Species diversity and stand structure as drivers of canopy complexity in southern African woodlands

John L. Godlee

26th August 2021

1 Abstract

Atmospheric CO₂ enrichment and human-induced climate change are expected to drive woody encroachment and an increase in tree cover across African savannas, with consequences for ecosystem function, particularly related to carbon dynamics. The patch dynamics of savanna-woodland mosaics are complex however, as woody growth is mediated by seasonal fire that is itself driven by properties of the woody overtstorey. It is unclear how variation in tree species composition and stand structure in this ecosystem affects woody canopy complexity, and how this might determine future vegetation dynamics. Here, I conducted a study of canopy structure in southern African savannas using terrestrial LiDAR, at sites in Bicuar National Park, Angola and Mtarure Forest Reserve, Tanzania, to explore relationships between tree species diversity, species composition, the spatial distribution of trees, variation in tree size and canopy complexity. I found consistent weak positive effects of species diversity on plot scale canopy complexity metrics related to canopy density, but a negative effect on metrics related to the spatial heterogeneity of the distribution of canopy material. Species diversity caused an increase in canopy height, canopy closure, and within-canopy structural complexity. However, stochasticity in neighbourhood scale woody structure masked species diversity effects at small spatial scales. Finally I found that spatial clustering of trees in space led to a reduction in canopy closure, even within clustered areas, suggesting that disturbance by fire and herbivory not only reduce canopy cover at the landscape scale, but also reduce canopy cover at smaller spatial scales But how does disturbance lead to clustering? Could just say "suggesting"

$_{22}$ 1 Introduction

2

3

5

6

8

10

11

12

13

14

15 16

17

18

19

20

21

Atmospheric CO₂ enrichment, coupled with climate change and changing disturbance regimes, is expected to drive woody encroachment, i.e. proliferation of trees in previously non-wooded 24 areas, and increased growth of trees in currently wooded areas, across the savanna biome over 25 the coming century (Criado et al., 2020; Mitchard & Flintrop, 2013; Stevens et al., 2016). 26 As atmospheric CO₂ concentrations increase, C₃ trees are expected to gain a competitive 27 edge over C₄ grasses due to differences in photosynthetic pathway and carbon use efficiency 28 (Buitenwerf et al., 2012), allowing trees to increase their productivity. This is expected to have 29 cascading effects on canopy cover, grass growth, and disturbance regime (Bond & Midgley, 30 2012). If realised, woody encroachment and woody densification will have significant effects 31 on the global carbon cycle, as more CO₂ is stored in woody biomass, as well as myriad other 32 effects on ecosystem structure (Donohue et al., 2013). Indeed, tropical savannas have been 33 identified as the fastest increasing component of the terrestrial carbon sink (Sitch et al., 2015). 34 Previous studies however, have reported wide variation in rates of woody encroachment and 35 densification (Mitchard & Flintrop, 2013), particularly in disturbance-prone savannas such as 36 miombo woodlands in southern Africa (Axelsson & Hanan, 2018), and it is unclear how the 37 fertilisation effect of atmospheric CO₂ enrichment will interact with other ecosystem properties 38 to alter vegetation structure (Körner, 2017; Reich et al., 2014). 39

- 40 Savanna vegetation is defined by the coexistence of trees and grasses (Scholes & Archer, 1997).
- 41 In the tropical mesic savannas of southern Africa, disturbance by fire and herbivory are the main

limitations on tree cover, preventing the competitive exclusion of shade-sensitive C₄ grasses where climatic conditions would otherwise allow for closed canopy forest (Sankaran et al., 2005). 43 C₄ grasses also provide the main fuel source for seasonal fires in these savannas (Frost, 1996), 44 producing a positive feedback where an increase in tree cover reduces grass fuel load, reducing 45 fire frequency and intensity, increasing tree cover, and so on (Staver & Koerner, 2015). As such, 46 even small perturbations in tree cover can lead to large changes in vegetation structure if critical 47 thresholds of tree cover are crossed (Hirota et al., 2011). Previous research has sought to identify 48 environmental factors which affect tree cover and its responses to atmospheric CO₂ enrichment, 49 but few have considered the functional role of the existing tree community and its effect on 50 ecosystem processes. 51

Canopy structure describes the spatial distribution and density of tree canopy foliage (Lowman & 52 Rinker, 2004). Canopy structural complexity, i.e. the spatial heterogeneity of foliage distribution 53 within the canopy, has been linked to increased net ecosystem productivity (Baldocchi & Wilson, 54 2001; Chen et al., 2012; Gough et al., 2019; Hardiman et al., 2011; Law et al., 2001; Morin, 2015), 55 increased resilience of productivity (Pretzsch, 2014), reduced understorey light penetration (Fotis 56 et al., 2018; Scheuermann et al., 2018), and greater moderation of understorey micro-climate 57 (Wright et al., 2017). Furthermore, in temperate and boreal forests, functional differences among 58 coexisting tree species in their vertical and horizontal canopy occupation provides a link between 59 species diversity, canopy structural complexity and canopy density, with canopy complexity 60 constituting a mechanism for observed positive biodiversity-ecosystem function effects in wooded 61 ecosystems (Barry et al., 2019; Pretzsch, 2014). In tropical savannas, tree species diversity might 62 therefore influence ecosystem-level woody thickening in response to elevated atmospheric CO₂, 63 where competition effects in diverse tree communities are reducted due to niche separation, and 64 can more effectively increase foliage density and reduce understorey light penetration, excluding 65 grass and thus reducing the probability of disturbance. 66

As well as the species diversity of trees, the spatial distribution and relative size of tree stems, 67 i.e. stand structure, is also expected to affect canopy structural complexity (Stark et al., 2015). 68 Heterogeneity in stem size, whether a result of species diversity, disturbance history or some other 69 factor, is expected to increase canopy complexity and canopy density as individuals of different sizes occupy different parts of the vertical canopy space (Panzou et al., 2020), and may differ in 71 light requirements (Charles-Dominique et al., 2018). Additionally, clustering of individuals in 72 space is expected to increase canopy structural heterogeneity across the wider savanna landscape, 73 but ultimately decrease total foliage density due to an increase in competitive interactions (Dohn 74 et al., 2017). Clustering may occur as a result of disturbance history, facilitation effects among 75 individuals in stressful environments (Ratcliffe et al., 2017), or due to other limitations on 76 establishment arising from growth strategy (). More diverse communities may allow greater stem 77 density and greater foliage density, as differences in canopy occupancy among species reduce 78 negative effects competition among individuals on growth (Gough et al., 2019). 79

Functional differences among floristic types of savanna may also drive variation in canopy 80 complexity, irrespective of species diversity. Some savanna trees form denser canopies than 81 others, as a result of variation in leaf size and branch architecture. Previous studies have 82 compared the branch architecture of ex-Acacia (e.g. Senegalia and Vachellia spp.) and miombo 83 (e.g. Julberardia, Brachystegia, and Isoberlinia) archetypal tree species. While ex-Acacia species 84 tend to inhabit drier, heavily grazed areas, miombo species tend to inhabit dystrophic wetter 85 areas structured heavily by fire (Ribeiro et al., 2020). These studies have shown that ex-Acacia 86 species develop sparser canopies, cagey branch architecture, and wider spreading crowns, while 87 dominant Fabaceae species from the miombo develop thicker, taller canopies, and can grow to 88 large trees (Archibald & Bond, 2003; Mugasha et al., 2013; Privette et al., 2004). Similarly, 89 dominant miombo Fabaceae species from the Detarioideae subfamily have been shown to exhibit wider crowns and grow taller than coexisting species from the Combretaceae family (). Shenkin

et al. (2020) showed that Fabaceae tree species from tropical forests exhibit wider and more voluminous tree crowns than other common families of tropical trees. Under identical stem densities, miombo woodland species may therefore exclude grass more effectively than ex-Acacia or Combretaceae species given these differences in growth form.

Canopy complexity is multi-dimensional and has previously been explained using a plethora 96 of simple metrics that originated in forest and community ecology (Kershaw et al., 2017). 97 Assessments of canopy complexity have most often modelled tree canopies as a series of ellipses 98 (2D), ellipsoids or cones (3D) based on field measurements with measuring tapes (Jucker et al., 99 2015), or used surrogate proxies for 3D canopy structure, due to its inherent complexity (Seidel et 100 al., 2011). Measurements of this kind are time consuming and yet remain an over-simplification of 101 canopy structure. Alternatively, canopy closure is often measured using indirect optical methods 102 which partition sky from canopy material, i.e. with hemispherical photography or the commonly 103 used LAI-2000, providing a 2D representation of the canopy but lacking information on vertical 104 canopy structure (Jonckheere et al., 2004). In recent years, particularly in temperate and boreal 105 forests, LiDAR (Light Detection And Ranging) has emerged as a suitable technology for rapidly 106 and precisely assessing canopy structure in 3D, conserving information on 3D structure of the 107 calibre that is required to understand it's complexities (Calders et al., 2020; Muir et al., 2018). 108 In tropical savannas, very few studies have used terrestrial LiDAR for vegetation analyses, and 109 in southern Africa all existing studies have been located at the Skukuza Flux Tower in Kruger 110 National Park, South Africa (Muumbe et al., 2021). Pioneering work describing the ecology of 111 southern African savannas placed large emphasis on canopy structural diversity as a mediator of 112 ecosystem function (Solbrig et al., 1996), but much of that understanding of savanna vegetation 113 structure was derived from traditional mensuration methods. Using terrestrial LiDAR to measure 114 canopy complexity in southern African savannas therefore offers a unique chance to validate 115 accepted theory and describe differences in ecosystem structure among savanna vegetation types 116 in finer detail than previously possible. 117

In this study I applied terrestrial LiDAR techniques to woodland-savanna mosaics at two sites in southern Africa, with the aim of increasing understanding of how various measures of tree canopy complexity relate to tree diversity and stand structure. I hypothesise that tree neighbourhoods with greater tree species diversity, greater heterogeneity in stem size, and greater heterogeneity in stem location, allow greater canopy complexity and foliage density. Thus, more diverse savannas might exhibit a higher potential woody biomass, greater productivity, and more effectively increase their growth under elevated atmospheric CO₂, promoting woody thickening. I also consider the functional differences in canopy architecture among tree communities and how this affects canopy closure and total canopy occupancy.

2 Materials and methods

128 2.1 Study sites

118

119

120

121

122

123

124

125

126

127

Field measurements were conducted at two sites, Bicuar National Park, in southwest Angola (S15.1°, E14.8°), and Mtarure Forest Reserve, in southeast Tanzania (S9.0°, E39.0°) (Figure 1).
At each site, 1 ha (100×100 m) plots were located in areas of savanna-woodland vegetation, across a gradient of stem density and a range of savanna floristic archetypes. In Angola, 15 plots were sampled, while in Tanzania, seven were sampled following the curtailment of fieldwork due to COVID-19 travel restrictions. Fieldwork was conducted between February and April at both sites, during the peak growth period of each site in order to capture the maximum foliage volume in the canopy.

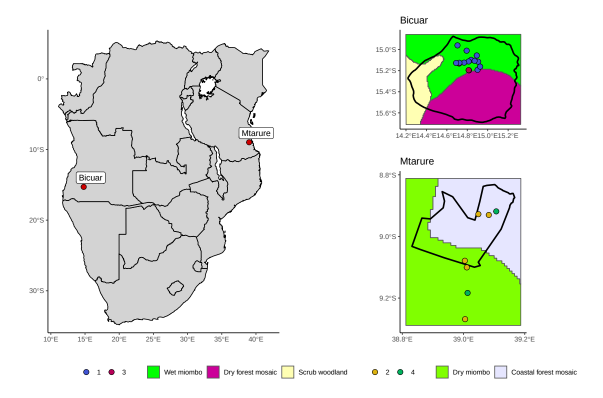


Figure 1: Location of study sites within southern Africa (left), and of 1 ha plots within each site (right). The black outlines in each site map denote the boundaries of protected areas which encompass the majority of study sites, Bicuar National Park in Angola (top), and Mtarure Forest Reserve in Tanzania (bottom). The background of each site map is a re-classified version of White's vegetation map (White, 1983). Points in site maps are shaded according to vegetation type identified by hierarchical clustering of tree genera abundances. Note that all maps are on different scales.

137 2.2 Field measurements

Within each 1 ha plot, each woody stem ≥ 5 cm stem diameter was identified to species, the 138 stem Diameter at Breast Height (DBH) was measured at 1.3 m above the ground, and the stem 139 location within the plot was recorded using tape measures. Each 1 ha plot was sampled by nine 140 10 m diameter circular subplots arranged in a regular grid, with a 15 m buffer from the plot edge 141 and 35 m between subplots. For each subplot, the distance and direction from the subplot centre of each stem >5 cm diameter with canopy material inside the subplot was recorded. Within each 143 subplot, a variable number of scans were recorded using a Leica HDS6100 phase-shift Terrestrial 144 Laser Scanner (TLS). The number and position of scans within a subplot was determined by the 145 arrangement of canopy material in the subplot, to minimise shadows within the canopy of the 146 subplot, and to maximise canopy penetration. The number of scans per subplot ranged between 147 one and five across both sites. Extended field methods and data analysis methods are described in Chapter 6. 149

150 2.3 Data analysis

151

175

2.3.1 TLS processing

Point clouds from scans in each subplot were registered and unified using Leica Cyclone (version 152 9.1), using five reflective cross targets visible to all scans as anchor points. Point clouds were 153 voxelised to cubic voxels of different sizes depending on the application of the data. Subplot 154 height profile estimation and gap fraction was conducted using 5 cm³ voxels, while whole plot 155 canopy rugosity was estimated using 50 cm³ voxels. Voxels were classified as 'filled' if they 156 intersected one or more points. Variation in voxel size reflects the spatial scale of each analysis, 157 and is bounded by the beam divergence of the scanner over longer distances (Cifuentes et al., 158 2014). Choosing voxels that are too small can result in pock-marked representations of surfaces 159 that are especially problematic when calculating larger scale canopy complexity metrics such as 160 canopy top roughness, while voxels that are too large can result in an over-estimation of plant 161 volume when estimating canopy foliage density at the subplot scale (Cifuentes et al., 2014; Seidel 162 et al., 2012). 163

The noise reduction algorithm from Rusu et al. (2008) was used to discard points based on 164 mean nearest neighbour distances, with a mean number of neighbours of eight, and a standard 165 deviation threshold of 1.96. This effectively removed 'ghost points' produced by partial beam 166 interceptions and also removed many erroneous returns caused by airborne dust particles, which 167 was common at these study sites. Raw points clouds for each subplot had a mean of ~2.9e+08 168 points, ~4.5e+07 points after voxelisation to 5 cm³, and ~2.1e+07 points after noise reduction. 169 Ground points were classified using the Progressive Morphological Filter (PMF) from Zhang 170 et al. (2003). Point cloud height was reclassified based on this revised ground layer by measuring 171 the vertical distance between the nearest ground point and each point. Points below 1.3 m 172 height above ground were discarded for calculations of foliage density, canopy cover, and canopy 173 complexity, as points below this threshold where often occupied by long grass. 174

2.3.2 Canopy complexity metrics

Ray-tracing was used to estimate canopy closure in each subplot, i.e. the proportion of the sky hemisphere occluded by plant material at the subplot centre from multiple TLS scans. Hemispherical images were created using the POV-Ray ray-tracing software (Persistence of Vision Pty. Ltd., 2004). Filled voxels were represented as black cubes filling the voxel volume, with a white sky box and no light source. A 'camera' with a 180° fisheye lens was placed at the

subplot centre within POV-Ray, at a height of 1.8 m pointing directly upwards. The images produced by POV-Ray were analysed using Hemiphot (ter Steege, 2018) to estimate canopy closure. Canopy closure estimates from the TLS were validated with hemispherical photographs taken at the same location and processed using the same method in Hemiphot, and compared using Pearson's correlation (r(195)=0.87, p<0.001). A plot level estimate of canopy closure was calculated as the mean of subplot canopy closure measurements. See Chapter 6 for expanded methods and explanation of the behaviour of the different canopy complexity metrics.

Effective Number of Layers (ENL) was calculated according to Ehbrecht et al. (2016) to measure vertical variation in subplot foliage density. ENL is calculated as the exponential Shannon index (i.e. the Hill number of order q=1) of foliage density among 50 cm vertical layers within each subplot:

$$ENL = \exp\left(-\sum_{i=1}^{N} p_i \times \ln p_i\right) \tag{1}$$

Where p_i is the proportion of filled voxels in the 50 cm layer i, and N is the total number of layers. ENL increases with canopy height and thus number of layers, and also with variation in foliage density among those layers, but not with increased total foliage density.

Total foliage density was calculated within each subplot as the area under the curve of the foliage height profile. Total foliage density was also calculated at the plot level as the sum of filled 50 cm³ voxels across the plot. Vertical variation in subplot foliage density was calculated by fitting a linear model to the cumulative foliage density profile, then calculating the sum of squared residuals of that model.

Plot level canopy surface models were extracted using the 99th percentile of canopy height in 10 cm² columns. A pit-filling algorithm provided by Khosravipour et al. (2014) was applied at 50 cm² resolution to reduce the effects of incomplete canopy penetration in dense canopies. Whole plot canopy complexity was measured by three metrics. Canopy top roughness was measured as the coefficient of variation (CV) of canopy height across the plot. Canopy rugosity was measured according to Hardiman et al. (2011), as the CV of vertical and horizontal foliage density within 0.5 m³ cubic bins. Finally, canopy height was calculated as the mean of the canopy surface roughness model across the plot.

2.3.3 Stand structure and diversity

195

196

197

198

199

200

201

204

205

206

207

For each subplot, an adapted version of the Iterative Hegyi index was used to estimate crowding, 209 as an alternative to stem density which does not adequately capture crowding at small spatial 210 scales when only a small number of trees are included in the sample (Hegyi, 1974). The CV of 211 stem diameter was calculated as a measure of the heterogeneity of tree size in the neighbourhood. 212 At the plot level, the regularity of species spatial distribution was estimated using the spatial 213 mingling index (von Gadow & Hui, 2002), which scores each tree based on whether it shares 214 species identity with its nearest neighbours. The spatial regularity of trees was estimated using 215 the uniform angle index (winkelmass) (von Gadow & Hui, 2002), which scores each tree based on 216 the angles between nearest neighbours. Additionally, the degree of spatial clustering of trees was measured using Voronoi tessellation, as the CV of Voronoi cell areas (Ong et al., 2012). Finally, 218 plot level tree density was calculated to estimate crowding at the plot scale. See Chapter 6 for 219 more information on the behaviour of the spatial mingling index and uniform angle index. 220

Species diversity at both the subplot and plot level was measured using the exponential Shannon index (i.e. the Hill number of order q = 1), calculated using tree species abundance (Jost, 2006). At the subplot level trees were included if they had canopy material inside the 10 m

diameter subplot, while at the plot level trees were included if the largest stem was inside the plot boundaries.

226 2.3.4 Statistical analysis

Non-metric Multi-dimensional Scaling (NMDS) was used to describe variation in species composition among plots, using genus-level basal area weighted abundance in each plot. Stems that could not be identified to genus were excluded from this analysis, which accounted for 0.2% of the total basal area recorded. Four distinct vegetation types, two from each site (Table 1), were identified using hirerarchical clustering of the four dominant NMDS ordination axes. Clusters were further described using Dufrêne-Legendre indicator species analysis and by ranking tree species according to abundance across all plots within each cluster.

Linear mixed effects models tested the effects of tree species diversity and stand structural 234 diversity on subplot canopy complexity metrics. Mixed models used a nested random intercept 235 structure to account for the sampling design of subplots within plots and plots within vegetation 236 types. Separate models were fitted for each canopy complexity metric, resulting in four models 237 at the subplot level. Effect sizes among fixed effects in maximal models were compared for each 238 canopy complexity metric, using the 95% confidence interval of the effect size to ascertain whether 239 a fixed effect was significant by whether the confidence interval overlapped zero (Nakagawa & 240 Cuthill, 2007). AIC values and Akaike weights of models with different combinations of fixed 241 effects were compared to determine which combination of diversity and structural metrics best 242 explained variation in each canopy complexity metric. 243

Path analysis was used to test whether tree species diversity influences canopy complexity 244 indirectly through its effect on stand structure, using the piecewiseSEM R package (Lefcheck, 245 2016). Two path analyses were conducted, one at the plot level and one at the subplot level. 246 Subplot path analysis investigated the direct effect of species diversity on canopy closure, as well as the indirect effect of diversity on canopy closure via the CV of basal area, with random 248 intercept terms for each vegetation type. The ex-Acacia vegetation type was represented by only 249 two plots and could not be included in this model due to lack of replication. Plot level path 250 analysis investigated the direct effects of species diversity and spatial mingling of species on 251 mean canopy height, as well as the indirect effects of these metrics on canopy height via tree density and basal area CV. Again, ex-Acacia plots were excluded from this path analysis. 253

Statistical analysis of the determinants of plot level canopy complexity metrics were conducted using linear models. Again, these models excluded the ex-Acacia vegetation type due to lack of replication. As with the subplot linear mixed models, predictor variable effect sizes were used to assess predictor variable significance, and comparison of candidate models using AIC, Akaike weights, and model R² values was used to determine which combination of predictors best explained each canopy complexity metric.

260 3 Results

261

3.1 Description of vegetation types

Indicator species analysis shows that the four identified vegetation types constitute common southern African savanna floristic archetypes (Table 2). Cluster 1, found in Bicuar National Park contains typical miombo species from the Detarioideae subfamily, such as *Julbernardia paniculata*. Cluster 1 is the most frequent vegetation type in this study, with 12 plots. Cluster 1 has the highest stem density, but lower AGB than Clusters 2 or 3, which contain larger individuals with disproportionately higher biomass. Cluster 2, found in Mtarure Forest Reserve, is dominated

Table 1: Description of the vegetation type clusters, identified using the Ward algorithm based on basal area weighted genus abundance. AGB = Above-Ground woody Biomass. Species richness, stem density and AGB are reported as the median among plots, with the interquartile range in parentheses.

Site	Cluster	N sites	Richness	Stem density (Stems ha ⁻¹)	AGB (t ha ⁻¹)
Bicuar	1	12	17(2)	642(194)	41(8.4)
Mtarure	2	5	23(4)	411(137)	72(11.9)
Bicuar	3	3	6(1)	196(55)	77(7.3)
Mtarure	4	2	12(2)	288(73)	9(0.2)

Table 2: Floristic description of the vegetation type clusters. Dominant species are the most abundant individuals across all plots per cluster. Indicator species are derived from Dufrêne-Legendre indicator species analysis with the three highest indicator values.

Cluster	Dominant species	Indicator species	Indicator value
1	Julbernardia paniculata	Strychnos spinosa	0.83
	Burkea africana	Combretum collinum	0.74
	Combretum collinum	Julbernardia paniculata	0.70
2	Diplorhynchus condylocarpon	Pteleopsis myrtifolia	1.00
	Pseudolachnostylis maprouneifolia	Diplorhynchus condylocarpon	0.89
	Gymnosporia senegalensis	Pseudolachnostylis maprouneifolia	0.81
3	Baikiaea plurijuga	Baikiaea plurijuga	0.94
	Baphia massaiensis	Baphia massaiensis	0.83
	Philenoptera nelsii	Philenoptera nelsii	0.45
4	Combretum apiculatum	Vachellia nilotica	0.99
	Burkea africana	Combretum apiculatum	0.70
	Bauhinia petersiana	Senegalia polyacantha	0.62

by Pteleopsis myrtifolia, a common miombo species from the Combretaceae family. Indeed, 268 Cluster 2 also contained other common miombo species shared with plots in Cluster 1, such as 269 Julbernardia globiflora and Pseudolachnostylis maprouneifolia, but these clusters remain distinct 270 due to biogeographic variation in endemic genera at the longitudinal extremes of the miombo 271 ecoregion represented by the two sites in this study. Cluster 3 represents Baikiaea woodland, 272 found on Kalahari sands in southern Angola. It is species poor and dominated by Baikiaea 273 plurijuga which forms large spreading canopy trees with high AGB. Other shrubby species that 274 coppice readily in response to disturbance by fire such as Baphia massaiensis are also common. 275 Cluster 4, found in Mtarure is a type of ex-Acacia woodland, dominated by Vachellia and 276 Senegalia spp. This vegetation type was not well represented in the study, with only two plots, 277 precluding its use in some multi-level statistical analyses due to lack of replication. Cluster 4 278 had far lower AGB than the other clusters (Table 1).

Differences in canopy structure among the four vegetation types are evident through observation 280 of canopy surface models for typical plots within each vegetation type (Figure 5), and by 281 comparing canopy complexity metrics (Figure 6). Cluster 1 shows many overlapping crowns 282 forming a nearly contiguous canopy surface, and the heighest plot foliage density of all clusters. 283 Though most trees in Cluster 1 have smaller crowns than those in Cluster 2, which also forms a 284 nearly contiguous canopy. The largest trees in Cluster 2 grow taller and have a wider spreading 285 canopy than those in other vegetation types. Cluster 3 shows two distinct size classes of tree, 286 the large Baikiaea plurijuqa forming clear isolated canopies, and much smaller scattered shrubby 287 individuals in the understorey. Cluster 4 shows many small shrubby individuals with irregular 288 canopy shapes, but a greater total crown area coverage than Cluster 3. 280

3.2 Bivariate relationships

290

291

292

293

294

295

296

297

298

309

Bivariate plots and linear models show that subplot species diversity, measured as the truenumbers equivalent of the Shannon diversity index of the tree neighbourhood around each 10 m diameter subplot, appears to have weak positive effects on subplot canopy layer diversity, canopy closure and foliage density (Figure 3, Table 3). The Hegyi crowding index had strong positive effects on all canopy complexity metrics, as expected. The effect of Hegyi crowding on subplot canopy complexity metrics was similar across all vegetation types (Table 6). Structural diversity, measured as the CV of subplot stem basal area had significant weak positive effects on total canopy foliage, layer diversity, and canopy closure.

At the plot level, effects of species diversity and stand structure on canopy complexity were 299 similarly weak, but not significant except for the effect on canopy height, which explained more 300 variance in canopy height than tree density (Figure 4, Table 3). The effect of spatial regularity of 301 stems on canopy closure, measured by uniform angle index, was clearly negative, while the effect 302 of spatial clustering of stems, measured by Voronoi cell area CV, was negligible. Additionally, 303 there was a non-significant negative effect of basal area CV on whole canopy rugosity. As 304 expected, tree density had strong positive and significant effects on foliage density and canopy 305 closure, and negative effects on canopy roughness and canopy rugosity. Cluster 4 represented 306 an outlier in plot level bivariate relationships, with low canopy closure, low canopy height, low 307 species diversity, and low variation in stem size. 308

3.3 Subplot mixed models

Linear mixed effects models showed that species diversity of the subplot neighbourhood contributed to both layer diversity and canopy closure (Table 4), despite their low R² in bivariate linear models, and low effect sizes in maximal linear mixed models (Figure 7). As also seen in the subplot bivariate relationships Figure 3, the Hegyi crowding index had strong positive

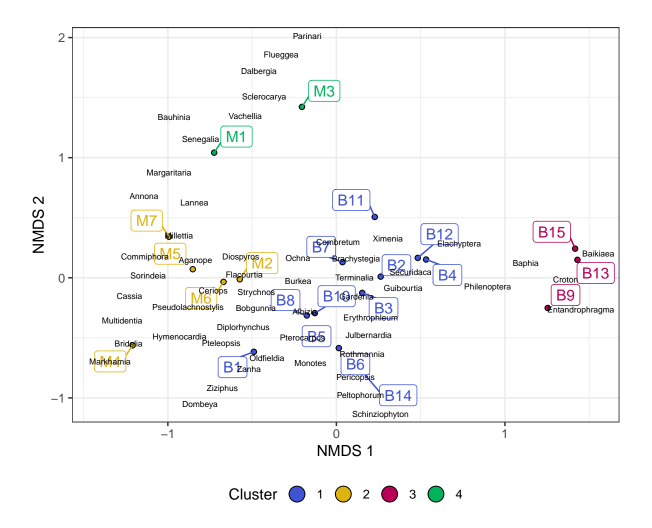


Figure 2: The first two axes of a Non-metric Multi-Dimensional Scaling (NMDS) analysis of tree genus diversity in each plot. Genus scores are labelled as black text, while plot scores are labelled as coloured points. Plots are shaded by vegetation types identified by hierarchical clustering: 1) B1-B8, B10-B12, B14, dominated by core miombo species such as *Julbernardia* spp., *Brachystegia* spp.; 2) M2, M5, M6, and M7, also dominated by core miombo genera with some genera not found in Bicuar National Park such as *Commiphora* and *Sorindeia*; 3) B9, B13 and B15, dominated by *Baikiaea plurijuga*; and 4) M1, M3, and M4, dominated by *Senegalia* spp., *Vachellia* spp., and *Combretum* spp.

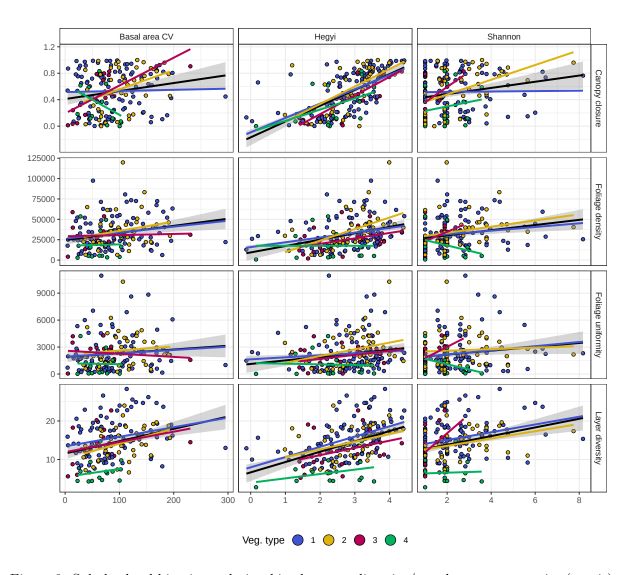


Figure 3: Subplot level bivariate relationships between diversity/stand structure metrics (x axis) and canopy complexity metrics (y axis). Points and linear model lines of best fit are coloured by vegetation type. Black lines of best fit are linear models including all plots, with a 95% confidence interval. See Table 6 for a comparison of linear model fits by vegetation type.

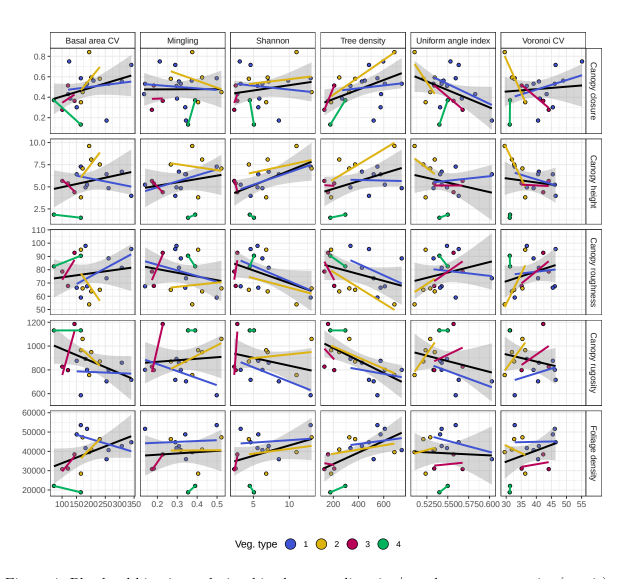


Figure 4: Plot level bivariate relationships between diversity/stand structure metrics (x axis) and canopy complexity metrics (y axis). Points and linear model lines of best fit are coloured by vegetation type. Black lines of best fit are linear models including all plots, with a 95% confidence interval. See Table 6 for a comparison of linear model fits by vegetation type.

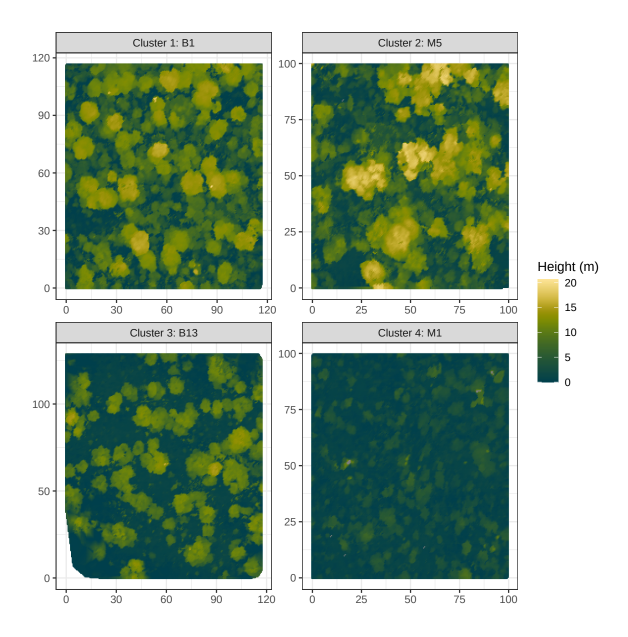


Figure 5: Representative canopy surface models for each vegetation type identified in the Non-metric Multi-dimensional Scaling (NMDS) clustering analysis. Panel titles show the plot name and the vegetation type.

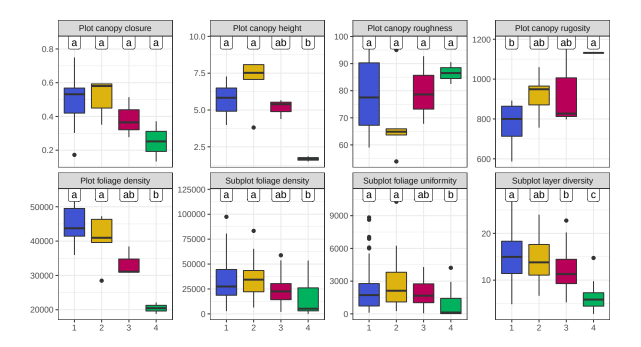


Figure 6: Box plots showing variation in canopy complexity metrics among the four vegetation types identified in the Non-metric Multi-dimensional Scaling (NMDS) clustering analysis. Thick lines show the median, boxes show the interquartile range (IQR), whiskers show $1.5 \times IQR$, and points show outliers beyond these limits. Labels above each box plot group vegetation types according to significant differences in pairwise Tukey's tests; vegetation types sharing a letter are not significantly different.

effects on all measured canopy complexity metrics, though these effects were non-significant for vegetation clusters 3 and 4. Heterogeneity of stem basal area had a significant positive effect on layer diversity and foliage density, but there was wide variation in vegetation type marginal effects for Clusters 3 and 4, due to low levels of replication. Cluster 3 had strong positive effects of species diversity on foliage uniformity and layer diversity. The random effects of vegetation type and plot identity described most of the variation in layer diversity and foliage density. Foliage uniformity was poorly explained by all combinations of fixed effects, with the best model only explaining 29%. All models were better than random effects only models according to AIC values.

3.4 Whole-plot multivariate linear models

While species diversity had varying effects on different plot level canopy complexity metrics, the confidence intervals on these effect sizes were wide (Figure 8). Species diversity had a significant positive effect on canopy height (β =3±0.96, p<0.05), a non-significant positive effect on canopy closure (β =0.07±0.085, p=0.41), but a negative effect on canopy surface roughness (β =-13±6.8, p=0.09) and whole canopy rugosity (β =-111±71, p=0.15). Spatial mingling of tree species had a positive effect on canopy surface roughness and canopy rugosity, but a strong negative effect on canopy height. Plot tree density had negligible effects on canopy complexity, except for canopy rugosity (β =-61±42, p=0.17), in contrast to the effect of Hegyi crowding on subplot canopy complexity. Spatially explicit measures of structural diversity, measured by the uniform angle index, Voronoi cell area CV, and basal area CV, had smaller effects on canopy complexity than species diversity, which were generally insignificant. One exception was the effect of uniform angle index, i.e. the spatial clustering of stems, on canopy closure, which was clearly negative,

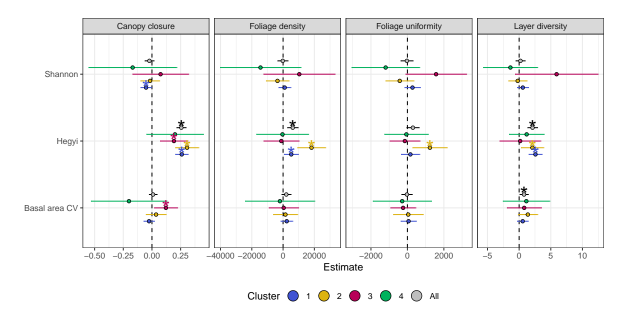


Figure 7: Standardised fixed effect slopes for each model of a canopy complexity metric. Slope estimates where the interval (± 1 standard error) does not overlap zero are considered to be significant effects, marked with asterisks. Points are coloured according to vegetation type.

though still insignificant (β =-0.08±0.043, p=0.1), the effect of Voronoi cell area CV on foliage density, which was positive (β =6199±3312, p=0.09), and the effect of basal area CV on canopy closure, which was positive (β =0.06±0.042, p=0.19).

Despite the weak effect sizes of species diverity on canopy complexity at the plot level, model selection showed that foliage density, canopy height and canopy roughness were better explained by models which included species diversity (Table 5). Additionally, the best models for canopy height and canopy roughness also included spatial mingling of tree species. The model for canopy roughness was only maginally better than a null model and the model did not have a significant p-value.

3.5 Path analysis

The subplot level path analysis investigating the indirect effect of subplot species diversity on canopy closure via the basal area CV showed that while species diversity had a strong positive significant effect on basal area variation, the effect of basal area variation on canopy closure remained negligible (Figure 9). The indirect effect of species diversity on canopy closure via basal area CV was -0.0016, while the direct effect was -0.078. The R² of this model was 0.47. As in the bivariate relationships and plot level linear models, species diversity had a weak positive significant effect on canopy closure, while the major driver of canopy closure was the Hegyi crowding index.

The plot level path analysis, which tested the effects of species diversity and species mingling on canopy height, showed that the main effect of species diversity on canopy height was direct (1.3*), while the indirect effects via basal area CV (0.0210), and tree density (-0.0294), remained small and insignificant. Shannon diversity had a strong positive effect on tree density. Species mingling had a moderately strong negative but insignificant direct effect on canopy height, as in the linear mixed models.

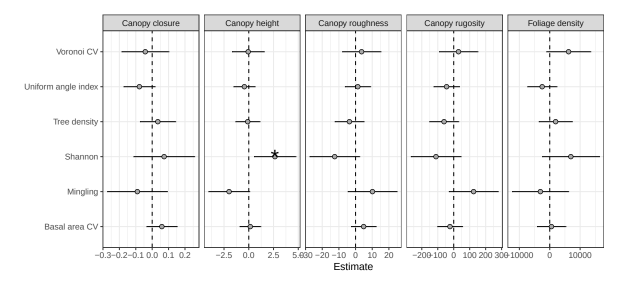


Figure 8: Standardised effect sizes for whole-plot canopy rugosity. Slope estimates where the interval (± 1 standard error) does not overlap zero are considered to be significant effects, marked with asterisks.

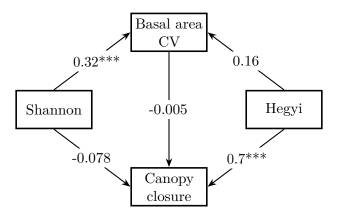


Figure 9: Directed Acyclic Graph showing standardised path coefficients of paths in the path analysis of the indirect effect of subplot species diversity (Shannon diversity index) on canopy closure via basal area CV. Asterisks define p-value thresholds: *<0.05, **<0.01, ***<0.001.

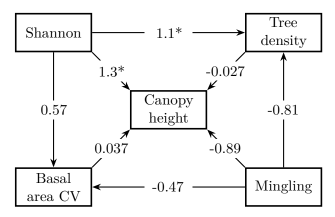


Figure 10: Directed Acyclic Graph showing standardised path coefficients of paths in the path analysis of the indirect effect of plot species diversity (Shannon diversity index) and species mingling on mean canopy height via stand structural metrics of basal area CV and tree density. Asterisks define p-value thresholds: *<0.05, **<0.01, ***<0.001.

3.6 Covariance of subplot and plot measures of canopy complexity

Plot and subplot canopy complexity metrics were highly correlated in many cases, with similar relationships among vegetation types (Figure 8). Most subplot and plot level canopy metrics covaried in a predictable manner. For example, increased canopy height led to an increase in canopy closure. Plot canopy height especially, tended to be strongly positively correlated with subplot canopy complexity metrics. Additionally, as canopy rugosity increased, many subplot canopy complexity and density metrics decreased. Subplot metrics varied greatly within plots, producing large uncertainty in plot level estimates of these metrics. All subplot level canopy complexity metrics positive correlated with each other (Figure 13). Plot level canopy complexity also generally correlated (Figure 12). Plot level measures of spatial heterogeneity in canopy structure, i.e. canopy surface roughness and canopy rugosity, were negatively correlated with measures of canopy density, i.e. foliage density, canopy closure, and canopy height. Measures of canopy spatial heterogeneity positively correlated with each other, as did measures of canopy density.

³⁷⁴ 4 Discussion

This study investigated relationships between tree species diversity, stand structure, and several metrics of tree canopy complexity using terrestrial LiDAR in southern African savannas, with a view to improving understanding of the biotic drivers of variation in canopy complexity and vegetation dynamics. Species diversity appeared to generally have weak positive effects on canopy complexity metrics related to canopy density at both the subplot and plot scales. Plots with greater species diversity produced taller tree canopies, with greater canopy closure and foliage density. Species diversity had negative effects on canopy surface roughness and canopy rugosity, canopy complexity metrics both related to the spatial heterogeneity of foliage distribution. The study did not however, find support for the hypothesis that increased heterogeneity in tree stem size causes an increase in canopy complexity, and only partial support for the hypothesis that greater heterogeneity in stem location causes increased canopy complexity.

4.1 Ecological consequences of a species diversity effect on canopy complexity

The result that species diversity increases metrics of canopy density suggests that diverse stands can more effectively close the tree canopy under a given set of environmental conditions. Increased canopy closure reduces light penetration to the ground (), reducing grassy fuel load, and so could promote woody densification in diverse stands under atmospheric CO₂ enrichment. Similarly, the finding that species diversity causes an increase in foliage density and canopy height suggests that more diverse stands could more effectively upregulate productivity in response to atmospheric CO₂ fertilisation, and maintain stands with greater woody biomass. Taller trees hold disproportionately higher biomass than shorter trees (King, 1990). In mesic savannas that are prone to disturbance by fire, increased growth rate and canopy height could increase the likelihood of trees escaping the "fire trap", and facilitate their growth to larger canopy trees (). This finding concurs with many previous studies, which have found that species diversity leads to greater woody productivity in both forests and savannas (). This study adds further information on the mechanisms underlying the species diversity effect on ecosystem function in savannas, that niche complementarity among species promotes greater canopy occupancy.

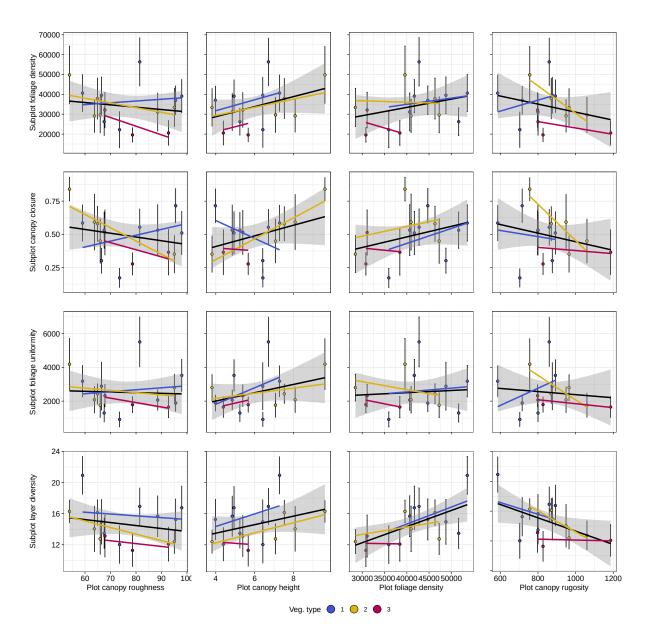


Figure 11: Bivariate plots comparing canopy structural metrics at the plot (x axis) and subplot scale (y axis). Each point represents the mean values of a single plot. Points and linear model fits are coloured according to vegetation type. The black linear model combines all vegetation types. Error bars on points are the standard deviation of mean subplot metrics across the plot. Note that because plot level canopy closure is calculated as the mean of subplot canopy closure, a comparison of subplot and plot canopy closure is not made in this figure.

4.2 Species diversity and variation in tree size

401

402

420

421

422

423

424

425

426

427

428

429

430

431

432

433

434

435

stem size were negligible, due to the lack of an effect of tree stem size on canopy complexity at 403 both the subplot and plot scale. This finding suggests that the effects of species diversity on 404 canopy complexity are not simply due to covariation with heterogeneity of tree stem size, which 405 could also arise due to demographic effects, rather they are due to differences among species 406 in their canopy dimensions and growth strategy. This strengthens support for the hypothesis 407 that species diversity genuinely increases canopy complexity. Previous studies in temperate and 408 boreal forests have suggested that increasing tree stem size diversity through active management 409 and thinning can offset productivity losses caused by reduced species diversity (Levick et al., 410 2009), but this study suggests that in southern African savannas at least, this is not the case. 411 Shannon diversity did however, have strong positive effects on tree stem size variation and tree 412 density. This can be interpreted as a niche complementarity effect, where diverse savannas are 413 able to support a greater density of trees by reducing the effects of intraspecific competition and 414 promoting canopy occupation in different vertical layers, by trees of different sizes. Although basal area CV was included in best models for subplot layer diversity and foliage density, subplot 416 level path analysis suggests that covariance between basal area CV and species diversity, and 417 the strong direct effect of species diversity on canopy complexity, means that any observed effect 418 of basal area CV on subplot canopy complexity is actually due to species diversity rather than 419

Path analysis showed that indirect effects of species diversity on canopy complexity via tree

Despite tree density having strong positive observed effects on canopy density, and negative effects on canopy heterogeneity in bivariate relationships, tree density only appeared in one of the best quality plot level mixed models, for canopy rugosity. Additionally, the effect sizes for tree density in maximal mixed models were small. At the subplot level however, the Hegyi crowding index, which can be seen as analagous to tree density at neighbourhood scales, was a strong determinant of canopy complexity. Tree density was shown to covary with basal area CV and Voronoi cell area CV at the plot scale, which described variation in tree size and degree of spatial clustering, respectively. This covariance may have led to the effect dropping out in the best models at the plot scale, while at the subplot scale there were no measures of spatial clustering included. Hegyi crowding and tree density are expected to positively correlate with resource availability, and negatively correlate with disturbance which causes tree mortality and reduces tree growth. The study sites sampled here did cover a gradient of tree density, but variation in tree density within vegetation types was small. As vegetation type was included as a random effect in mixed models, this may have led to tree density not having a larger effect size in the models.

Variation in tree size caused positive canopy complexity effects for within-canopy structural
metrics such as layer diversity and canopy surface roughness, but had negligible effects on canopy
density. This is in line with other studies in forest ecosystems, which report that variation in
tree size increases total canopy volume occupancy by increasing the number of canopy layers,
but does not necessarily result in a concomitant increase in canopy closure, as the resulting
canopies are often more sparse, due to increased competition for light (**Beland**). Both path
analyses also support this conclusion, where species diversity was found to cause an increase in
stand structural diversity, but this did not extend to an increase in canopy closure.

44 4.3 Spatial clustering of stems

basal area CV itself.

Spatial regularity of stems, measured using the uniform angle index, caused a clear decrease in canopy closure, with similar behaviour across vegetation types. Uniform angle index was also included in the best multivariate model predicting canopy closure. Concurrently, spatial

clustering of stems, measured by Voronoi cell area CV was included in the best model for 448 foliage density but had a positive effect on this canopy complexity metric. This finding is 449 expected, as spatial clustering results in reduced canopy cover in areas outside clusters, and 450 a non-compensatory increase in canopy closure within clusters, due to competition among 451 individuals (). In contrast, changes in spatial regularity of trees do not imply changes in the 452 distance of stems, only their layout, but does spatial regularity does covary with spatial clustering. 453 Clustering of trees in savannas can result from positive feedback effects from disturbance by fire 454 and herbivory (), which result in a patchwork of open areas alongside areas with a dense canopy 455 (). 456

Covariation of canopy complexity metrics 4.4 457

All subplot canopy complexity metrics positively covaried. For rapid assessments of canopy 458 complexity, simple canopy closure measurements are sufficient to characterise canopy complexity, 459 at the spatial scale of an individual tree neighbourhood. In closed-canopy forest ecosystems, 460 measures of canopy spatial heterogeneity often correlate with canopy closure (). In the savannas 461 studied here however, the sparser and discontinuous canopy results in a negative relationship. 462

Plot vs. subplot scale results 463

464

465

467

468

469

471

472

473

474

475

476

477

478

479

481

482

483

485

486

487

488

489

491

The standardised effect sizes of species diversity on canopy complexity metrics were generally greater at the plot level than at the subplot level. While positive and significant relationships between species diversity and subplot canopy complexity metrics were observed in the subplot bivariate models, subplot linear mixed effects models did not show strong species diversity effects, and were instead dominated by the effect of crowding. This finding suggests a large degree of stochastic variability in canopy complexity within plots, that can mask species effects at smaller spatial scales. The prevalence of disturbance events such as fire and damage by 470 elephants in southern African woodlands, as well as tree-fall, small-scale variability in edaphic factors, and stochastic tree germination all contribute to heterogeneity in canopy complexity (). While disturbances are controlled to some extent by stand structure and composition, due to the stochastic nature of disturbance events, a snapshot study such as this cannot capture the average disturbance regime and there is therefore a great deal of noise in models predicting canopy complexity from species diversity. The contrast in strength of species diversity effects at the subplot and plot level demonstrates the importance of large sample units, a high degree of spatial replication, and ideally a longer time scale when measuring canopy complexity, especially in disturbed systems, to effectively account for the inherent heterogeneity in the system ().

Variation among vegetation types 4.6 480

Bivariate relationships showed that some of the observed species diversity effect on canopy closure and foliage density may be driven by vegetation type, and the variation in species diversity among vegetation types. The linear mixed model framework however, which accounted for differences among vegetation types, still shows weak species diversity effects even after vegetation type is controlled for, strengthening the validity of the result. Canopy complexity metrics also differed among vegetation types, but significant differences among vegetation types only occurred in a few cases. Canopy density and total foliage volume was lowest in ex-Acacia plots, as expected, while the highest canopy density occurred in miombo plots in clusters 1 and 2. Miombo woodlands frequently have contiguous canopies with overlapping individual tree canopies (), while ex-Acacia savannas show greater negative density dependence of individuals, forming patchy canopies with simpler vertical profiles (). From this result it is suggested that

under identical disturbance regimes, ex-Acacia savannas may not be as effective at closing their canopy to exclude grasses. Particularly, the lower maximum tree height of trees in the ex-Acacia plots may preclude these savannas from forming a multi-layer canopy that may be necessary to allow increased biomass areal density ().

While vegetation types differed in mean values for stand structural and species diversity metrics, 496 variation in these metrics produced results of similar direction and magnitude among vegetation 497 types in most cases, suggesting that ecosystem processes driving canopy complexity are similar, 498 even across these varied savanna types. Small sample sizes for Baikiaea and ex-Acacia vegetation 499 however, led to wide errors on most relationships especially at the plot level, such that it is impossible to draw deep conclusions about the behaviour of these vegetation types. Variation in 501 mean values of canopy complexity metrics among vegetation types is likely driven by species 502 identity and variation in physiological limits on tree physiognomy (), though species composition 503 itself is driven by environmental factors and disturbance regime (). 504

$_{505}$ 5 Conclusion

6 References

- Archibald, S. & W. J. Bond (2003). 'Growing tall vs growing wide: tree architecture and allometry of Acacia karroo in forest, savanna, and arid environments'. In: *Oikos* 102.1, pp. 3–14. DOI: 10.1034/j.1600-0706.2003.12181.x.
- Axelsson, C. R. & N. P. Hanan (2018). 'Rates of woody encroachment in African savannas reflect water constraints and fire disturbance'. In: *Journal of Biogeography* 45.6, pp. 1209–1218. DOI: 10.1111/jbi.13221.
- Baldocchi, D. D. & K. B. Wilson (2001). 'Modeling CO2 and water vapor exchange of a temperate
 broadleaved forest across hourly to decadal time scales'. In: *Ecological Modelling* 142.1-2,
 pp. 155–184. DOI: 10.1016/s0304-3800(01)00287-3.
- Barry, K. E., L. Mommer, J. van Ruijven, C. Wirth, A. J. Wright, Y. Bai, J. Connolly, G. B. D. Deyn, H. de Kroon, F. Isbell et al. (2019). 'The Future of Complementarity: Disentangling Causes from Consequences'. In: *Trends in Ecology & Evolution* 34.2, pp. 167–180. DOI: 10.1016/j.tree.2018.10.013.
- Bond, W. J. & G. F. Midgley (2012). 'Carbon dioxide and the uneasy interactions of trees and savannah grasses'. In: *Philosophical Transactions of the Royal Society B: Biological Sciences* 367.1588, pp. 601–612. DOI: 10.1098/rstb.2011.0182.
- Buitenwerf, R., W. J. Bond, N. Stevens & W. S. W. Trollope (2012). 'Increased tree densities in South African savannas: >50 years of data suggests CO₂ as a driver'. In: Global Change Biology 18.2, pp. 675–684. DOI: 10.1111/j.1365-2486.2011.02561.x.
- Calders, K., J. Adams, J. Armston, H. Bartholomeus, S. Bauwens, L. P. Bentley, J. Chave, F. M. Danson, M. Demol, M. Disney et al. (2020). 'Terrestrial laser scanning in forest ecology: Expanding the horizon'. In: *Remote Sensing of Environment* 251, p. 112102. DOI: 10.1016/j.rse.2020.112102.
- Charles-Dominique, T., G. F. Midgley, K. W. Tomlinson & W. J. Bond (2018). 'Steal the
 light: shade vs fire adapted vegetation in forest-savanna mosaics'. In: New Phytologist 218.4,
 pp. 1419–1429. DOI: 10.1111/nph.15117.
- Chen, J. M., G. Mo, J. Pisek, J. Liu, F. Deng, M. Ishizawa & D. Chan (2012). 'Effects of foliage clumping on the estimation of global terrestrial gross primary productivity'. In: Global Biogeochemical Cycles 26, GB1019. DOI: 10.1029/2010gb003996.
- Cifuentes, R., D. V. der Zande, J. Farifteh, C. Salas & P. Coppin (2014). 'Effects of voxel size
 and sampling setup on the estimation of forest canopy gap fraction from terrestrial laser

- scanning data'. In: Agricultural and Forest Meteorology 194, pp. 230-240. DOI: 10.1016/j. agrformet.2014.04.013.
- Criado, M. G., I. H. Myers-Smith, A. D. Bjorkman, C. E. R. Lehmann & N. Stevens (2020).
 'Woody plant encroachment intensifies under climate change across tundra and savanna biomes'.
 In: Global Ecology and Biogeography 29.5, pp. 925–943. DOI: 10.1111/geb.13072.
- Dohn, J., D. J. Augustine, N. P. Hanan, J. Ratnam & M. Sankaran (2017). 'Spatial vegetation patterns and neighborhood competition among woody plants in an East African savanna'. In:

 Ecology 98.2, pp. 478–488. DOI: 10.1002/ecy.1659.
- Donohue, R. J., M. L. Roderick, T. R. McVicar & G. D. Farquhar (2013). 'Impact of CO₂
 fertilization on maximum foliage cover across the globe's warm, arid environments'. In:
 Geophysical Research Letters 40.12, pp. 3031–3035. DOI: 10.1002/grl.50563.
- Ehbrecht, M., P. Schall, J. Juchheim, C. Ammer & D. Seidel (2016). 'Effective number of layers:

 A new measure for quantifying three-dimensional stand structure based on sampling with
 terrestrial LiDAR'. In: Forest Ecology and Management 380, pp. 212–223. DOI: 10.1016/j.
 foreco.2016.09.003.
- Fotis, A. T., T. H. Morin, R. T. Fahey, B. S. Hardiman, G. Bohrer & P. S. Curtis (2018).

 'Forest structure in space and time: Biotic and abiotic determinants of canopy complexity and
 their effects on net primary productivity'. In: Agricultural and Forest Meteorology 250-251,
 pp. 181–191. DOI: 10.1016/j.agrformet.2017.12.251.
- Frost, P. (1996). 'The ecology of miombo woodlands'. In: *The miombo in transition: woodlands*and welfare in Africa. Ed. by B. Campbell. Bogor, Indonesia: Center for International Forestry
 Research, pp. 11–55.
- Gough, C. M., J. W. Atkins, R. T. Fahey & B. S. Hardiman (2019). 'High rates of primary production in structurally complex forests'. In: *Ecology* 100.10. DOI: 10.1002/ecy.2864.
- Hardiman, B. S., G. Bohrer, C. M. Gough, C. S. Vogel & P. S. Curtis (2011). 'The role of canopy
 structural complexity in wood net primary production of a maturing northern deciduous
 forest'. In: *Ecology* 92.9, pp. 1818–1827. DOI: 10.1890/10-2192.1.
- Hegyi, F. (1974). 'A simulation model for managing jack-pine stands'. In: Royal College of
 Forestry, editor. Stockholm, Sweden: Royal College of Forestry, pp. 74–90.
- Hirota, M., M. Holmgren, E. H. Van Nes & M. Scheffer (2011). 'Global resilience of tropical forest and savanna to critical transitions'. In: *Science* 334, pp. 232–235. DOI: 10.1126/science. 1210657.
- Jonckheere, I., S. Fleck, K. Nackaerts, B. Muys, P. Coppin, M. Weiss & F. Baret (2004). 'Review of methods for in situ leaf area index determination'. In: *Agricultural and Forest Meteorology* 121.1-2, pp. 19–35. DOI: 10.1016/j.agrformet.2003.08.027.
- Jost, L. (2006). 'Entropy and diversity'. In: *Oikos* 113.2, pp. 363–375. doi: 10.1111/j.2006. 0030-1299.14714.x.
- Jucker, T., O. Bouriaud & D. A. Coomes (2015). 'Crown plasticity enables trees to optimize canopy packing in mixed-species forests'. In: *Functional Ecology* 29.8, pp. 1078–1086. DOI: 10.1111/1365-2435.12428.
- Kershaw, J. A., M. J. Ducey, T. W. Beers & B. Husch (2017). Forest Mensuration. Chichester, UK: John Wiley & Sons. ISBN: 9781118902035.
- Khosravipour, A., A. K. Skidmore, M. Isenburg, T. Wang & Y. A. Hussin (2014). 'Generating
 Pit-free Canopy Height Models from Airborne LiDAR'. In: *Photogrammetric Engineering & Remote Sensing* 80.9, pp. 863–872. DOI: 10.14358/pers.80.9.863.
- King, D. A. (1990). 'The Adaptive Significance of Tree Height'. In: *The American Naturalist* 135.6, pp. 809–828. Doi: 10.1086/285075.
- Körner, C. (2017). 'A matter of tree longevity'. In: *Science* 355.6321, pp. 130–131. DOI: 10.1126/ science.aal2449.

- Law, B. E., A. Cescatti & D. D. Baldocchi (2001). 'Leaf area distribution and radiative transfer in 587 open-canopy forests: implications for mass and energy exchange'. In: Tree Physiology 21.12-13, 588 pp. 777-787. DOI: 10.1093/treephys/21.12-13.777. 589
- Lefcheck, J. S. (2016). 'piecewiseSEM: Piecewise structural equation modeling in R for ecology, 590 evolution, and systematics'. In: Methods in Ecology and Evolution 7.5, pp. 573–579. DOI: 591 10.1111/2041-210X.12512. 592
- Levick, S. R., G. P. Asner, T. Kennedy-Bowdoin & D. E. Knapp (2009). 'The relative influ-593 ence of fire and herbivory on savanna three-dimensional vegetation structure'. In: Biological 594 Conservation 142.8, pp. 1693-1700. DOI: 10.1016/j.biocon.2009.03.004. 595
- Lowman, M. D. & H. B. Rinker (2004). Forest Canopies. Physiological Ecology. Burlington MA, 596 USA: Elsevier Science. ISBN: 9780080491349. 597
- Mitchard, E. T. A. & C. M. Flintrop (2013). 'Woody encroachment and forest degradation in 598 sub-Saharan Africa's woodlands and savannas 1982-2006'. In: Philosophical Transactions of the 599 Royal Society B: Biological Sciences 368.1625, p. 20120406. DOI: 10.1098/rstb.2012.0406. 600
- Morin, X. (2015). 'Species richness promotes canopy packing: a promising step towards a better 601 understanding of the mechanisms driving the diversity effects on forest functioning'. In: 602 Functional Ecology 29.8, pp. 993-994. DOI: 10.1111/1365-2435.12473. 603
- Mugasha, W. A., O. M. Bollandsås & T. Eid (2013). 'Relationships between diameter and height 604 of trees in natural tropical forest in Tanzania'. In: Southern Forests: a Journal of Forest 605 Science 75.4, pp. 221–237. DOI: 10.2989/20702620.2013.824672. 606
- Muir, J., S. Phinn, T. Eyre & P. Scarth (2018). 'Measuring plot scale woodland structure using 607 terrestrial laser scanning'. In: Remote Sensing in Ecology and Conservation 4.4, pp. 320–338. 608 DOI: 10.1002/rse2.82. 609
- Muumbe, T. P., J. Baade, J. Singh, C. Schmullius & C. Thau (2021). 'Terrestrial Laser Scanning 610 for Vegetation Analyses with a Special Focus on Savannas'. In: Remote Sensing 13.3, p. 507. 611 DOI: 10.3390/rs13030507. 612
- Nakagawa, S. & I. C. Cuthill (2007). 'Effect size, confidence interval and statistical significance: a 613 practical guide for biologists'. In: Biological Reviews 82.4, pp. 591–605. DOI: 10.1111/j.1469-614 185x.2007.00027.x. 615
- Ong, M. S., Y. C. Kuang & M. P.-L. Ooi (2012). 'Statistical measures of two dimensional point set uniformity'. In: Computational Statistics & Data Analysis 56.6, pp. 2159–2181. DOI: 617 10.1016/j.csda.2011.12.005. 618
- Panzou, G. J. L., A. Fayolle, T. Jucker, O. L. Phillips, S. Bohlman, L. F. Banin, S. L. Lewis, 619 K. Affum-Baffoe, L. F. Alves, C. Antin et al. (2020). 'Pantropical variability in tree crown 620 allometry'. In: Global Ecology and Biogeography 30.2, pp. 459-475. DOI: 10.1111/geb.13231. 621
- Persistence of Vision Pty. Ltd. (2004). Persistence of Vision Raytracer (Version 3.7). [Computer 622 software. 623
- Pretzsch, H. (2014). 'Canopy space filling and tree crown morphology in mixed-species stands 624 compared with monocultures'. In: Forest Ecology and Management 327, pp. 251–264. DOI: 625 http://dx.doi.org/10.1016/j.foreco.2014.04.027. 626
- Privette, J., Y. Tian, G. Roberts, R. Scholes, Y. Wang, K. Caylor, P. Frost & M. Mukelabai (2004). 627 'Vegetation structure characteristics and relationships of Kalahari woodlands and savannas'. 628 In: Global Change Biology 10.3, pp. 281–291. DOI: 10.1111/j.1365-2486.2004.00740.x. 629
- Ratcliffe, S., C. Wirth, T. Jucker, F. van der Plas, M. Scherer-Lorenzen, K. Verheyen, E. Allan, 630 R. Benavides, H. Bruelheide, B. Ohse et al. (2017). 'Biodiversity and ecosystem functioning 631 relations in European forests depend on environmental context'. In: Ecology Letters 20, 632 pp. 1414-1426. DOI: http://dx.doi.org/10.1111/ele.12849. 633
- Reich, P. B., S. E. Hobbie & T. D. Lee (2014). 'Plant growth enhancement by elevated CO2 634 eliminated by joint water and nitrogen limitation'. In: Nature Geoscience 7.12, pp. 920–924. 635 DOI: 10.1038/ngeo2284. 636

- Ribeiro, N. S., P. L. Silva de Miranda & J. Timberlake (2020). 'Biogeography and Ecology 637 of Miombo Woodlands'. In: Miombo Woodlands in a Changing Environment: Securing the 638 Resilience and Sustainability of People and Woodlands. Ed. by N. S. Ribeiro, Y. Katerere, 639 P. W. Chirwa & I. M. Grundy. Springer International Publishing, pp. 9–53. DOI: 10.1007/978-640 3-030-50104-4_2.
- Rusu, R. B., Z. C. Marton, N. Blodow, M. Dolha & M. Beetz (2008). 'Towards 3D Point cloud based object maps for household environments'. In: Robotics and Autonomous Systems 56.11, 643 pp. 927-941. DOI: 10.1016/j.robot.2008.08.005. 644

641

- Sankaran, M., N. P. Hanan, R. J. Scholes, J. Ratnam, D. J. Augustine, B. S. Cade, J. Gignoux, 645 S. I. Higgins, X. Le Roux, F. Ludwig et al. (2005). 'Determinants of woody cover in African 646 savannas'. In: Nature 438.8, pp. 846-849. DOI: http://dx.doi.org/10.1038/nature04070. 647
- Scheuermann, C. M., L. E. Nave, R. T. Fahey, K. J. Nadelhoffer & C. M. Gough (2018). 'Effects of canopy structure and species diversity on primary production in upper Great Lakes forests'. 649 In: Oecologia 188.2, pp. 405–415. DOI: 10.1007/s00442-018-4236-x. 650
- Scholes, R. J. & S. R. Archer (1997). 'Tree grass interactions in savannas'. In: Annual Review of 651 Ecology and Systematics. 652
- Seidel, D., S. Fleck & C. Leuschner (2012). 'Analyzing forest canopies with ground-based 653 laser scanning: A comparison with hemispherical photography'. In: Agricultural and Forest 654 Meteorology 154-155, pp. 1-8. DOI: 10.1016/j.agrformet.2011.10.006. 655
- Seidel, D., S. Fleck, C. Leuschner & T. Hammett (2011). 'Review of ground-based methods to 656 measure the distribution of biomass in forest canopies'. In: Annals of Forest Science 68.2, 657 pp. 225–244. DOI: 10.1007/s13595-011-0040-z. 658
- Shenkin, A., L. P. Bentley, I. Oliveras, N. Salinas, S. Adu-Bredu, B. H. Marimon-Junior, B. S. 659 Marimon, T. Peprah, E. L. Choque, L. T. Rodriguez et al. (2020). 'The Influence of Ecosystem and Phylogeny on Tropical Tree Crown Size and Shape'. In: Frontiers in Forests and Global 661 Change 3. DOI: 10.3389/ffgc.2020.501757. 662
- Sitch, S., P. Friedlingstein, N. Gruber, S. D. Jones, G. Murray-Tortarolo, A. Ahlström, S. C. 663 Doney, H. Graven, C. Heinze, C. Huntingford et al. (2015). 'Recent trends and drivers of 664 regional sources and sinks of carbon dioxide'. In: Biogeosciences 12.3, pp. 653–679. DOI: 10.5194/bg-12-653-2015.
- Solbrig, O. T., E. Medina & J. F. Silva (1996). Biodiversity and Savanna Ecosystem Processes. 667 Berlin, Germany: Springer-Verlag. 668
- Stark, S. C., B. J. Enquist, S. R. Saleska, V. Leitold, J. Schietti, M. Longo, L. F. Alves, P. B. 669 Camargo & R. C. Oliveira (2015). 'Linking canopy leaf area and light environments with tree 670 size distributions to explain Amazon forest demography'. In: Ecology Letters 18.7, pp. 636–645. 671 DOI: 10.1111/ele.12440. 672
- Staver, A. C. & S. E. Koerner (2015). 'Top-down and bottom-up interactions determine tree 673 and herbaceous layer dynamics in savanna grasslands'. In: Trophic Ecology: Bottom-up and 674 Top-Down Interactions Across Aquatic and Terrestrial Systems. Ed. by K. J. La Pierre & 675 T. C. Hanley. Cambridge, United Kingdom: Cambridge University Press, pp. 86–106. 676
- Stevens, N., C. E. R. Lehmann, B. P. Murphy & G. Durigan (2016). 'Savanna woody encroachment is widespread across three continents'. In: Global Change Biology 23.1, pp. 235–244. DOI: 678 10.1111/gcb.13409. 679
- ter Steege, H. (2018). Hemiphot.R: Free R scripts to analyse hemispherical photographs for 680 canopy openness, leaf area index and photosynthetic active radiation under forest canopies. 681
- Unpublished report. Leiden, The Netherlands: Naturalis Biodiversity Center. URL: https: 682 //github.com/Naturalis/Hemiphot. 683
- von Gadow, K. & G. Hui (2002). Characterising forest spatial structure and diversity. Ed. by 684 L. Bjoerk. Lund, Sweden, pp. 20–30. 685

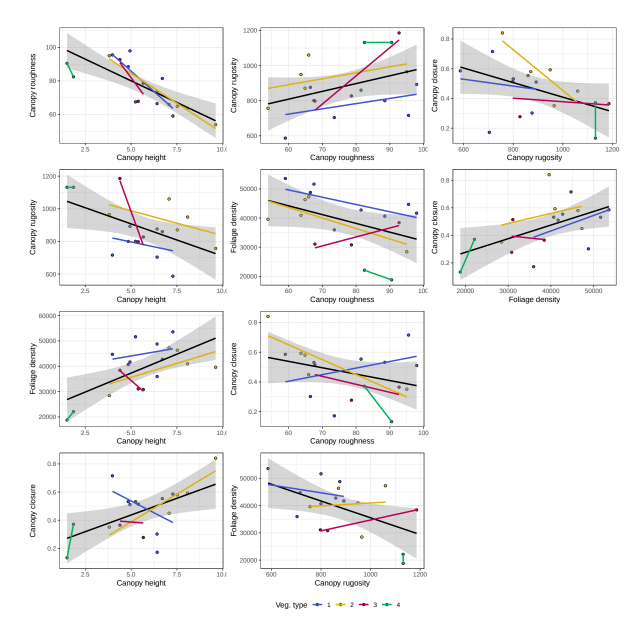


Figure 12: Bivariate scatter plots of plot level canopy complexity metrics.

White, F. (1983). The Vegetation of Africa: A descriptive memoir to accompany the UN ESCO/AETFAT/UNSO vegetation map of Africa. Paris, France: UNESCO. DOI: 10.2307/
 2260340.

Wright, A. J., W. D. A. Wardle, W. R. Callaway & A. Gaxiola (2017). 'The overlooked role of facilitation in biodiversity experiments'. In: Trends in Ecology & Evolution 32, pp. 383–390.
 DOI: 10.1016/j.tree.2017.02.011.

692

693

694

695

Zhang, K., S.-C. Chen, D. Whitman, M.-L. Shyu, J. Yan & C. Zhang (2003). 'A progressive morphological filter for removing nonground measurements from airborne LIDAR data'. In: *IEEE Transactions on Geoscience and Remote Sensing* 41.4, pp. 872–882. DOI: 10.1109/tgrs. 2003.810682.

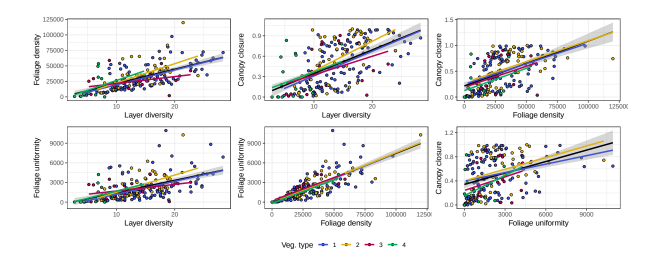


Figure 13: Bivariate scatter plots of subplot level canopy complexity metrics.

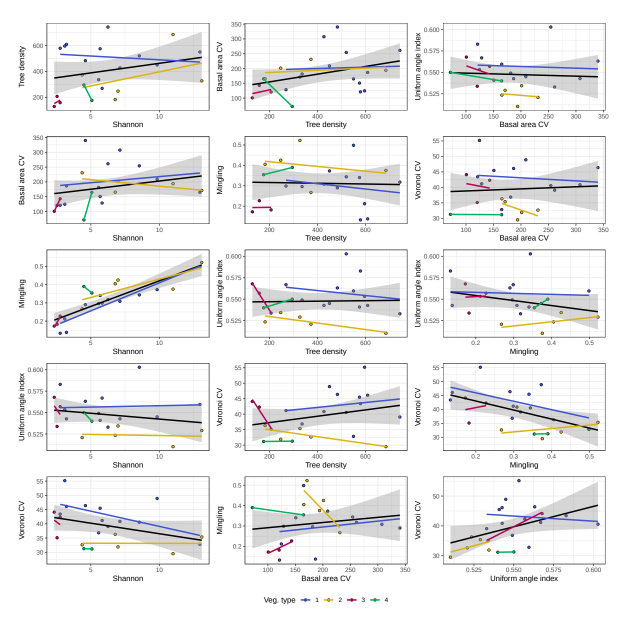


Figure 14: Bivariate scatter plots of plot level diversity and stand structural metrics.

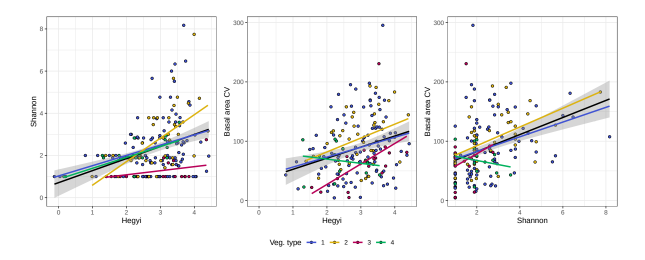


Figure 15: Bivariate scatter plots of subplot level diversity and stand structural metrics.

Table 6: Summary statistics of bivariate linear models comparing canopy complexity metrics with diversity and stand structural metrics, grouped by vegetation type. Note that models plot level canopy complexity metrics could not be fitted for Cluster 4, as this cluster only contained two plots. Slope refers to the slope of the predictor term in the model, \pm 1 standard error. T is the t-value of the slope of the predictor term in the model, Asterisks indicate the p-value of these terms (***<0.001, **<0.01, *<0.05).

Response	Predictor	Cluster	Slope	F	\mathbb{R}^2	T
		1	$7.3e + 01 \pm 3.7e + 01$	4.0(2,97)	0.04	1.99*
D-1:	D1 CV	2	$1.1e + 02 \pm 7.9e + 01$	2.1(2,38)	0.05	1.44
Foliage density	Basal area CV	3	$1.4e + 01 \pm 7.2e + 01$	0.0(2,14)	0.00	0.20
		4	$1.6e + 01 \pm 2.0e + 02$	0.0(2,12)	0.00	0.08
		1	$5.9e+03\pm2.1e+03$	8.2(2,102)	0.07	2.86**
Foliage density	Hegyi	2	$1.4e + 04 \pm 3.6e + 03$	15.2(2,40)	0.28	3.90***
ronage density	negyi	3	$6.6e + 03 \pm 3.0e + 03$	4.8(2,23)	0.17	2.18*
		4	$1.5e + 01 \pm 5.5e + 03$	0.0(2,13)	0.00	0.00
		1	$2.2e+03\pm1.3e+03$	2.8(2,102)	0.03	1.67
T-1:	Shannon	2	$3.8e + 03 \pm 2.4e + 03$	2.6(2,39)	0.06	1.61
Foliage density	Shannon	3	$1.1e+04\pm6.5e+03$	3.1(2,20)	0.13	1.77
		4	$-6.5e + 03 \pm 6.5e + 03$	1.0(2,13)	0.07	-1.01
		1	$1.7 \text{e-}04 \pm 6.0 \text{e-}04$	0.1(2,97)	0.00	0.28
Canopy closure	Basal area CV	2	$2.9e-03\pm1.1e-03$	6.9(2,39)	0.15	2.62*
Canopy closure	Dasar area C v	3	$4.2e-03\pm1.1e-03$	15.1(2,14)	0.52	3.89**
		4	$-4.6e-03\pm3.0e-03$	2.2(2,12)	0.16	-1.50
		1	$2.2 \text{e-} 01 \pm 2.8 \text{e-} 02$	62.3(2,102)	0.38	7.89***
Canopy closure	Цотті	2	$2.6e-01\pm5.1e-02$	27.0(2,41)	0.40	5.19***
Canopy closure	Hegyi	3	$2.8e-01\pm4.0e-02$	50.7(2,23)	0.69	7.12***
		4	$1.7e-01\pm 8.0e-02$	4.5(2,13)	0.26	2.12
		1	$3.1e-03\pm2.2e-02$	0.0(2,102)	0.00	0.14
Canany alagues	Shannon	2	$1.1e-01\pm3.2e-02$	12.1(2,40)	0.23	3.48**
Canopy closure	SHAIIIIOH	3	$2.3e-01\pm1.4e-01$	2.9(2,20)	0.13	1.69

		4	$6.7e-02\pm1.1e-01$	0.4(2,13)	0.03	0.60
				. , ,	0.03	0.92
		$\frac{1}{2}$	$3.7e+00\pm4.0e+00$ $4.5e+00\pm7.4e+00$	0.9(2,97)	0.01	0.92 0.61
Foliage uniformity	Basal area CV	$\frac{2}{3}$		0.4(2,38)	0.01 0.02	-0.59
			$-3.5e+00\pm5.9e+00$	0.4(2,14)		
		4	$-9.3e-01\pm1.5e+01$	0.0(2,12)	0.00	-0.06
		1	$2.2e+02\pm2.3e+02$	1.0(2,102)	0.01	0.98
Foliage uniformity	Hegyi	2	$7.5e + 02 \pm 3.7e + 02$	4.0(2,40)	0.09	2.00
ronage annormity	1108,11	3	$4.5e + 02 \pm 2.6e + 02$	2.9(2,23)	0.11	1.72
		4	$-7.5e + 01 \pm 4.0e + 02$	0.0(2,13)	0.00	-0.19
		1	$2.3e+02\pm1.4e+02$	2.6(2,102)	0.02	1.61
Foliage uniformity	Channon	2	$8.6e + 01 \pm 2.2e + 02$	0.1(2,39)	0.00	0.38
ronage uniformity	Shaimon	3	$1.3e+03\pm5.1e+02$	6.1(2,20)	0.23	2.48*
		4	$-5.9e + 02 \pm 4.7e + 02$	1.6(2,13)	0.11	-1.27
		1	$2.5e-02\pm9.3e-03$	7.1(2,97)	0.07	2.66**
T 1: '4	D 1 CV	2	$3.9e-02\pm1.4e-02$	8.0(2,38)	0.17	2.83**
Layer diversity	Basal area CV	3	$2.7e-02\pm2.3e-02$	1.3(2,14)	0.09	1.15
		4	$2.1e-02\pm3.1e-02$	0.5(2,12)	0.04	0.67
		1	$2.7e + 00 \pm 4.9e - 01$	29.1(2,102)	0.22	5.39***
T 1	TT .	2	$2.0e+00\pm7.5e-01$	7.1(2,40)	0.15	2.66*
Layer diversity	Hegyi	3	$1.9e + 00 \pm 1.0e + 00$	3.6(2,23)	0.13	1.89
		4	$1.1e+00\pm 8.5e-01$	1.8(2,13)	0.12	1.33
		1	$1.0e + 00 \pm 3.4e - 01$	8.7(2,102)	0.08	2.95**
		$\frac{1}{2}$	$9.5e-01\pm4.3e-01$	4.8(2,39)	0.11	2.18*
Layer diversity	Shannon	3	$4.9e + 00 \pm 1.8e + 00$	7.2(2,20)	0.26	2.68*
		$\overline{4}$	$1.8e-01\pm1.1e+00$	0.0(2,13)	0.00	0.16
		1	$1.2e-01\pm6.9e-02$	2.9(2,6)	0.33	1.72
		2	$-3.2e-01\pm2.9e-01$	1.2(2,3)	0.29	-1.10
Canopy roughness	Basal area CV	3	$3.5e-01\pm4.7e-01$	0.6(2,1)	0.36	0.74
		4	0.00 01±1.70 01	0.0(2,1)	0.00	0.11
		1	$2.6e-01\pm1.2e+00$	0.0(2,6)	0.01	0.22
		$\frac{1}{2}$	$4.6e + 00 \pm 1.9e + 00$	6.1(2,3)	0.67	2.48
Canopy roughness	Voronoi CV	3	$1.8e + 00 \pm 1.9e + 00$	1.0(2,1)	0.49	0.99
		4	1.00 00±1.00 00	1.0(2,1)	0.40	0.00
		1	$-4.2e+01\pm5.7e+01$	0.5(2,6)	0.08	-0.74
		$\frac{1}{2}$	$1.6e + 01 \pm 9.7e + 01$	$0.0(2,0) \\ 0.0(2,3)$	0.08	-0.74 0.17
Canopy roughness	Mingling	$\frac{2}{3}$	$3.5e+02\pm2.5e+02$	2.0(2,1)	0.67	1.42
		4	3.9e+02±2.9e+02	2.0(2,1)	0.07	1.42
			4.20.00 4.5-00	0.0(9.6)	0.19	0.06
		1	$-4.3e-02\pm4.5e-02$ $-5.9e-02\pm3.1e-02$	0.9(2,6) $3.6(2,3)$	$0.13 \\ 0.54$	-0.96 -1.89
Canopy roughness	Tree density	$\frac{2}{3}$	$-5.9e-02\pm3.1e-02$ $-1.8e-01\pm2.6e-01$	* ' '	0.54 0.31	-1.89 -0.68
		3 4	-1.0e-U1±2.0e-U1	0.5(2,1)	0.51	-0.08
			0.9-100117 100	1 7(0.0)	0.00	1 20
		1	$-2.3e+0.0\pm1.7e+0.0$	1.7(2.6)	0.22	-1.32
Canopy roughness	Shannon	2	$-1.4e + 00 \pm 2.4e + 00$	0.4(2,3)	0.11	-0.60
_		$\frac{3}{4}$	$3.4e + 01 \pm 4.7e + 01$	0.5(2,1)	0.34	0.72
		4				
		1	$-7.4e + 01 \pm 2.6e + 02$	0.1(2,6)	0.01	-0.28

Canopy roughness Uniform angle index

Canopy height Basal area CV			2	$4.1e+02\pm9.5e+02$	0.2(2,3)	0.06	0.43
Canopy height Basal area CV			3	$4.4e + 02 \pm 5.7e + 02$	0.6(2,1)	0.37	0.76
Canopy height Basal area CV			4				
Canopy height Basal area CV			1	$-6.5e-03\pm6.1e-03$	1.1(2,6)	0.16	-1.07
Canopy height	G 1 1 1 1	D 1 CIV	2	$4.3e-02\pm4.0e-02$	* ' '	0.28	1.08
$ \begin{array}{c} \text{Canopy height} \\ Can$	Canopy height	Basal area CV		$-3.1e-02\pm8.7e-03$	* ' '	0.92	-3.51
$ \begin{array}{c} \text{Canopy height} \\ \text{Canopy height} \\ \text{Canopy height} \\ \text{Voronoi CV} \\ & 2 \\ & 3 \\ & -1.8e-02\pm1.4e-01 \\ & 4 \\ $					(
Canopy height vorono CV 3 -1.8e-02±1.4e-01 0.0(2,1) 0.02 -0.13			1	-1.0e-01±8.6e-02	1.5(2,6)	0.20	-1.21
Canopy height Mingling 1	Canany baimbt	Vananai CV	2	$-7.0e-01\pm2.0e-01$	12.7(2,3)	0.81	-3.57*
$ \text{Canopy height} \text{Mingling} \begin{array}{c} 1 \\ 2 \\ -3.3 \text{e} + 00 \pm 3.8 \text{e} + 00 \\ 3 \\ -2.3 \text{e} + 00 \pm 1.3 \text{e} + 01 \\ 4 \\ \end{array} \begin{array}{c} 0.1(2.3) \\ 0.1(2.3) \\ 0.02 \\ -0.25 \\ 0.00 \\ -0.25 \\ 0.00 \\ -0.09 \\ -0.09 \\ 0.00 \\ -0.09 \\ -0.00 \\ $	Canopy neight	voronoi Cv	3	$-1.8e-02\pm1.4e-01$	0.0(2,1)	0.02	-0.13
$ \begin{array}{c} \text{Canopy height} \\ \text{Canopy height} \\ Paramote of the properties of the pr$			4				
Canopy height Free density			1	$6.8e + 00 \pm 3.8e + 00$	3.2(2,6)	0.34	1.78
Canopy height Pree density	Canany haight	Mingling		$-3.3e+00\pm1.3e+01$	0.1(2,3)	0.02	-0.25
$ \text{Canopy height} \text{Tree density} \begin{array}{c} 1 \\ 2 \\ 3 \\ 3 \\ -1.0e - 03 \pm 1.7e - 02 \\ 4 \\ \end{array} \begin{array}{c} 0.0(2,6) \\ 0.00 \\ -0.09 \\ -0.06 \\ -0.06 \\ \end{array} \\ \text{Canopy height} \text{Tree density} \begin{array}{c} 1 \\ 2 \\ 3 \\ -1.0e - 03 \pm 1.7e - 02 \\ -0.0(2,1) \\ 0.00 \\ -0.06 \\ \end{array} \\ \end{array} \begin{array}{c} 0.06 \\ -0.06 \\ -0.06 \\ -0.06 \\ \end{array} \\ \text{Canopy height} \text{Shannon} \begin{array}{c} 1 \\ 2 \\ 2 \\ 1.7e - 01 \pm 3.3e - 01 \\ 3 \\ -3.0e + 00 \pm 9.0e - 01 \\ 4 \\ \end{array} \begin{array}{c} 0.3(2,3) \\ 0.08 \\ 0.52 \\ 3 \\ -3.0e + 00 \pm 9.0e - 01 \\ 11.1(2,1) \\ 0.92 \\ -3.32 \\ \end{array} \\ \text{Canopy height} \text{Uniform angle index} \begin{array}{c} 1 \\ 2 \\ 2 \\ -7.2e + 01 \pm 1.3e + 02 \\ 3 \\ 3 \\ -6.0e - 02 \pm 3.9e + 01 \\ 4 \\ \end{array} \begin{array}{c} 0.2(2,6) \\ 0.04 \\ 0.02,3) \\ 0.09 \\ -0.56 \\ 0.00 \\ 0.00 \\ 0.00 \\ \end{array} \\ \text{Canopy closure} \text{Basal area CV} \begin{array}{c} 1 \\ 2 \\ 2 \\ 3.5e - 03 \pm 3.5e - 03 \\ 3 \\ 1.9e - 03 \pm 5.3e - 03 \\ 3 \\ 1.9e - 03 \pm 5.3e - 03 \\ 0.1(2,1) \\ 0.11 \\ 0.11 \\ 0.35 \\ \end{array} \begin{array}{c} 0.03 \\ 0.53 \\ 0.56 \\ -0.2e \pm 4.6e - 03 \\ 0.96 \\ -0.2e \pm 3.9e - 03 \\ 0.1(2,1) \\ 0.11 \\ 0.13 \\ 0.97 \\ -0.539 \\ \end{array} \\ \text{Canopy closure} \text{Mingling} \begin{array}{c} 1 \\ 9.3e - 03 \pm 8.2e - 03 \\ 3 \\ -2.5e - 02 \pm 4.6e - 03 \\ 2.90(2,1) \\ 0.00 \\ 0.0(2,1) \\ 0.00 \\ 0.01 \\ 0.01 \\ 0.01 \\ 0.02 \\ -0.63 \\ 3 \\ 3.6e - 02 \pm 4.1e + 00 \\ 0.0(2,1) \\ 0.00 \\ 0.02 \\ \end{array} \\ \text{Canopy closure} \text{Mingling} \begin{array}{c} 1 \\ 1 \\ -1.6e - 01 \pm 5.1e - 01 \\ 0.1(2,10) \\ 0.01 \\ 0.02 \\ -0.63 \\ 3 \\ 3.6e - 02 \pm 4.1e + 00 \\ 0.0(2,1) \\ 0.00 \\ 0.02 \\ \end{array} \\ \text{Canopy closure} \text{Tree density} \begin{array}{c} 1 \\ 1 \\ 2 \\ 8.5e - 04 \pm 2.4e - 04 \\ 4 \\ 12.2(2,3) \\ 0.80 \\ 3.50^* \\ 3.30e - 03 \pm 4.3e - 06 \\ 499683.9(2,1) \\ 1.00 \\ 0.02 \\ -0.45 \\ 0.00 \\ 0.02 \\ -0.45 \\ 0.00 \\ 0.02 \\ -0.45 \\ 0.00 \\ 0.02 \\ -0.45 \\ 0.00 \\ 0.02$	Canopy neight	winging	3	$-2.3e+01\pm9.3e-01$	619.2(2,1)	1.00	-24.88*
$ \begin{array}{c} \text{Canopy height} \\ \text{Canopy closure} \\ C$			4				
			1	$-3.5e-04\pm3.8e-03$	0.0(2,6)	0.00	-0.09
Canopy height Shannon $ \begin{array}{c} 3 \\ 4 \\ \\ 1 \\ 2.8e-01\pm 1.1e-01 \\ 3 \\ 3 \\ -3.0e+00\pm 9.0e-01 \\ 4 \\ \\ 2 \\ 1.7e-01\pm 3.3e-01 \\ 3 \\ -3.0e+00\pm 9.0e-01 \\ 11.1(2,1) \\ 0.92 \\ -3.32 \\ \\ 4 \\ \\ \\ Canopy height \\ Canopy closure \\ Canopy closu$	Canany haight	Tron donaity	2	$8.6e-03\pm4.0e-03$	4.7(2,3)	0.61	2.16
$ \text{Canopy height} \text{Shannon} \begin{cases} 1 \\ 2 \\ 2 \\ 1.7e-01\pm3.3e-01 \\ 3 \\ -3.0e+00\pm9.0e-01 \\ 4 \end{cases} \begin{array}{c} 0.3(2,3) \\ 0.08 \\ 0.52 \\ 0.3(2,3) \\ 0.08 \\ 0.52 \\ 0.32 \\ 0.08 \\ 0.52 \\ 0.32 \\ 0.08 \\ 0.52 \\ 0.32 \\ 0.09 \\ 0.00 $	Canopy neight	free density	3	$-1.0e-03\pm1.7e-02$	0.0(2,1)	0.00	-0.06
$ \begin{array}{c} \text{Canopy height} \text{Shannon} \\ & 2 \\ & 3 \\ & -3.0 \text{e} + 00 \pm 9.0 \text{e} - 01 \\ & 11.1(2,1) \\ & 0.92 \\ & -3.32 \\ & & & & & & & & & & \\ & & & & & & & $			4				
Canopy height Shannon 3 -3.0e+00±9.0e-01 11.1(2,1) 0.92 -3.32 4 Canopy height Uniform angle index $\begin{bmatrix} 1 & 1.0e+01\pm2.1e+01 & 0.2(2,6) & 0.04 & 0.49 \\ 2 & -7.2e+01\pm1.3e+02 & 0.3(2,3) & 0.09 & -0.56 \\ 3 & 6.0e-02\pm3.9e+01 & 0.0(2,1) & 0.00 & 0.00 \end{bmatrix}$ Canopy closure Basal area CV $\begin{bmatrix} 1 & 3.6e-04\pm6.9e-04 & 0.3(2,10) & 0.03 & 0.53 \\ 2 & 3.5e-03\pm3.5e-03 & 1.0(2,3) & 0.24 & 0.98 \\ 3 & 1.9e-03\pm5.3e-03 & 0.1(2,1) & 0.11 & 0.35 \end{bmatrix}$ Canopy closure Voronoi CV $\begin{bmatrix} 1 & 9.3e-03\pm8.2e-03 & 1.3(2,10) & 0.11 & 1.13 \\ 2 & -6.6e-02\pm7.9e-03 & 69.7(2,3) & 0.96 & -8.35** \\ 3 & -2.5e-02\pm4.6e-03 & 29.0(2,1) & 0.97 & -5.39 \end{bmatrix}$ Canopy closure Mingling $\begin{bmatrix} 1 & -1.6e-01\pm5.1e-01 & 0.1(2,10) & 0.01 & -0.31 \\ 2 & -6.9e-01\pm1.1e+00 & 0.4(2,3) & 0.12 & -0.63 \\ 3 & 7.6e-02\pm4.1e+00 & 0.0(2,1) & 0.00 & 0.02 \end{bmatrix}$ Canopy closure Tree density $\begin{bmatrix} 1 & 1.4e-04\pm4.0e-04 & 0.1(2,10) & 0.01 & 0.36 \\ 2 & 8.5e-04\pm2.4e-04 & 12.2(2,3) & 0.80 & 3.50* \\ 3 & 3.0e-03\pm4.3e-06 & 499683.9(2,1) & 1.00 & 706.88*** \\ 4 & & & & & & & & & & & & & & & & & &$			1	$2.8\text{e-}01\pm1.1\text{e-}01$	7.1(2,6)	0.54	2.66*
Canopy height Uniform angle index $\begin{bmatrix} 3 & -3.0e+00\pm 9.0e-01 \\ 4 & & & & \\ 1 & 1.0e+01\pm 2.1e+01 \\ 2 & -7.2e+01\pm 1.3e+02 \\ 3 & 6.0e-02\pm 3.9e+01 \\ 4 & & & \\ \end{bmatrix} 0.0(2,1) 0.00 0.00 0.00$ Canopy closure Basal area CV $\begin{bmatrix} 1 & 3.6e-04\pm 6.9e-04 \\ 2 & 3.5e-03\pm 3.5e-03 \\ 3 & 1.9e-03\pm 5.3e-03 \\ 4 & & & \\ \end{bmatrix} 0.1(2,1) 0.11 0.35 0.53 0.24 0.98 0.1(2,1) 0.11 0.35 0.24 0.98 0.1(2,1) 0.11 0.35 0.24 0.98 0.1(2,1) 0.11 0.35 0.24 0.98 0.1(2,1) 0.11 0.35 0.24 0.98 0.1(2,1) 0.11 0.35 0.24 0.98 0.1(2,1) 0.11 0.35 0.24 0.98 0.1(2,1) 0.11 0.35 0.24 0.98 0.1(2,1) 0.11 0.35 0.25 0.25e-02\pm 4.6e-03 0.1(2,1) 0.11 0.35 0.25 0.25e-02\pm 4.6e-03 0.1(2,1) 0.97 0.25 0.25 0.25e-02\pm 4.6e-03 0.1(2,1) 0.97 0.25 0.25 0.25e-02\pm 4.6e-03 0.1(2,1) 0.01 0.01 0.36 0.25 0.25 0.25e-02\pm 4.1e+00 0.0(2,1) 0.00 0.02 0.02 0.02 0.02 0.02 0.02 0.0$	Canany baimbt	Charanan	2	$1.7e-01\pm3.3e-01$	0.3(2,3)	0.08	0.52
$ \text{Canopy height} \text{Uniform angle index} \begin{cases} 1 & 1.0\text{e} + 01 \pm 2.1\text{e} + 01 & 0.2(2,6) & 0.04 & 0.49 \\ 2 & -7.2\text{e} + 01 \pm 1.3\text{e} + 02 & 0.3(2,3) & 0.09 & -0.56 \\ 3 & 6.0\text{e} - 02 \pm 3.9\text{e} + 01 & 0.0(2,1) & 0.00 & 0.00 \\ 4 & & & & & & \\ \\ \text{Canopy closure} \end{cases} \text{Basal area CV} \begin{cases} 1 & 3.6\text{e} - 04 \pm 6.9\text{e} - 04 & 0.3(2,10) & 0.03 & 0.53 \\ 2 & 3.5\text{e} - 03 \pm 3.5\text{e} - 03 & 1.0(2,3) & 0.24 & 0.98 \\ 3 & 1.9\text{e} - 03 \pm 5.3\text{e} - 03 & 0.1(2,1) & 0.11 & 0.35 \\ 4 & & & & & \\ \\ \text{Canopy closure} \end{cases} \text{Voronoi CV} \begin{cases} 1 & 9.3\text{e} - 03 \pm 8.2\text{e} - 03 & 1.3(2,10) & 0.11 & 1.13 \\ 2 & -6.6\text{e} - 02 \pm 7.9\text{e} - 03 & 69.7(2,3) & 0.96 & -8.35^{**} \\ 3 & -2.5\text{e} - 02 \pm 4.6\text{e} - 03 & 29.0(2,1) & 0.97 & -5.39 \\ 4 & & & & \\ \\ \text{Canopy closure} \end{cases} \text{Mingling} \begin{cases} 1 & -1.6\text{e} - 01 \pm 5.1\text{e} - 01 & 0.1(2,10) & 0.01 & -0.31 \\ 2 & -6.9\text{e} - 01 \pm 1.1\text{e} + 00 & 0.4(2,3) & 0.12 & -0.63 \\ 3 & 7.6\text{e} - 02 \pm 4.1\text{e} + 00 & 0.0(2,1) & 0.00 & 0.02 \\ 4 & & & & \\ \\ \text{Canopy closure} \end{cases} \text{Tree density} \begin{cases} 1 & 1.4\text{e} - 04 \pm 4.0\text{e} - 04 & 0.1(2,10) & 0.01 & 0.36 \\ 2 & 8.5\text{e} - 04 \pm 2.4\text{e} - 04 & 12.2(2,3) & 0.80 & 3.50^* \\ 3 & 3.0\text{e} - 03 \pm 4.3\text{e} - 06 & 499683.9(2,1) & 1.00 & 706.88^{****} \\ 4 & & & & \\ \\ \text{Canopy closure} \end{cases} \begin{cases} 1 & -7.6\text{e} - 03 \pm 1.7\text{e} - 02 & 0.2(2,10) & 0.02 & -0.45 \\ 4 & & & & \\ \\ \end{array} \begin{cases} 2 & 8.5\text{e} - 03 \pm 3.0\text{e} - 02 & 0.1(2,3) & 0.03 & 0.28 \\ \end{cases} \end{cases}$	Canopy neight	Snannon	3	$-3.0e + 00 \pm 9.0e - 01$	11.1(2,1)	0.92	-3.32
Canopy height Uniform angle index			4				
Canopy closure Basal area CV $ \begin{array}{c} 3 \\ 4 \\ \end{array} \begin{array}{c} 6.0e-02\pm3.9e+01 \\ 4 \\ \end{array} \begin{array}{c} 0.0(2,1) \\ 0.00 \\ \end{array} \begin{array}{c} 0.00 \\ 0.00 \\ \end{array} \\ 0.00$			1	$1.0e+01\pm2.1e+01$	0.2(2,6)	0.04	0.49
Canopy closure Basal area CV $ \begin{array}{c} 3 \\ 4 \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\$	C	II:fl. : l	2	$-7.2e + 01 \pm 1.3e + 02$	0.3(2,3)	0.09	-0.56
Canopy closure Basal area CV $ \begin{bmatrix} 1 & 3.6e-04\pm6.9e-04 & 0.3(2,10) & 0.03 & 0.53 \\ 2 & 3.5e-03\pm3.5e-03 & 1.0(2,3) & 0.24 & 0.98 \\ 3 & 1.9e-03\pm5.3e-03 & 0.1(2,1) & 0.11 & 0.35 \end{bmatrix} $ Canopy closure Voronoi CV $ \begin{bmatrix} 1 & 9.3e-03\pm8.2e-03 & 1.3(2,10) & 0.11 & 1.13 \\ 2 & -6.6e-02\pm7.9e-03 & 69.7(2,3) & 0.96 & -8.35** \\ 3 & -2.5e-02\pm4.6e-03 & 29.0(2,1) & 0.97 & -5.39 \end{bmatrix} $ Canopy closure Mingling $ \begin{bmatrix} 1 & -1.6e-01\pm5.1e-01 & 0.1(2,10) & 0.01 & -0.31 \\ 2 & -6.9e-01\pm1.1e+00 & 0.4(2,3) & 0.12 & -0.63 \\ 3 & 7.6e-02\pm4.1e+00 & 0.0(2,1) & 0.00 & 0.02 \end{bmatrix} $ Canopy closure Tree density $ \begin{bmatrix} 1 & 1.4e-04\pm4.0e-04 & 0.1(2,10) & 0.01 & 0.36 \\ 2 & 8.5e-04\pm2.4e-04 & 12.2(2,3) & 0.80 & 3.50* \\ 3 & 3.0e-03\pm4.3e-06 & 499683.9(2,1) & 1.00 & 706.88*** \\ 4 & & & & & & & \end{bmatrix} $	Canopy neight	Omform angle index	3	$6.0e-02\pm3.9e+01$	0.0(2,1)	0.00	0.00
Canopy closure Basal area CV $ \begin{array}{c} 2 \\ 3 \\ 3 \\ 1.9e-03\pm5.3e-03 \\ 4 \\ \end{array} \begin{array}{c} 1.0(2,3) \\ 0.24 \\ 0.98 \\ 0.1(2,1) \\ 0.11 \\ 0.35 \\ \end{array} $ $ \begin{array}{c} 0.24 \\ 0.98 \\ 0.35 \\ 0.1(2,1) \\ 0.11 \\ 0.35 \\ \end{array} $ $ \begin{array}{c} 0.35 \\ 0.1(2,1) \\ 0.11 \\ 0.35 \\ \end{array} $ $ \begin{array}{c} 0.35 \\ 0.1(2,1) \\ 0.11 \\ 0.35 \\ \end{array} $ $ \begin{array}{c} 0.11 \\ 0.35 \\ 0.1(2,1) \\ 0.11 \\ 0.35 \\ \end{array} $ $ \begin{array}{c} 0.11 \\ 0.35 \\ 0.12 \\ 0.96 \\ -8.35 $			4				
Canopy closure Basal area CV $ \begin{array}{c} 3 \\ 4 \\ \end{array} \begin{array}{c} 1.9 \text{e-}03 \pm 5.3 \text{e-}03 \\ 4 \\ \end{array} \begin{array}{c} 0.1(2,1) \\ \end{array} \begin{array}{c} 0.11 \\ 0.35 \\ \end{array} \\ \end{array} $ Canopy closure Voronoi CV $ \begin{array}{c} 1 \\ 2 \\ 3 \\ -6.6 \text{e-}02 \pm 7.9 \text{e-}03 \\ 3 \\ -2.5 \text{e-}02 \pm 4.6 \text{e-}03 \\ \end{array} \begin{array}{c} 0.97(2,3) \\ 29.0(2,1) \\ 0.97 \\ -5.39 \\ \end{array} \begin{array}{c} 0.96 \\ -8.35 \text{**} \\ -8.5 \text{e-}02 \pm 4.6 \text{e-}03 \\ \end{array} \\ \text{Canopy closure} \end{array} $ Mingling $ \begin{array}{c} 1 \\ 2 \\ -6.9 \text{e-}01 \pm 5.1 \text{e-}01 \\ 2 \\ -6.9 \text{e-}01 \pm 1.1 \text{e+}00 \\ 3 \\ 7.6 \text{e-}02 \pm 4.1 \text{e+}00 \\ 0.0(2,1) \\ 0.00 \\ 0.002 \\ \end{array} \begin{array}{c} 0.01 \\ 0.00 \\ 0.02 \\ \end{array} $ Canopy closure $ \begin{array}{c} 1 \\ 1.4 \text{e-}04 \pm 4.0 \text{e-}04 \\ 3 \\ 3.0 \text{e-}03 \pm 4.3 \text{e-}04 \\ \end{array} \begin{array}{c} 0.1(2,10) \\ 0.01 \\ 0.01 \\ 0.00 \\ 0.02 \\ \end{array} \begin{array}{c} 0.36 \\ 0.36 \\ 3.50 \text{*} \\ 3.0 \text{e-}03 \pm 4.3 \text{e-}06 \\ \end{array} \begin{array}{c} 0.9683.9(2,1) \\ 1.00 \\ 0.02 \\ -0.45 \\ 0.2(2,10) \\ 0.02 \\ -0.45 \\ 0.28 \\ \end{array} $			1	$3.6e-04\pm6.9e-04$	0.3(2,10)	0.03	0.53
Canopy closure Voronoi CV $ \begin{array}{c} 3 & 1.9e - 0.3 \pm 5.3e - 0.3 & 0.1(2,1) & 0.11 & 0.35 \\ 4 & & & & & & & & \\ 1 & 9.3e - 0.3 \pm 8.2e - 0.3 & 1.3(2,10) & 0.11 & 1.13 \\ 2 & -6.6e - 0.2 \pm 7.9e - 0.3 & 69.7(2,3) & 0.96 & -8.35 ** \\ 3 & -2.5e - 0.2 \pm 4.6e - 0.3 & 29.0(2,1) & 0.97 & -5.39 \\ 4 & & & & & \\ Canopy closure & Mingling & 1 & -1.6e - 0.1 \pm 5.1e - 0.1 & 0.1(2,10) & 0.01 & -0.31 \\ 2 & -6.9e - 0.1 \pm 1.1e + 0.0 & 0.4(2,3) & 0.12 & -0.63 \\ 3 & 7.6e - 0.2 \pm 4.1e + 0.0 & 0.0(2,1) & 0.00 & 0.02 \\ 4 & & & & & \\ Canopy closure & Tree density & 1 & 1.4e - 0.4 \pm 4.0e - 0.4 & 0.1(2,10) & 0.01 & 0.36 \\ 2 & 8.5e - 0.4 \pm 2.4e - 0.4 & 12.2(2,3) & 0.80 & 3.50 * \\ 3 & 3.0e - 0.3 \pm 4.3e - 0.6 & 499683.9(2,1) & 1.00 & 706.88 *** \\ 4 & & & & & \\ Canopy closure & Shannon & 2 & 8.5e - 0.3 \pm 3.0e - 0.2 & 0.1(2,3) & 0.03 & 0.28 \\ \end{array} $	Canany alaguna	Dagal area CV	2	$3.5e-03\pm3.5e-03$	1.0(2,3)	0.24	0.98
Canopy closure Voronoi CV	Canopy closure	Dasar area Cv	3	$1.9e-03\pm5.3e-03$	0.1(2,1)	0.11	0.35
Canopy closure Voronoi CV $ \begin{array}{c} 2 & -6.6 \text{e-}02 \pm 7.9 \text{e-}03 & 69.7(2,3) & 0.96 & -8.35^{**} \\ 3 & -2.5 \text{e-}02 \pm 4.6 \text{e-}03 & 29.0(2,1) & 0.97 & -5.39 \\ 4 & & & & & & \\ & & & & & & \\ & & & &$			4				
Canopy closure Voronoi CV $ \begin{array}{c} 3 \\ 4 \\ \hline \\ Canopy closure \end{array} \begin{array}{c} -2.5 \text{e-} 02 \pm 4.6 \text{e-} 03 \\ 4 \\ \hline \\ Canopy closure \end{array} \begin{array}{c} 29.0(2,1) \\ 4 \\ \hline \\ Canopy closure \end{array} \begin{array}{c} 0.97 \\ -5.39 \\ \hline \\ 1 \\ -1.6 \text{e-} 01 \pm 5.1 \text{e-} 01 \\ 2 \\ -6.9 \text{e-} 01 \pm 1.1 \text{e+} 00 \\ 3 \\ 7.6 \text{e-} 02 \pm 4.1 \text{e+} 00 \end{array} \begin{array}{c} 0.1(2,10) \\ 0.4(2,3) \\ 0.12 \\ -0.63 \\ 0.00 \\ 0.02 \\ \hline \\ Canopy closure \end{array} $			1	9.3e-03±8.2e-03	1.3(2,10)	0.11	1.13
Canopy closure Mingling $ \begin{array}{c} 3 \\ 4 \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\$	Canany alaguna	Vananai CV	2	$-6.6e-02\pm7.9e-03$	69.7(2,3)	0.96	-8.35**
Canopy closure Mingling	Canopy closure	voronoi C v	3	$-2.5e-02\pm4.6e-03$	29.0(2,1)	0.97	-5.39
Canopy closure Mingling			4				
Canopy closure Minging $3 - 7.6e-02\pm 4.1e+00 - 0.0(2,1) - 0.00 - 0.02$ Canopy closure Tree density $1 - 1.4e-04\pm 4.0e-04 - 0.1(2,10) - 0.01 - 0.36$ $2 - 8.5e-04\pm 2.4e-04 - 12.2(2,3) - 0.80 - 3.50*$ $3 - 3.0e-03\pm 4.3e-06 - 499683.9(2,1) - 1.00 - 706.88***$ $4 - 7.6e-03\pm 1.7e-02 - 0.2(2,10) - 0.02 - 0.45$ Canopy closure Shappon $2 - 8.5e-03\pm 3.0e-02 - 0.1(2,3) - 0.03 - 0.28$			1	$-1.6e-01\pm5.1e-01$	0.1(2,10)	0.01	-0.31
Canopy closure Tree density $ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	Canony alasuma	Mingling	2	$-6.9e-01\pm1.1e+00$	0.4(2,3)	0.12	-0.63
Canopy closure Tree density $ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	Canopy closure	wiingiing	3	$7.6e-02\pm4.1e+00$	0.0(2,1)	0.00	0.02
Canopy closure Tree density $ \begin{array}{ccccccccccccccccccccccccccccccccccc$			4				
Canopy closure Tree density $ \begin{array}{ccccccccccccccccccccccccccccccccccc$			1	$1.4e-04\pm4.0e-04$	0.1(2,10)	0.01	0.36
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	Conony alasma	Tron dong!t	2	$8.5e-04\pm2.4e-04$	* ' '	0.80	3.50*
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	Canopy closure	rree density		$3.0e-03\pm4.3e-06$,	1.00	706.88***
Canopy closure Shappon 2 $8.5e-03\pm3.0e-02$ $0.1(2,3)$ 0.03 0.28			4		,		
Canopy closure Shappon 2 $8.5e-03\pm3.0e-02$ $0.1(2,3)$ 0.03 0.28			1	$-7.6e-03\pm1.7e-02$	0.2(2,10)	0.02	-0.45
Canony closure Shannon	Comonus -1	Chamman	2	$8.5e-03\pm3.0e-02$, ,	0.03	0.28
	Canopy closure	энаннон		$1.9e-01\pm5.2e-01$	` ' '	0.12	0.37

		4				
		1	$-3.9e+00\pm2.3e+00$	2.9(2,10)	0.23	-1.71
Canany alaguna	Uniform angle index	2	$-1.2e + 01 \pm 9.3e + 00$	1.7(2,3)	0.36	-1.30
Canopy closure	Omform angle index	3	$-6.9e + 00 \pm 3.9e - 01$	306.2(2,1)	1.00	-17.50*
		4		, ,		
		1	$-4.5e + 01 \pm 2.9e + 01$	2.3(2,6)	0.28	-1.52
77.14	D 1 CT1	2	$1.5e + 02 \pm 1.4e + 02$	1.1(2,3)	0.27	1.05
Foliage density	Basal area CV	3	$1.8e + 02 \pm 8.9e + 01$	4.2(2,1)	0.81	2.06
		4	1.00 02 100 01	112(2,1)	0.01	
		1	$3.5e + 0.1 \pm 5.0e + 0.02$	0.0(2,6)	0.00	0.07
		2	$-7.7e + 02 \pm 1.5e + 03$	0.3(2,3)	0.08	-0.51
Foliage density	Voronoi CV	3	$2.7e + 02 \pm 8.7e + 02$	0.1(2,1)	0.09	0.31
		4	2	011(= ,1)	0.00	0.01
		1	$4.5e + 03 \pm 2.5e + 04$	0.0(2,6)	0.01	0.18
		2	$8.0e + 0.02 \pm 4.7e + 0.04$	0.0(2,3)	0.00	0.02
Foliage density	Mingling	3	$1.5e + 0.00 \pm $	54.1(2,1)	0.98	7.35
		4	1.00 00 ± 2.00 04	04.1(2,1)	0.50	1.00
		1	$8.8e + 00 \pm 2.0e + 01$	0.2(2,6)	0.03	0.45
		2	$1.1e+01\pm2.1e+01$	0.2(2,3) $0.3(2,3)$	0.08	0.51
Foliage density	Tree density	3	$-1.3e+0.1\pm1.1e+0.2$	0.0(2,3) $0.0(2,1)$	0.01	-0.12
		$\frac{3}{4}$	-1.90 01±1.10 02	0.0(2,1)	0.01	0.12
		1	$2.5e + 02 \pm 8.1e + 02$	0.1(2,6)	0.02	0.31
		2	$5.0e + 02 \pm 0.1e + 02$ $5.0e + 02 \pm 1.2e + 03$	0.2(2,3)	0.02	0.42
Foliage density	Shannon	3	$1.8e + 04 \pm 9.1e + 03$	3.9(2,1)	0.80	1.98
		4	1.06+04±9.16+03	3.9(2,1)	0.00	1.90
		1	$-1.1e + 05 \pm 1.0e + 05$	1.3(2,6)	0.18	-1.15
		2	$1.2e + 05 \pm 4.7e + 05$	0.1(2,3)	0.02	0.25
Foliage density	Uniform angle index	3	$4.3e + 04 \pm 2.5e + 05$	0.1(2,3) $0.0(2,1)$	0.03	0.18
		$\frac{3}{4}$	4.90 0412.90 00	0.0(2,1)	0.05	0.10
		1	-1.0e-01±6.1e-01	0.0(2,6)	0.00	-0.17
		2	$-2.2e+00\pm2.2e+00$	1.1(2,3)	0.26	-1.03
Canopy rugosity	Basal area CV	3	$8.7e + 00 \pm 5.4e + 00$	2.6(2,1)	0.73	1.62
		$\frac{3}{4}$	0.7e+00±3.4e+00	2.0(2,1)	0.15	1.02
		1	$7.9e + 00 \pm 8.2e + 00$	0.9(2,6)	0.13	0.96
		2	$3.5e+00\pm0.2e+00$ $3.5e+01\pm1.3e+01$	6.8(2,3)	0.69	2.61
Canopy rugosity	Voronoi CV	3	$1.8e + 01 \pm 4.2e + 01$	0.2(2,1)	0.05	0.42
		$\frac{3}{4}$	1.06+01±4.26+01	0.2(2,1)	0.10	0.42
		1	$-5.9e + 02 \pm 3.6e + 02$	2.7(2,6)	0.31	-1.63
		2	$8.5e + 02 \pm 5.2e + 02$	2.7(2,3) $2.7(2,3)$	0.31 0.47	1.63
Canopy rugosity	Mingling	$\frac{2}{3}$	$7.2e+03\pm1.7e+03$	17.6(2,1)	0.47 0.95	4.19
		3 4	1.26+00±1.76+00	11.0(4,1)	0.30	4.13
		1	-1.9e-01±3.4e-01	0.3(2,6)	0.05	-0.56
		2	$-4.6e-01\pm 3.4e-01$	4.9(2,3)	0.62	-2.22
Canopy rugosity	Tree density	3	$-1.2e + 00 \pm 5.4e + 00$	0.0(2,1)	0.02	-2.22 -0.22
			-1.46〒UU上り.46 〒 UU	0.0(2,1)	0.00	-0.22
		4	0.4 +04+4.0 +04	F 0/0 3\	0.17	0.01
		1	$-2.4e + 01 \pm 1.0e + 01$	5.3(2,6)	0.47	-2.31
Canopy rugosity	Shannon					

		2 3 4	$6.4e+00\pm1.8e+01$ $8.5e+02\pm5.4e+02$	0.1(2,3) 2.5(2,1)	0.04 0.71	0.35 1.57
Canopy rugosity	Uniform angle index	1 2 3 4	$-2.6e+03\pm1.6e+03$ $1.0e+04\pm4.1e+03$ $3.4e+03\pm1.2e+04$	$ 2.5(2,6) \\ 6.1(2,3) \\ 0.1(2,1) $	0.30 0.67 0.07	-1.58 2.47 0.28

Table 3: Summary statistics of bivariate linear models comparing canopy complexity metrics with diversity and stand structural metrics across all vegetation types. Slope refers to the slope of the predictor term in the model, \pm 1 standard error. T is the t-value of the slope of the predictor term in the model, Asterisks indicate the p-value of these terms (***<0.001, **<0.01, *<0.05).

Response	Predictor	Slope	F	\mathbb{R}^2	Т
	Basal area CV	$8.7e + 01 \pm 3.0e + 01$	8.6(2,167)	0.05	2.93**
Foliage density	Hegyi	$7.8e + 03 \pm 1.6e + 03$	25.5(2,184)	0.12	5.05***
	Shannon	$3.2e+03\pm1.1e+03$	8.9(2,180)	0.05	2.98**
	Basal area CV	$1.2e-03\pm4.8e-04$	6.3(2,168)	0.04	2.52*
Canopy closure	Hegyi	$2.4e-01\pm2.1e-02$	132.8(2,185)	0.42	11.52***
	Shannon	$4.7e-02\pm1.7e-02$	7.3(2,181)	0.04	2.70**
	Basal area CV	$4.1e+00\pm3.0e+00$	1.9(2,167)	0.01	1.37
Foliage uniformity	Hegyi	$4.0e + 02 \pm 1.6e + 02$	6.2(2,184)	0.03	2.49*
	Shannon	$2.2e + 02 \pm 1.1e + 02$	4.1(2,180)	0.02	2.04*
	Basal area CV	$3.2 \text{e-} 02 \pm 7.6 \text{e-} 03$	17.6(2,167)	0.10	4.20***
Layer diversity	Hegyi	$2.7e + 00 \pm 3.9e - 01$	46.8(2,184)	0.20	6.84***
	Shannon	$1.1e+00\pm2.7e-01$	16.8(2,180)	0.09	4.10***
	Basal area CV	$3.0 \text{e-} 02 \pm 5.0 \text{e-} 02$	0.4(2,16)	0.02	0.60
	Voronoi CV	$7.5e-01\pm 5.9e-01$	1.6(2,16)	0.09	1.26
Canopy roughness	Mingling	$-2.8e+01\pm3.3e+01$	0.7(2,16)	0.04	-0.86
Canopy roughness	Tree density	$-2.6e-02\pm1.7e-02$	2.3(2,16)	0.12	-1.51
	Shannon	$-1.9e + 00 \pm 9.5e - 01$	4.0(2,16)	0.20	-2.01
	Uniform angle index	$1.6e + 02 \pm 1.6e + 02$	1.0(2,16)	0.06	0.98
	Basal area CV	$7.1e-03\pm7.3e-03$	0.9(2,16)	0.06	0.97
	Voronoi CV	$-4.7e-02\pm9.1e-02$	0.3(2,16)	0.02	-0.52
Canopy height	Mingling	$3.8e+00\pm4.8e+00$	0.6(2,16)	0.04	0.79
Canopy neight	Tree density	$4.3e-03\pm2.5e-03$	3.1(2,16)	0.16	1.76
	Shannon	$3.3e-01\pm1.3e-01$	6.0(2,16)	0.27	2.45*
	Uniform angle index	$-2.2e+01\pm2.4e+01$	0.8(2,16)	0.05	-0.90
	Basal area CV	$8.5 \text{e-}04 \pm 5.7 \text{e-}04$	2.2(2,20)	0.10	1.50
	Voronoi CV	$2.4e-03\pm5.8e-03$	0.2(2,20)	0.01	0.41
Canopy closure	Mingling	$7.2e-03\pm3.7e-01$	0.0(2,20)	0.00	0.02
Canopy closure	Tree density	$4.7e-04\pm1.9e-04$	6.3(2,20)	0.24	2.50*
	Shannon	$1.0e-02\pm1.2e-02$	0.7(2,20)	0.04	0.86
	Uniform angle index	$-3.4e + 00 \pm 1.7e + 00$	3.9(2,20)	0.16	-1.98
	Basal area CV	$5.8e + 01 \pm 3.2e + 01$	3.3(2,16)	0.17	1.80
	Voronoi CV	$5.8e + 02 \pm 4.1e + 02$	2.1(2,16)	0.11	1.43
Foliage density	Mingling	$6.6e + 03 \pm 2.3e + 04$	0.1(2,16)	0.01	0.29
romage density	Tree density	$3.0e+01\pm1.0e+01$	8.6(2,16)	0.35	2.93**
	Shannon	$1.1e + 03 \pm 6.9e + 02$	2.5(2,16)	0.13	1.57
	Uniform angle index	$-2.1e+04\pm1.1e+05$	0.0(2,16)	0.00	-0.18
	Basal area CV	$-1.0e + 00 \pm 5.3e - 01$	3.7(2,16)	0.19	-1.92
	Voronoi CV	$-6.0e + 00 \pm 7.0e + 00$	0.7(2,16)	0.04	-0.86
Canopy rugosity	Mingling	$1.3e + 02 \pm 3.8e + 02$	0.1(2,16)	0.01	0.33
Carropy ragosity	Tree density	$-5.2e-01\pm1.7e-01$	10.0(2,16)	0.38	-3.16**
	Shannon	$-1.3e+01\pm1.2e+01$	1.2(2,16)	0.07	-1.11
	Uniform angle index	$-1.8e + 03 \pm 1.9e + 03$	0.9(2,16)	0.06	-0.97

Table 4: Explanatory variables included in the best model for each subplot canopy structure variable. ΔAIC shows the difference in model AIC value compared to a null model which included only the random effects of vegetation type and plot. Positive ΔAIC values >2 indicate that the model is of better quality than the null model. R^2_c is the R^2 of the best model, while R^2_m is the R^2 of the model fixed effects only.

Response	Hegyi	Shannon	Basal area CV	$\Delta { m AIC}$	${ m R^2_{ c}}$	R^2_{m}
Layer diversity	✓	✓	√	37.0	0.50	0.17
Foliage density	\checkmark		\checkmark	47.6	0.27	0.09
Foliage uniformity	\checkmark			13.1	0.28	0.02
Canopy closure	\checkmark	\checkmark		101.9	0.60	0.46

Table 5: Explanatory variables included in the best linear model for each plot-level canopy complexity metric. ΔAIC shows the difference in model AIC value compared to a null model. Positive ΔAIC values >2 indicate that the model is of better quality than the null model.

Response	Shannon	Tree density	Basal area CV	Mingling	Uniform angle index	Voronoi CV	$\Delta { m AIC}$	\mathbb{R}^2	Prob.
Foliage density	✓					√	5.8	0.42	< 0.05
Canopy closure			\checkmark		\checkmark		5.8	0.42	< 0.05
Canopy height	\checkmark			\checkmark			8.2	0.49	< 0.01
Canopy roughness	\checkmark			\checkmark			2.5	0.30	0.07
Canopy rugosity		\checkmark			\checkmark		6.9	0.45	< 0.05