625 ha (25 m × 25 m) because they correspond to a resolution close to the GEDI footprint size. The data sets were used at one, two or three of the aforementioned resolutions depending on the original plot size and the availability of stem maps or subplots (Table 1, full table in Supporting Information Appendix S1). For each of the field sites, we calculated S for the entire data set and for each plot at each plot size (Table 2). Only live trees with a DBH ≥ 10 cm were included, to ensure consistency among data sets, and we included all plots of each resolution in which more than 80% of the trees were identified to at least the genus level.

2.2 | Lidar data sets

Each of the field data sets had coincident discrete return airborne laser scanning (ALS) data, or full-waveform lidar data from the Land Vegetation and Ice Sensor (LVIS), collected over the field plots within 5 years of field data collection. We used the GEDI simulator (Hancock et al., 2019) to create lidar waveforms from the ALS data over the field plots. The ALS data were originally collected with a variety of airborne instruments, but the GEDI simulator ensures a reliable GEDI-like waveform with minimal influence of the original instrument-specific characteristics. In this way, all lidar information could be processed consistently across all study sites ensuring a reliable inter-comparison of canopy structure metrics derived from the waveforms and allowing for easy transfer of the developed models to future on-orbit GEDI data. Lidar waveforms were simulated with a 22-m ground footprint (Gaussian distribution of laser energy, σ = 5.5 m). Lidar waveform locations were determined by filling each field plot, using the original field plot size and shape, with footprint centre locations 6.25 m from the plot edge and 5 m between footprint centre locations (Figure 2). This allowed a reliable measure of canopy structure to be acquired for each plot by averaging lidar metrics from all waveforms inside...
the plot, instead of using single waveforms in the plot centre and evaluating structure–richness relationships based on such potentially unrepresentative waveforms. The following information was extracted from each simulated lidar waveform using mature and published algorithms: canopy height (expressed as the 98th percentile of the relative height metric; RH98), total plant area index (PAI), and plant area index at a 1-m vertical resolution (Drake, Dubayah, Knox, Clark, & Blair, 2002; Hancock et al., 2019; Marselis et al., 2018; Tang et al., 2012). The 1-m vertical profile was used to compare the canopy structure across the study sites. It was aggregated into a 10-m vertical profile, summing all PAI values in each 10-m vertical bin, to be used in the structure–richness analyses. We chose to use the PAI profile because it is a biophysical variable describing the amount of plant material along the vertical forest axis, thus, directly indicating the occupation of vertical space. Marselis et al. (2019) previously showed this information relates well to tree species richness in Africa. The average of each of the resulting metrics from all waveforms within each plot was

<table>
<thead>
<tr>
<th>Country</th>
<th>Project code</th>
<th>No. native plots</th>
<th>Total area (ha)</th>
<th>Source/additional information</th>
</tr>
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<tr>
<td><strong>Oceania</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Australia</td>
<td><em>rob</em></td>
<td>1</td>
<td>25</td>
<td>Bradford, Metcalfe, Ford, Liddell, and McKeown (2014)</td>
</tr>
<tr>
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<td></td>
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<td></td>
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</tr>
<tr>
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<td><em>dan</em></td>
<td>1</td>
<td>50</td>
<td><a href="https://forestgeo.si.edu/sites/asia/danum-valley">https://forestgeo.si.edu/sites/asia/danum-valley</a></td>
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<tr>
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<td></td>
</tr>
<tr>
<td>DRC</td>
<td><em>mal</em></td>
<td>21</td>
<td>21</td>
<td>Bastin et al. (2015)</td>
</tr>
<tr>
<td>DRC</td>
<td><em>yan</em></td>
<td>9</td>
<td>9</td>
<td>Kearsley et al. (2013)</td>
</tr>
<tr>
<td>Gabon</td>
<td><em>rab</em></td>
<td>1</td>
<td>25</td>
<td><a href="https://forestgeo.si.edu/sites/africa/rabi">https://forestgeo.si.edu/sites/africa/rabi</a>; Memiaghe, Lutz, Korte, Alonso, and Kenfack (2016); Engone Obiang et al. (2019)</td>
</tr>
<tr>
<td>Gabon</td>
<td><em>mon</em></td>
<td>12</td>
<td>12</td>
<td>Fatoyinbo et al. (2017)</td>
</tr>
<tr>
<td>Gabon</td>
<td><em>mab</em></td>
<td>10</td>
<td>10</td>
<td>Bastin et al. (2015); Labrière et al. (2018)</td>
</tr>
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<td><strong>South America</strong></td>
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<td></td>
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</tr>
<tr>
<td>Brazil</td>
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<td>s12</td>
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<td>3.36</td>
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<tr>
<td><strong>Central America</strong></td>
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</tr>
<tr>
<td>Costa Rica</td>
<td>cha</td>
<td>3</td>
<td>2</td>
<td><a href="http://neoselvas.wordpress.uconn.edu/costa-rica/">http://neoselvas.wordpress.uconn.edu/costa-rica/</a></td>
</tr>
<tr>
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<td>bci</td>
<td>1</td>
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<td><a href="https://forestgeo.si.edu/sites/neotropics/barro-colorado-island">https://forestgeo.si.edu/sites/neotropics/barro-colorado-island</a>; Lobo and Dalling (2013)</td>
</tr>
</tbody>
</table>
TABLE 2 The total number of species identified at each study site and the average ($\bar{x}$) and standard deviation (SD) of the species richness for each of the three plot sizes expressed as $\bar{x} \pm$ SD [including only live trees with diameter at breast height (DBH) ≥ 10 cm]

<table>
<thead>
<tr>
<th>Country</th>
<th>Project code</th>
<th>Total no. species</th>
<th>Total sampled area used (ha)</th>
<th>Species richness 1.0 ha</th>
<th>Species richness 0.25 ha</th>
<th>Species richness 0.0625 ha</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oceania</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Australia</td>
<td>rob</td>
<td>205</td>
<td>25</td>
<td>98 ± 10</td>
<td>56 ± 8</td>
<td>27 ± 5</td>
</tr>
<tr>
<td>Southeast Asia</td>
<td></td>
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</tr>
<tr>
<td>Malaysia</td>
<td>dan</td>
<td>260</td>
<td>6</td>
<td>117 ± 13</td>
<td>51 ± 7</td>
<td>19 ± 4</td>
</tr>
<tr>
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<td>sep</td>
<td>517</td>
<td>32</td>
<td>102 ± 22</td>
<td>53 ± 11</td>
<td>-</td>
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<tr>
<td>Africa</td>
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<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>DRC</td>
<td>mal</td>
<td>116</td>
<td>21</td>
<td>37 ± 11</td>
<td>20 ± 7</td>
<td>-</td>
</tr>
<tr>
<td>DRC</td>
<td>yan</td>
<td>232</td>
<td>9</td>
<td>50 ± 23</td>
<td>24 ± 13</td>
<td>10 ± 6</td>
</tr>
<tr>
<td>Gabon</td>
<td>rab</td>
<td>234</td>
<td>25</td>
<td>84 ± 8</td>
<td>42 ± 6</td>
<td>17 ± 4</td>
</tr>
<tr>
<td>Gabon</td>
<td>lop</td>
<td>118</td>
<td>9.5</td>
<td>32 ± 22</td>
<td>17 ± 10</td>
<td>8 ± 4</td>
</tr>
<tr>
<td>Gabon</td>
<td>mon</td>
<td>146</td>
<td>12</td>
<td>32 ± 15</td>
<td>15 ± 9</td>
<td>7 ± 5</td>
</tr>
<tr>
<td>Gabon</td>
<td>mab</td>
<td>196</td>
<td>10</td>
<td>55 ± 8</td>
<td>-</td>
<td>-</td>
</tr>
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<td>South America</td>
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<tr>
<td>Peru</td>
<td>tam</td>
<td>517</td>
<td>6</td>
<td>171 ± 13</td>
<td>70 ± 9</td>
<td>24 ± 5</td>
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<tr>
<td>Brazil</td>
<td>s11</td>
<td>91</td>
<td>1.44</td>
<td>-</td>
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<td>17 ± 3</td>
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<tr>
<td>Brazil</td>
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<td>135</td>
<td>3.36</td>
<td>-</td>
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<td>16 ± 4</td>
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<td>Costa Rica</td>
<td>lsv</td>
<td>216</td>
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<td>-</td>
<td>48 ± 8</td>
<td>19 ± 5</td>
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<tr>
<td>Costa Rica</td>
<td>cha</td>
<td>81</td>
<td>2</td>
<td>58</td>
<td>28 ± 5</td>
<td>13 ± 4</td>
</tr>
<tr>
<td>Panama</td>
<td>bci</td>
<td>220</td>
<td>50</td>
<td>87 ± 8</td>
<td>42 ± 6</td>
<td>17 ± 3</td>
</tr>
</tbody>
</table>

computed to represent the canopy structure for each plot at each spatial resolution.

2.3 | Canopy structure across the tropics

To evaluate the canopy characteristics across the different study sites, we calculated the median plant area volume density profile (composed of the PAI values for each 1-m vertical bin), using all simulated lidar waveforms for each study site. In addition to the median (50th percentile), we calculated the 10th, 30th, 70th and 90th percentiles of the PAI values in the same 1-m vertical bins, to provide a representative distribution of the canopy structure across each study site.

2.4 | Species–area relationships across the tropics

We created species–area relationships, calculating the mean and standard deviation of $S$ for plot sizes ranging between 0.01 and 50 ha, to assess how species richness changes by plot size across our study sites. Each of the original field plots was filled with as many non-overlapping subplots as possible at 17 spatial resolutions (0.01, 0.0225, 0.04, 0.09, 0.16, 0.25, 0.36, 0.64, 1.0, 2.25, 4.00, 6.25, 9.00, 12.25, 16.0, 25.0, 50.0 ha) with each tree assigned to a subplot at each resolution. The plot sizes used at each study site depended on the original plot size and the availability of stem maps (Supporting Information Appendix S1). We visualized the mean and standard deviation of $S$ for each plot size at each study site to evaluate the differences in species–area curves across the tropics.

2.5 | Structure–richness analysis

To evaluate the existence of a relationship between vertical canopy structure and tree species richness across the tropics, we developed models at three scales: local, regional and pan-tropical, because many historical and environmental drivers of (tree) species diversity have stronger or weaker relationships depending on the scale of observation (Gaston, 2000; Keil & Chase, 2019) as do different ecosystem functions (Chisholm et al., 2013). Definitions of the scales are presented in the following sections.

2.5.1 | Local analysis

The local analysis focused on the structure–richness relationship within large (25 or 50 ha) plots. We used data from adjacent field plots to evaluate the relationship between $S$ and the canopy structure expressed as canopy height (RH98), total PAI and vertical canopy profile...
2.5.2 Regional and pan-tropical analysis

The regional analysis was focused on the structure–richness relationship based on non-adjacent plots across study sites within the same biogeographical zone. We evaluated different combinations of study sites at three spatial resolutions (Table 3). To prevent the large plots from dominating the regional and pan-tropical analyses, we thinned their contribution to both the regional and pan-tropical data sets. From the 25-ha plots, we selected 1.0-ha plots at each corner, and from the 50-ha plots, we selected all corner and the middle plots along the long sides of the plot (six 1.0-ha plots total). To avoid mixing local and regional effects, we employed a Monte-Carlo simulation approach in which we drew different samples from the full regional data set. In each Monte-Carlo run, we randomly sampled one plot at the given resolution from each original plot location (especially important at the 0.25 and 0.0625 ha resolutions at which up to 16 plots exist at the location of each original 1.0-ha plot) and applied a cross-validation (80/20) or leave-one-out cross validation (if n ≤ 25) approach. In the cross-validation, we again performed a two-step approach: first, we performed variable selection on the Poisson regression model choosing the model with lowest BIC (using the bestglm (McLeod, Xu, & Lai, 2020) package in R Core Team (2020)), and then, built the predictive model with the chosen variables. We applied the model to the test data and calculated the model performance statistics for each fold according to Piñeiro et al. (2008).

The pan-tropical analysis focused on the structure–richness relationship combining the information from all 15 study sites across all tropical regions, in other words, it was a special case of the regional analysis in which data from all sites were included. Thus, the same methods were applied as in the regional analysis.

3 RESULTS

3.1 Vertical forest structure across the tropics

The vertical canopy structure of forests, in terms of the vertical distribution of plant material, varies between tropical regions (Figure 3). Maximum canopy height in our study sites in the Neotropics and Central Africa is typically around 40 m, and slightly lower in Australia, while canopy heights in Southeast Asia exceed 60 m. Many sites show a distinct understorey layer and a decrease in plant material through the canopy. Relative to the understorey, the canopy layer sharply declines in vegetation density (sep and dan, Malaysia) or steadily declines along the vertical axis (bci, Panama; rab, Gabon; mal, DRC; rob, Australia). This vertical distribution of declining vegetation is exacerbated in degraded forests: in s11, s12 (Brazil) and mon (Gabon), where the bulk of the vegetation exists close to the forest floor at c. 5 m height, but remnant trees in some plots may reach 40 m. Other sites, especially undisturbed ones, have distinct canopy layers. In tam (Peru) and in the old-growth forest in lsv (Costa Rica) there are multiple peaks of high-density vegetation across the vertical strata of the forest. The profiles of yan (DRC) and lop (Gabon) are characterized by a multiple-peak pattern, with one peak 20–30 m in the canopy and another within 5 m of the ground, reflecting the inherent structure of the forest–savanna mosaic. The less disturbed mab (Gabon) forest shows high variability in canopy structure between plots (e.g. the wide shaded area in Figure 3).

3.2 Species–area relationships

The number of species increases with plot size, but the rate of increase varies across study sites (Figure 4). For example, in rob
<table>
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</table>
(Australia) 82–117 species occur in a 1.0-ha plot compared to 16–44 species in 0.0625-ha plots. By contrast, tam (Peru) contains 154–185 species/ha, but only 11–35 species in a 0.0625-ha plot, similar to rob. Thus, species’ composition of adjacent 0.0625-ha plots in tam must be more dissimilar from each other than adjacent 0.0625-ha plots in rob (Australia), in other words, the β-diversity of the plots in tam is higher than in rob. The species–area curves vary in shape across study sites, with the highest total species richness in tam and lowest species richness in the African sites (Figure 4). Curves that are initially steep and decrease in slope at larger plot sizes indicate a high α-diversity but a lower β-diversity (e.g. when the area is increased, the same species are encountered).

3.3 | Structure–richness relationships

Pulling together the information on tree species richness and canopy structure (RH98 and total PAI), species richness generally increases with increasing canopy height and increasing total plant area index across the tropics (Figure 5).
The cross-validation results of the local models reveal weak structure–richness relationships. Of the three large plots (25 and 50 ha), only the models for bci (50 ha) show evidence of a significant relationship between the predicted and observed values ($R^2 = .32$ at 1.0 ha, Supporting Information Appendix S2). Even though species richness within all three large plots can be predicted with a root mean squared error between 7% and 20% of the mean species richness, the low RMSD% found only indicates that the predictions at the local scale are close to the mean species richness; however, in rab and rob the canopy structure is insensitive to the local variation in tree species richness (see for example Supporting Information Appendix SI2).

Regional structure–richness models generally show much better performance (Figure 6) than the local models in terms of the variance in species richness that can be explained with the canopy structure information (mostly significant models and higher $R^2$ values). However, prediction error (as percentage of the mean species richness) is generally higher, partly due to the larger range in species richness in these regional data sets. Regions of Africa and South America (Table 3) show the best model performance whereas regions including the Costa Rica data sets show much poorer performance (regions indicated with centralamerica). Results from an additional analysis on the compositional similarity (Bray–Curtis; Faith, Minchin, & Belbin, 1987; Supporting Information Appendix S3) of the Costa Rica data set showed that, even though species richness varies in Costa Rica (Table 2), the plots share many species, that is, the composition is similar. In the africa and southamerica data sets the variation in species richness is accompanied by a much larger variation in species composition (Supporting Information Appendix S3). The variation of the model performance for seasia is very high because of the low number of plots available for this region and at the 0.25-ha resolution it was not possible to create a significant model for > 95% of the Monte-Carlo iterations (Table 3). The model performance does not provide clear results on the effect of the different resolutions, given the overlapping error bars for models in the same region at multiple resolutions and the inability to create each regional model at each spatial resolution (Figure 6).

Pan-tropical structure–richness models show varying performance across the spatial resolutions with mean $R^2$ ranging between .25 and .39 and RMSD% between 66% and 43% for the plot sizes from 1.0 and 0.0625 ha (Figure 7). However, the error bars of the model performance at different resolutions are overlapping, indicating that no resolution has a statistically better performance. Around 39% of the variation in tree species richness can be explained using canopy structure metrics alone at the 0.25-ha resolution at the pan-tropical scale. Sites with extremely high values of observed species richness are generally predicted poorly (Supporting Information Appendix S4).

4 | DISCUSSION

4.1 | Structure–richness relationships across scales

In this study, we explored the relationships between vertical canopy structure and tree species richness at different resolutions across local, regional and pan-tropical scales, using a total of 15 study sites with coincident lidar and field data across the tropics. We found weak relationships between canopy structure and tree species richness at the local scale and the strongest relationship at the regional scales in Africa and South America. We also found significant relationships between canopy structure and tree species richness combining the data from all study sites across the tropics.

At the local scale, within one large plot inside one forest type, the variation in the canopy structure is determined mostly by variability in growth structure within the same species (the 25 and 50 ha plots have a similar composition throughout the plot, Supporting Information Appendix S1 and S3). For example, an adult tree of species X may range in height from 20–40 m, so even though the canopy structure may differ between two plots of similar composition, the difference is not attributed to a difference in species composition.
Furthermore, if a 20 m and 40 m tree of species X exist in the same plot, due to the difference in canopy structure the model may predict a species richness of 2 based on variation in structure. On the contrary, as area increases it is more likely that the difference in structure is caused by a difference in composition. Do keep in mind that structure can also change due to other variables such as topography, soil and microclimate. Individuals of most tropical forest species are spatially aggregated (Condit et al., 2000) so the composition of two adjacent plots is more similar than the composition of two more distant plots. This is the case for bci, where a 50-ha area with a species richness gradient was sampled (Fricker, Wolf, Saatchi, & Gillespie, 2015) and included in the local analysis, which led to more successful prediction of species richness based on structure. Within the 25-ha plots sampled at rab and rob, the variation in composition is smaller and no significant structure–richness relationships were found (Supporting Information Appendix S3).

Increasing the scale, we found that regions consisting of sites exhibiting a large variation in species composition among plots, but with a similar biogeographical history, show a much stronger structure–richness relationship. However, we note that model performance
differed quite drastically across regions. The forest in lsv, Costa Rica, consists of largely similar species composition, whereas species composition is much more varied in regions where the structure–richness models perform better (South America, Africa), supporting the result from local scale models that species richness can be better predicted from canopy structure in areas with greater \( \beta \)-diversity.

At the pan-tropical scale, we find a significant relationship between canopy structure and tree species richness across all spatial resolutions. At the intermediate resolution (0.25 ha) this relationship appears to be slightly stronger than at the higher and lower resolutions, but no significant difference was found. However, the observed difference may be attributed to the lower sensitivity of species richness to rare species at smaller plot sizes. For example, tam (Peru) plots have very high species richness at the 1.0-ha resolution (Table 2), whereas at the 0.0625-ha resolution the species richness ranges between 11 and 35 species, which is still higher than most other sites but much less than at the 1.0-ha plot size. Because the 1.0-ha plot size captures more rare species in each plot, the 1.0-ha pan-tropical model predictions for tam contain highly erroneous predictions that are not present in 0.0625-ha models (Supporting Information Appendix SI4). Rare species do not contribute much to the canopy structure, thereby complicating the relationship between structure and richness at a scale at which they contribute largely to species richness numbers.

4.2 | Limitations

This research could be significantly improved by using more coincident lidar and field data to thoroughly evaluate the existence and strength of the structure–richness relationship across all tropical regions. However, the collection of such data is costly and...
time-consuming. Here, we were able to exploit 15 independently collected data sets (Supporting Information Appendix S1), but data gaps exist, especially in the Amazon basin, high biomass forests of Central Africa, the mainland of Southeast Asia, New Guinea and Australia as well as the dry tropics and montane ecosystems. Apart from the spatial representation problem, the low number of plots for certain regions likely influences the observed variability in model performance. The pan-tropical models (with n ≥ 90) show more stable performance than models of regions with low numbers of plots (e.g. seasia). A training data set that does not fully represent the range of structure in the full data set can lead to biased predictions for some of the test plots. Such errors are exacerbated by the logarithmic link model in Poisson regression because errors can increase exponentially. Even so, negative predictions are possible with linear regression and the risk of underestimating tree species richness is higher for diverse areas. Hence, we chose to use Poisson regression, knowing that it may lead to extreme predictions in some cases that should be accounted for when operationalizing this method.

Species diversity can be identified in many different ways (Colwell, 2009; Gotelli & Colwell, 2001) and there are risks and pitfalls using just one metric. In this study, we only used ‘species richness’ (S), defined by the number of different tree species in a defined area (the plot, with different sizes), as this metric is easy to interpret and a prediction of the number of species/area can probably be used most directly by ecosystem managers. Hereby, we did not control for the number of stems in the plot, nor for the abundance of the different species. Such information can be considered, for example, by using the Shannon diversity index or rarefaction curves. Moreover, depending on the type of metric, a different model may need to be selected to describe the structure–richness relationship as different metrics are related differently to canopy structure information. For example, a generalized linear regression with a Poisson error distribution, as used here, is more suitable for estimated tree species richness values as these are count data, whereas a linear model with a Gaussian error distribution will be better suited for estimating Shannon diversity. Hence, we chose to focus on one metric of diversity to test the structure–richness relationships, while acknowledging other metrics may provide better, worse, or more useful predictions of tree species diversity and these should be considered in the future.

This study serves as a first attempt to study the pan-tropical structure–richness relationship and should be improved and further developed when more data become available. Additionally, the characteristics of each data set differed widely because all data were collected by different researchers and institutions. We accounted for this as much as possible by using data sets only at reliable plot and subplot resolutions, including only trees ≥ 10 cm DBH and including only plots with less than 20% of unidentified trees at the genus level. Nonetheless, we acknowledge that the quality of the species identification varied and may have affected our models as species identification in the tropics can be challenging due to the vast variety of tree species and the fact that new species are still encountered. Species identification of new and existing data could be improved using more botanists or genetic tests in the lab, which has been done for some of the data sets used here, but is not yet feasible for all data sets. Additionally, including information on species for trees with DBH ≥ 10 cm omits the (large) diversity found in the understory. Fricker et al. (2015) showed that especially this diversity variation in small trees related well to the canopy structure. Future research should examine if these findings are consistent across the tropics.

The availability of stem maps and subplots in each study site determined the spatial resolutions at which data sets could be used. This resulted in the inclusion of different data sets for each region (Table 3). This makes the comparison of model performance in the same region at different resolutions unreliable because the models were not always built on the same data (plots and study sites), but we weighed this decision to maximize the sizes of the data sets used to build the structure–richness models. Hence, no conclusion can be drawn about the optimal resolution for the structure–richness relationships.

Accurate geolocation of field plots is key for the development of reliable species-richness models (Fricker et al., 2015). However, geolocation of field plots in tropical forests can be challenging due to difficulties receiving a reliable GPS signal under dense canopy. This should be taken into account, especially when evaluating the performance of models built with small field plots, where the effects of such geolocation errors will be larger (Réjou-Méchain et al., 2014).

We included data from a range of forest stages, including old-growth forest, successional stages, disturbed forest and even low tree density savanna sites. The relationships we found are partially driven by this gradient (Figure 5). However, we deemed it essential to include data from across this range of forest types, because if this method is to be operationalized using canopy structure information from across the tropics, we will encounter all these different stages of forest (Lewis et al., 2015). We acknowledge that climatic, edaphic and topographic variables could also impact tree species richness across the tropics, such as mean annual temperature and precipitation (Keil & Chase, 2019) and slope and elevation (Robinson et al., 2018). However, in this study, we specifically focused on the relationship between canopy structure and tree species diversity, in light of the recently launched GEDI mission. We recognize that including such information on topographic and environmental variables may further improve the mapping of tree species richness across the tropics.

### 4.3 Future research & applications

Our results provide confidence regarding the existence of regional and pan-tropical structure–richness relationships that may be used to map pan-tropical tree species richness. The most accurate predictions seem to be achieved at the regional scale when adequate data are available and when forested areas are grouped by regions of similar biogeographical history. However, in the absence of such data it
may be of more immediate interest to further develop pan-tropical models, as these have been shown here to explain up to 39% of variation in tree species richness. At the time of writing, GEDI is collecting canopy structure information close to the finest resolution tested here (0.0625 ha) and thus, these data may be well suited for mapping tree species richness across the tropics. GEDI is a sampling mission in which lidar waveforms with 25-m diameter footprints are collected across eight tracks with 600-m between-track spacing, 60-m along-track spacing (Figure 8). By the end of its nominal 2-year mission, GEDI will have sampled roughly 4% of total land area.

The footprint-level GEDI information on vertical canopy structure is stored in the Level-2 data products that are publicly available from the NASA Land Processes Distributed Active Archive Center (LPDAAC; https://lpdaac.usgs.gov; Dubayah, Hofton, et al., 2020; Dubayah, Luthcke, et al., 2020; Dubayah, Tang, et al., 2020). GEDI gridded data products will have a 1-km² or finer resolution (Dubayah, Blair, et al., 2020). Our local scale models show that predictions of adjacent 0.0625-ha plots (or in the future, footprints) are on average correct, but they will not detect local nuances in species richness within forests of uniform composition. We suggest that the species richness predictions could potentially be used in a similar way as gridded GEDI data products by estimating the average number of species/0.0625 ha within a 1-km² cell, as such information may still be of interest to local land managers. Given the variable species-area relationships, it is not easy to translate species richness predictions at 0.0625-ha resolution to the expected number of tree species in 1 km². Also, the amount of variance in species richness explained is limited. Therefore, we propose two future research avenues of interest: fusion with spectral and/or radar data and using an environmental framework. Both spectral data and radar data have previously been shown to predict some of the variance in tree species richness (Bae et al., 2019; Bongalov et al., 2019; Boody & Cutler, 2006; Marselis et al., 2019; Schäfer et al., 2016; Wolf et al., 2012) and may improve our models and allow for more accurate predictions of tree species richness across the tropics and the creation of wall-to-wall data products at higher spatial resolution. Especially data from the hyperspectral imager suite (Matsunaga et al., 2013) instrument, that is soon to be launched to the International Space Station, the radar BIOMASS mission (Le Toan et al., 2011), the Ice, Cloud and land Elevation Satellite 2 mission (Duncanson et al., 2020), the TerraSAR-X add-on for Digital Elevation Measurement mission (Q), Saarela, Armstrong, Stahl, & Dubayah, 2019) and Landsat (Saarela et al., 2018), may be highly relevant for such applications. Alternatively, we believe that the inclusion of structural data within previously developed environmental and biogeographical frameworks will help to predict tree species diversity (Keil & Chase, 2019) as such frameworks already display intrinsic differences in tree species diversity. Such frameworks could benefit from GEDI lidar data providing information on the occupation of the vertical niche space and likely improve predictions of tree species richness across the tropics, which could then be compared to existing predictions such as from Slik et al. (2015). Moreover, it has previously been shown that lidar data can provide interesting information about the diversity of other taxa as well (Huang, Swatantran, Dubayah, & Goetz, 2014; Rappaport, Royle, & Morton, 2020) and future avenues for using lidar data to provide information on a holistic measure of species diversity, including many taxa, could be of incredible value.
CONCLUSIONS

In this study, we evaluated the existence of local, regional and pantropical relationships between vertical canopy structure and tree species richness in the tropics at three spatial resolutions: 1.0, 0.25 and 0.0625 ha. Full-waveform lidar data provide detailed information on the differences in vertical canopy structure between forests across the tropics. Our results show that canopy structure can explain a significant percentage of variation in tree species richness across different biogeographical regions. A full set of regional structure–richness models will most likely aid accurate pan-tropical species richness mapping, but the development of such a set of models is contingent on the availability of sufficient coincident field and lidar data across the tropics. Using one single predictive model at a pantropical scale, 39% of the variation in tree species richness could be explained using the vertical canopy structure. Given this canopy structure is measured directly from GEDI waveforms at the footprint level, this provides an interesting avenue for mapping tree species richness at high spatial resolution. Alternatively, canopy structure information from GEDI could be included in existing modelling frameworks, combining structural with spectral, environmental and topographic information to create more accurate tree species richness predictions.

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DATA AVAILABILITY STATEMENT

Most of the field and lidar data used in this study are available and can be downloaded directly from the internet. Otherwise the data
sets can be requested as described below. We have grouped the data in four groups: (a) LVIS lidar data, (b) ALS lidar data, (c) field data and (d) GEDI lidar data.

(a) LVIS lidar data

The LVIS data for the rab, lop, mon and mab study sites can be downloaded from the NASA data archive at the following https://doi.org/10.3334/ORNLDAAC/1591

The LVIS data for the cha and lsv study sites are available on the following website: https://lvis.gsfc.nasa.gov/Data/Maps/CR2005Map.html

(b) ALS lidar data

The ALS data over rob are available through the auscover data portal ftp://qld.auscover.org.au/airborne_validation/lidar/robsoncs_creek/

The ALS data over si1 and si2 can be downloaded from the sustainable landscapes data portal http://www.paisagenslidar.cnptia.embrapa.br/webgis/

The ALS data over yan and mal are available through ArcGIS online at https://www.arcgis.com/home/item.html?id=a6095e77541d4ad88dc6f0945639dd089

The ALS data over bci can be downloaded directly using the following download link: http://www.life.illinois.edu/dalling/lidar_data.tgz

The ALS data over tam are not publicly available online as they are actively supporting external research projects. However, anyone interested in working with this data can contact Chris Hopkinson (c.hopkinson@uleth.ca) or Ross Hill (rhill@bournemouth.ac.uk) to request access.

The ALS data over dan and sep are currently in the process of being made available through the Centre for Environmental Data Analysis (CEDA) https://www.ceda.ac.uk/

(c) Field data

Field data from rob have been published through the Terrestrial Ecosystem Research Network (TERN) data portal linked from https://supersites.tern.org.au/supersites/fnrq-robson

The dan, rob and bci field data are available on request through the Forestgeo website at https://forestgeo.si.edu/explorer-data: https://forestgeo.si.edu/explorer-data/rabitermconditionsrequest-form, https://forestgeo.si.edu/explorer-data/barro-colorado-island-termconditionsrequest-forms, https://forestgeo.si.edu/explorer-data/danum-valley-termconditionsrequest-forms

The sep, lop, tam and yan field data are all available upon request through forestplots.net and can be found under the project names ‘sepilok’, ‘lope’, ‘tambopata’ and ‘yangambi’ at https://www.forestplots.net/en/

The mon field data are archived through the NASA data archiving center and available at https://doi.org/10.3334/ORNLDAAC/1580

The si1 and si2 data were available through the data portals of the sustainable landscapes projects and can be found under the field data from the São Félix do Xingu region collected in 2011 and 2012 in the following data portal: http://www.paisagenslidar.cnptia.embrapa.br/webgis/

(d) GEDI lidar data

The different lidar data products from GEDI used to create Figure 8 can be downloaded through https://doi.org/10.5067/GEDI/GEDI01_B.001, https://doi.org/10.5067/GEDI/GEDI02_A.001 and https://doi.org/10.5067/GEDI/GEDI02_B.001

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REFERENCES


**BIOSKETCH**

Suzanne Marselis received her PhD degree in Geographical Sciences at the University of Maryland. She has a broad interest in the application of remote sensing data for mapping different aspects of biodiversity across the tropics and has an extensive background in the use of lidar data for mapping three-dimensional canopy structure.

**SUPPORTING INFORMATION**

Additional supporting information may be found online in the Supporting Information section.