- <sup>1</sup> Title: An assessment of the biodiversity ecosystem function rela-
- 2 tionship in southern African woodlands
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- providers and the field assistance they received when collecting plot data.

# 17 2 Biosketch

- SEOSAW (A Socio-Ecological Observatory for Southern African Woodlands, https://seosaw.
- 19 github.io) aims to understand the response of southern African woodlands to global change.
- The goal of SEOSAW is to produce novel analyses of the determinants of ecosystem structure and
- <sup>21</sup> function for the southern Africa region, based on syntheses of plot data. Additionally the group
- 22 hopes to develop infrastructure for a long-term regional plan for plot remeasurement in the south-
- 23 ern African region. While working on a multitude of diverse projects in the dry tropics at large,
- 24 all authors have a broad interest in community ecology and ecosystem assemblage in southern
- 25 African woodlands.

# 26 Blinded Main Text File

- Title: An assessment of the biodiversity ecosystem function rela-
- 28 tionship in southern African woodlands
- 29 Running title: Ecosystem function in southern African woodlands

## 30 Abstract

31 Aim: Positive correlations between tree biodiversity and ecosystem function have been widely

documented, but the nature of the relationship in highly disturbed and ecophysiologically stressful

33 systems is less clear in southern African savanna/woodlands, which experience high levels of dis-

turbance and ecophysiological stress, is less clear. Here, we explore the relationship between tree

species diversity and aboveground biomass across southern African savannas and woodlands, with

respect to gradients in stem density, resource availability and disturbance through fire, to build a

37 general understanding of the biodiversity - ecosystem function relationship in this understudied

38 ecological context.

Location: Southern African savannas and woodlands

40 **Time period:** 2010-2019

41 Major taxa studied: Trees

Methods: We used a network of 1767 savanna/woodland plots located across the southern African

sub-continent in which each tree >10 cm diameter was measured and identified to species level.

We used structural equation modelling and path analysis to determine the relationship between

tree species diversity and aboveground woody biomass, accounting for the interactive effects of re-

source availability and along a gradient of stem density.

Results: A positive effect of tree species diversity on aboveground biomass was demonstrated,

48 observed largely as an indirect effect of increasing woodland structural diversity. We also found

that the effect of tree species diversity on biomass increases with stem density. Finally, we found

that resource availability affects biomass in southern African woodlands largely indirectly via its

51 effect on species diversity.

52 Main conclusions: The study underlines the close association between tree diversity, ecosystem

53 structure and function of highly disturbed southern African savannas and woodlands. Our results

54 demonstrate the importance of including environmental conditions and vegetation type in models

to accurately describe a general relationship between biodiversity and ecosystem function at a re-

gional level. Biodiversity loss predominantly by human actions in southern Africa may have detri-

57 mental outcomes for ecosystem function, particularly in species poor Baikiaea woodlands, which

showed the strongest biodiversity - ecosystem function relationship.

# 4 Introduction

60 Scientific interest in the relationship between species diversity and ecosystem function springs

from both an interest in the factors which structure ecological communities (), and a more ap-

62 plied interest in determining the effect of global biodiversity loss on ecosystem form and func-

63 tion (). Numerous studies have shown relationships between biodiversity and ecosystem function

64 (e.g. Liang et al. 2016; ?; Cardinale et al. 2009). The strength and direction of these observed

65 Biodiversity-Ecosystem Function Relationships (BEFRs) varies depending on the ecosystem be-

66 ing studied, the ecosystem function(s) of interest (Hector and Bagchi, 2007), and the inclusion of

environmental covariates in statistical models (Vilà et al., 2005), but there appears to be a gener-

alisable positive correlation between biodiversity and ecosystem function (Liang et al., 2016). Over

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the past decade, many observational studies of the BEFR have been conducted, mostly in wet
    tropical and temperate forests, and grasslands (Chen et al., 2011). These studies support early
70
    findings from small scale experimental studies predominantly in grassland patches, which began in
71
    earnest during the 1990s as concern grew over the global loss of biodiversity (Tilman and Down-
    ing, 1994; Tilman et al., 2014).
73
    Ecosystem functions can be defined in broad terms as rate processes and properties of ecosystems
    which describe the degree of biotic activity within an ecosystem (Jax, 2005). This includes ba-
75
    sic processes of primary production such as gross primary productivity and atmospheric nitrogen
76
    fixation, but can be extended to indirect measures of function such as resistance of productivity
77
    to disturbance, and further to ecosystem properties which themselves influence process, such as
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    trophic complexity and total vegetative biomass. The frequently reported and intuitive relation-
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    ship between biodiversity and ecosystem function invokes three main mechanisms which drive the
    relationship (Tilman et al., 2014): 1) niche complementarity, whereby communities with greater
81
    biodiversity fill a greater breadth of realised niche space and avoid competition due to differences
82
    in their traits, 2) selection effects, whereby communities with greater biodiversity are more likely
83
    to include a species which contributes highly to the measured ecosystem function, and 3) facili-
84
    tation effects, whereby communities with greater biodiversity are more likely to include combina-
85
    tions of species which together increase the others' functional contribution.
    Compared to other forested forest ecosystems, dry tropical woodlands and savannas are highly
87
    structured by disturbance, mainly through fire and in Africa notably by herbivory also and her-
88
    bivory, with African savannas possessing large herbivores absent from other savannas (Sankaran
    et al., 2008; ?). Disturbance via human activities such as timber extraction and charcoal process-
90
    ing is also common in African woodlands, often causing high levels of disturbance in localised
91
    areas (). High levels of disturbance may weaken the role of competition in determining local
92
    species distribution and allow weak competitors to co-exist where they would normally be ex-
93
    cluded (Grime, 1979; Keddy, 1990). This means that interspecific competition and therefore the
94
    effect of niche complementarity, which contributes the majority of the observed biodiversity effect
95
    on ecosystem function in temperate and wet tropical forests (Wright et al., 2017; Poorter et al.,
96
    2015; van der Sande et al., 2017), may not be as apparent in dry woodland/savanna ecosystems.
97
    Instead, stress tolerance and the functional contribution of more abundant species (selection ef-
98
    fects) may be the predominant forces which influence ecosystem functions (Lasky et al., 2014;
99
    Tobner et al., 2016). Similarly, more diverse species assemblages may lead to facilitation effects
100
    between certain species combinations in environments which are more hostile to growthunder lim-
101
    iting environmental conditions such as low water availability () or high maximum temperature ().
102
    Across European forests Ratcliffe et al. (2017) found stronger positive relationships between tree
103
    species richness and various ecosystem functions in more arid environments. They suggest that in
104
    dry ecosystems, facilitative effects and selection effects may be more important than niche com-
105
    plementarity in driving the relationship between species diversity and ecosystem function. This
106
    potential mismatch in the contribution of different mechanisms to the BEFR between dry tropical
    woodlands and other forested ecosystems demands further investigation in order to characterise a
108
    generalisable BEFR.
109
    The representation of dry tropical ecosystems in the BEFR literature is poor compared to other
    ecosystems. Clarke et al. (2017) conducted a meta-analysis of 182 published BEFR studies, find-
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    ing that only 13% were conducted in the tropics generally, with 42% of those being conducted
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    in the wet tropical forests of Costa Rica, which hold many endemic species and unique ecosys-
113
    tem assemblages (Barthlott et al., 2005). A severe lack of study in dry tropical ecosystems, es-
114
    pecially given the potential mismatch in BEFR mechanism described above, suggests that a fo-
    cus on these ecosystems could greatly strengthen our understanding of a general BEFR and its
    environmental determinants. Savannas and woodlands are the dominant vegetation type across
117
    the southern African region, spanning >4 million km<sup>2</sup> (?Ratnam et al., 2011; Ryan et al., 2016)
118
    (Figure 1). The carbon stored in this vegetation is comparable to that found in the wet forests of
119
    the Congo basin and is of global importance to the carbon cycle (Houghton et al., 2009; Mayaux
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et al., 2008). Climatic conditions and biogeography vary across southern African vegetation, re-
121
    sulting in a diverse range of savanna and woodland tree species assemblages, which retain the
122
    common features of an open tree canopy and an understorey generally dominated by C4 grass
123
    species. Southern African savannas and woodlands (SAWs) are highly diverse, thought to har-
    bour ~8500 plant species of which there are >300 tree species (Frost, 1996), and have been iden-
125
    tified by previous studies as a priority for conservation efforts (Byers, 2001; Mittermeier et al.,
126
    2003). Many conservation projects in the region currently aim to conserve biodiversity and woody
127
    biomass stocks simultaneously under the directive of the United Nations REDD+ programme or
128
    the similar Forest Carbon Partnership Facility (FCPF) (Hinsley et al., 2015). Despite these efforts
129
    however, human actions are driving rapid changes in biodiversity, with largely un-quantified conse-
130
    quences for ecosystem structure and function.
131
    A small number of studies in SAWs, all of which were restricted in the spatial scope to a small re-
132
    gion of miombo woodland, have found that above ground woody carbon/biomass stocks correlate
133
    positively with tree species richness (McNicol et al., 2018; Shirima et al., 2015; Mutowo and Mur-
134
    wira, 2012). The results of these fine scale studies concur with similar studies in other biomes ().
135
    Studies of the BEFR often find that at fine scales, biodiversity shows a strong effect on ecosystem
136
    function, but at broad scales biodiversity effects pale in significance compared to abiotic factors
137
    such as climate (Pasari et al., 2013). Due to the highly variable environmental conditions within
    which SAWs occur (Frost, 1996), with wide variation in precipitation, diurnal and annual temper-
139
    ature range and given the potential importance of environment and biogeography in defining the
140
    strength and form of a relationship between biodiversity and above ground woody biomass (), it is
141
    important to sample across geographic and environmental gradients to gain understanding of the
142
    spatial variation in the relationship between biodiversity and biomass.
143
    In forests, climatic variation is known to affect both woody biomass (Michaletz et al., 2014, 2018)
144
    and tree species diversity independently (Spasojevic et al., 2014). It is important therefore to ac-
145
    count for climatic factors and understand how they interact with biomass and biodiversity to ef-
146
    fectively model and correctly attribute the effects of biodiversity on woody biomass in analyses
    at broad spatial scales. Sankaran et al. (2005) used data from 854 African woodland field sites to
148
    show that below a threshold of ~650 mm MAP, precipitation sets sets the upper limit for woody
149
    cover in savannas, which is positively correlated with biomass (Chisholm et al., 2013; Prado-Junior
150
    et al., 2016). Similarly, Condit et al. (2013) found that dry season intensity was the main determi-
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    nant of tree species distribution and abundance evenness in a wet Panamanian tropical forest.
152
    Solbrig et al. (1996) writes that SAWs possess structurally diverse tree canopies, with trees occu-
153
    pying distinct layers of the canopy at different growth stages and among species. This structural
154
    diversity may be one mechanism through which tree species diversity influences woody biomass.
155
    Kunz et al. (2019) found that crown complementarity and crown plasticity both increased with
    species richness in a seasonally dry subtropical forest. They also found that trees growing in species
157
    rich neighbourhoods exhibited enhanced biomass production. Occupation of multiple canopy lay-
158
    ers allows a more full canopy with a greater total foliage density, enhancing productivity and al-
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    lowing greater standing woody biomass in a smaller area via a form of niche complementarity.
160
    This mechanism however, which has been supported by experiments and observational studies in
161
    temperate and wet tropical ecosystems (Hardiman et al., 2011; Stark et al., 2012), may not be rel-
    evant in savannas, which are structured by disturbance rather than competition. Instead, distur-
163
    bance history may override the effects of tree species diversity on structural diversity nullifying the
164
    effects of tree species diversity on structural diversity.
165
    High levels of disturbance in SAWs may moderate the observable BEFR through its effect on ecosys-
166
    tem composition. Fire disturbance in forests has been linked to abundance dependent mortal-
167
    ity among smaller tree stems (Roques et al., 2001; Staver et al., 2009; Bond and Keeley, 2005).
168
    Some species in the regional species pool may be excluded from woodland plots with high levels
169
    of disturbance if they are unable to escape the fire bottleneck and grow to become a large tree.
    Selection effects may therefore be more important in maximising ecosystem function in distur-
171
    bance prone woodlands. If the regional species pool contains a large number of species, it is more
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likely that one of them will possess the necessary growth strategy to grow to a large tree with high biomass under an intense disturbance regime. 174 In this study, we made the first known regional estimation of the biodiversity-ecosystem func-175 tion relationship across southern African savannas and woodlands (SAWs), using inventory plots 176 which span environmental and biogeographical gradients (Figure 1). We used aboveground woody 177 biomass of trees and compared the relative effects of tree species diversity with that of environ-178 mental factors known to affect ecosystem productivity and biomass accumulation, namely water availability, energy input and soil fertility. We also investigated the potential moderating effects 180 of environmental covariates on the relationship between tree species diversity and biomass. We 181 incorporated vegetation type via clustering of plot level tree species composition, as a factor in 182 our analyses to understand how tree species composition as well as diversity affected ecosystem 183 function and assess the generality of our results. We used Structural Equation Modelling (SEM) 184 and path analysis as a preferred method to simultaneously account for environmental and biotic 185 factors, which may interact their effect which may have interacting effects on ecosystem structure 186 and therefore biomass. Initially, we made three hypotheses: (1) water availability and soil fertil-187 ity will indirectly positively affect woody biomass via an increase in tree species diversity, (2) the 188 effect size the strength of the effect of tree species diversity on woody biomass will increase with 189 plot level stem stocking density, due to an increased importance of niche complementarity as com-190 petition increases, and (3) tree species diversity will increase tree structural diversity, which will 191 provide an indirect path by which tree diversity increases woody biomass. 192

## 5 Materials and methods

# 5.1 Study location

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The study used 1767 woodland monitoring plots from the larger SEOSAW network (SEOSAW, 195 2019) located across 10 countries within southern Africa in the so-called miombo woodland eco-region miombo 196 ecoregion (Figure 1, ?). The study region spans area space the core climate space of the region, 197 with a precipitation gradient from ~460 mm y<sup>-1</sup> in southern Mozambique and southern Zimbabwe 198 to ~1700 mm y<sup>-1</sup> in northern Zambia, Malawi and northern Mozambique. The 2D convex hull of Mean Annual Precipitation (MAP) and Mean Annual Temperature (MAT) of the study sites cov-200 ers 94.4% of the pixel-wise climate space of the miombo woodland ecoregion as defined by (?), 201 using WorldClim estimates of temperature and precipitation between the year 1970 and 2000 with 202 a pixel size of 30 arc seconds (0.86 km<sup>2</sup> at the equator) (Fick and Hijmans, 2017). 203 Plots were chosen from a larger pool of 5395 plots based on the quality and completeness of data 204 collection, and plot setup. Plot vegetation was identified under the broad term of "savanna", which 205 includes "woodland", "savanna woodland", and "tree savanna", variously defined in other areas of 206 the scientific literature and here referred to collectively as southern African woodlands (SAWs) 207 (Ratnam et al., 2011; Hill and Hanan, 2011). Plots with evidence of farming, human resource extraction or experimental treatments such as prescribed burning or herbivore exclusion were ex-209 cluded from the initial pool. Only plots >0.1 hectares were used in analysis, as area based biomass 210 estimation from small plots is highly influenced by rare large trees (Stegen et al., 2011), leading 211 to inaccurate estimates. Only plots with a stem density >10 stems ha<sup>-1</sup> (>10 cm stem diameter) 212 were used, to ensure all plots were within woodland rather than "grassy savanna", which are considered a separate biome with very different species composition (Parr et al., 2014). Many plots provided by the 2005-2008 Zambian Integrated Land Use Assessment (?) were ar-215 ranged in clusters of up to four 20x50 m plots, 20 metres apart. Plots within each cluster were aggregated before the plot dataset filtering described above and treated as a single plot in analy-217 ses. 218 After the initial plot data cleaning described above, we conducted an outlier removal procedure of

plots with rare tree species composition. We used the outlier() function from the dave R pack-

age (Wildi, 2017), which uses a nearest neighbour criterion for each plot in species abundance ordination space and a threshold value for the minimum nearest neighbour distance to identify outliers. We set the threshold value to remove the top 5% of plots with the largest nearest neighbour
distances in multidimensional species composition space (Otto, 2013), thus removing 87 plots (Appendix A).

#### 5.2 Data collection

226

We considered only trees and shrubs in our calculations of above-ground woody biomass (AGB), 227 including woody species such as palms and cycads which are functionally tree-like, but excluding 228 lianas, which fill a different ecological niche (Selaya and Anten, 2008). Only stems >10 cm DBH 229 (Diameter at Breast Height, 1.3 m) were included in analyses. Many plots in the dataset did not include data on stems < 10 cm DBH. For those plots with stem measurements < 10 cm DBH, those 231 small stems only accounted for a median average of 2.2% of the plot level AGB. 232 All stems >10 cm DBH were measured within each plot resulting in a total of 93,242 stems with measurements. A tree may be comprised of multiple stems, but for this analysis each stem is treated 234 as an individual. For each stem we measured species, DBH and tree height to the top of the high-235 est branch material. Height was measured through a variety of means including laser rangefind-236 ers, manual clinometers and measuring sticks. When DBH could not be measured at 1.3 m due 237 to trunk abnormalities, it was measured at the closest regular portion of the trunk to 1.3 m. The 238 height of this measurement was recorded and used to estimate the DBH<sub>e</sub> at 1.3 m using a cubic polynomial regression, with parameters estimated using a test dataset from (Ryan C., unpub-240 lished) (Appendix B). 241 AGB for each plot was calculated using Equation 1, taken from Chave et al. (2014). Wood density estimates were taken from the global wood density database for each species where possible 243 (Chave et al., 2009; Zanne et al., 2009). Wood density for species without species level estimates 244 was estimated from the mean of their respective genus. 245

$$AGB = 0.0673 \times (\rho D^2 H)^{0.976} \tag{1}$$

Where  $\rho$  is the species level mean wood density, D is the DBH<sub>e</sub> at 1.3 m, and H is the tree height. 246 Climatic data were collected from the ECMWF ERA5 dataset, generated using Copernicus Cli-247 mate Change Service Information (Copernicus Climate Change Service, 2017). Values of Mean 248 Annual Temperature (MAT)MAT and Mean Annual Precipitation (MAP)MAP were calculated 249 from daily data between 2000 and 2018, then averaged across years to provide a single mean an-250 nual estimate per plot. Temperature seasonality (TS) and precipitation seasonality (PS) were both 251 calculated as the mean of the coefficient of variation of daily mean temperature and precipitation, 252 respectively, for each of the 18 years of available data. Soil fertility data was extracted from the 253 ISRIC gridded soil information data product at 250 m resolution, taking the grid cell value for 254 each plot centre (Hengl et al., 2017). We extracted Cation Exchange Capacity (CEC), percentage 255 soil organic carbon by volume (Org. C%), and percentage soil sand content by volume (Sand %). These data are a modelled product derived from various remotely sensed and directly measured 257 data sources. 258

#### 259 5.3 Data analysis

# 260 5.3.1 Species diversity and structural diversity metrics

Estimated tree species richness was calculated for each plot using ChaoRichness() from the iNEXT package in R (Hsieh et al., 2016). This procedure extrapolates a species rarefaction curve to its predicted asymptote and uses this value as its estimated species richness value. Extrapolated species richness accounts for variation in plot size (0.1-10 ha) and therefore sampling effort among plots.

Larger plots will tend to encompass more individuals, and therefore more species (Dengler, 2009).

To measure tree species abundance evenness, the Shannon Equitability index  $(E_{H'})$  (Smith and

Wilson, 1996) (??) was calculated as the ratio of the estimated Shannon diversity index to the

natural log of estimated species richness. Abundance evenness allows for greater niche complementarity at small scales due to an increased spatial heterogeneity of functional traits. We calculated

tree structural diversity for each plot by calculating the coefficient of variation of DBH (DBH CV)

and tree height (Height CV).

#### 5.3.2 Vegetation clusters

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Plots were assigned to vegetation type groups based on tree species composition. Groups were defined in Fayolle et al. (2018) in an Africa wide analysis of floristic units using plot data in savannas and woodlands with tree species diversity and relative abundance data. Group identification was conduscted using unconstrained correspondence analysis and ordination, followed by clustering based on dominant ordination axes. Plot data used in this study occurred in four vegetation type groups. See Table 1 for a description of each vegetation cluster and Figure 1 for the spatial distribution of plots from each of these clusters.

Table 1: Description of the biogeographical clusters (C1-C5) to which each plot in the study was assigned. Indicator species were generated using Dufrene-Legendre indicator species analysis (Dufrêne and Legendre, 1997) implemented with indval() from the labdsv R package (Roberts, 2019) and represent species which define the given cluster. Dominant species were identified by choosing the species with the largest AGB contribution within each cluster. Numeric values of species richness, stems ha<sup>-1</sup> and AGB are medians and interquartile ranges (75th percentile - 25th percentile).

Cluster	Dominant species	Indicator species	N plots	Species rich.	Stems ha <sup>-1</sup>	AGB (t ha <sup>-1</sup> )
Marginal miombo	$Julbernadia \ { m spp}.$	$Diplor hynchus\ condylocarpon$				
	$Brachystegia\ spic iform is$	is Burkea africana		11(11.2)	152(135.2)	32.9(30.57)
	$Baikeaea\ plurijuga$	$Pseudola chnostylis\ maproune ifolia$				
Core miombo	Julbernadia spp.	Julbernardia paniculata				
	Brachystegia  spp.	stegia spp. Isoberlinia angolensis		18(17.5)	193(174.1)	44.8(41.43)
	$Is oberlinia\ angolensis$	$Brachystegia\ longifolia$	chystegia longifolia			
Baikiaea	Spirostachys africana	Baikiaea plurijuga			162(152)	
	Senegalia  spp.	$Senegalia\ ataxacantha$	226	10(10)		45.3(47.36)
	$Euclea\ racemosa$	$Combretum\ collinum$				
Mopane	Colophospermum mopane	Colophospermum mopane	00	7(9.9)	190(155.7)	41.5(36.93)
		$Combretum \ \mathrm{spp.}$	99 7(8.2)		190(199.7)	41.0(30.93)

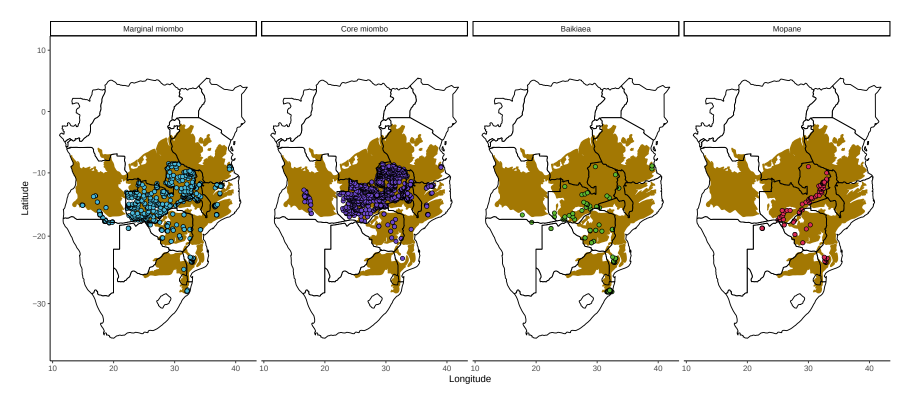


Figure 1: The locations of the 1767 plots used in this study, as points by geographic location with respect to the distribution of miombo woodland vegetation according to ?. Each panel shows plots categorized by their vegetation type as defined by the vegetation types in Table 1.

## 5.3.3 Structural Equation Modelling

Structural Equation Models (SEM) investigated the determinants of AGB. All SEMs were constructed and analysed in the lavaan package (Rosseel, 2012) in R version 3.6.0 (R Core Team, 2019). SEM was used because of its suitability for modelling complex causal interactions in ecological systems (Lee, 2007). A key aspect to our decision to use SEMs is that they can explicitly model and partition variance to indirect effects, which is challenging in standard multiple regressions. Using SEMs also allowed us to describe theoretical latent constructs which have been suggested to act upon biodiversity and biomass/productivity in previous studies despite these factors not having single observable measures in our dataset. Structural equation modelling is also necessary to properly account for potential feedback mechanisms between aspects of climate and tree species diversity, which could otherwise increase the chances of Type I error and wrongly attribute inference due to covariance of explanatory variables when using conventional regression analyses (Nachtigall et al., 2003).

Prior to analysis, we specified a conceptual model with factors expected to affect AGB: moisture availability, soil fertility, tree species diversity, tree structural diversity and stem density (Figure 2).

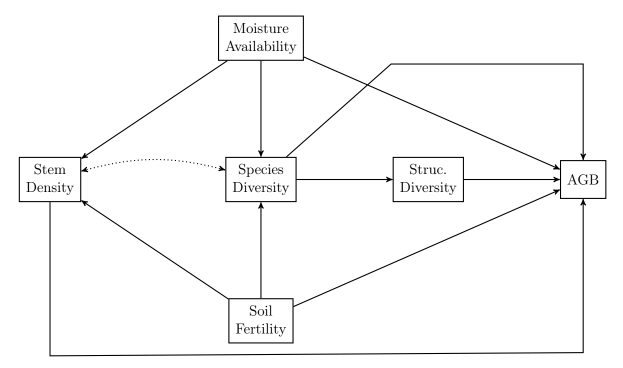


Figure 2: Conceptual Directed Acyclic Graph (DAG) showing the theoretical relationships between environmental factors, tree species diversity, tree structural diversity, tree stem density, and AGB. Hypothesised paths of causation are depicted as arrows from predictor to response. Correlations are depicted as curved dotted arrows.

Observed variables were transformed to achieve normality where necessary and standardised to Z-scores prior to analysis (Appendix C). Standardisation put each latent variable on the same scale, with a mean of zero and a standard deviation of one. Standardisation allows path regression coefficients to be easily compared between paths in the same model to assess their relative effect size, and eliminates confusion in model interpretation arising from the observed variables being on different scales (Beaujean, 2014). Standardisation also controls for variables with different orders of magnitude which could otherwise prevent adequate model estimation from the covariance matrix in lavaan. To ensure that observed variables within a latent variable had consistent directions of influence, some observed variables were reversed by multiplying by -1. For example, soil fertility is expected to decrease as soil sand content increases, so soil percentage sand content was reversed

for model fitting. Precipitation seasonality (PS), temperature seasonality (TS), and mean annual temperature (MAT) were also reversed in this way to account for the direction of their effect on moisture availability.

The factor loadings of the observed variable assumed to contribute most to each latent variable were set to 1 as per convention, with other observed variables being allowed to vary (Beaujean, 2014). We tested the robustness of our assumptions with a chi-squared test of all possible combinations of observed variable factor loadings set to 1, while ensuring no factor loadings were in excess of 1. We found no significant difference between model specifications (p>0.05). Full Information Max-Likelihood (FIML) was used in each model to estimate the values of missing data in each latent variable (Cham et al., 2017).

We assessed the role of tree structural diversity and species diversity in determining AGB via a 316 simple mediation model which allowed species diversity to influence AGB both directly and in-317 directly via structural diversity. To account for variation in stem density which may covary with 318 species diversity we also included it as an predictor in our model. To explore variation in the model 319 among woodland vegetation types, we fit the model both at the regional scale and for each vegetation cluster separately. We compared unstandardised path coefficients among these vegetation 321 cluster scale models to understand the effect that vegetation type has on the relationship between 322 tree species diversity, structural diversity, stem density and AGB. Path coefficients show the effect 323 of a path with other paths of inference held constant. Models were estimated using the "MLM" 324 estimator, because it is robust to multivariate non-normality (Shapiro, 1983). Model fit was eval-325 uated using the robust Comparative Fit Index (CFI), the robust Tucker Lewis Index (TLI), the 326 Root Mean Squared Error (RMSEA) and the R<sup>2</sup> coefficient of determination for AGB. We crit-327 ically assess model fit in each case, taking into consideration the recommendations of Hu and 328 Bentler (1999) which define threshold values of acceptability for these model fit indices: CFI >0.85, 329 TLI > 0.85, RMSEA < 0.15, alongside our judgement of the model estimates. 330

To explore the hypothesis that complementarity effects increase in strength as stem density increases, we repeatedly sub-sampled the available plot dataset to create 50 datasets of similar size with varying median stem density. We used each of these datasets separately to fit the model including only tree species and structural diversity latent variables to predict AGB. We excluded the effect of stem density on AGB and the correlation between stem density and species diversity from this model as we were deliberately controlling stem density in our subsampling. We then examined how the unstandardised path coefficients for each path in the SEM varied according to the median stem density of subsampled dataset.

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We incorporated environmental covariates into our model to understand the relative effects of moisture availability and soil fertility on AGB both directly and indirectly via species diversity and stem density. We compared standardised path coefficients between paths in the model to understand the relative contribution of each path to explain variance in AGB. Vegetation type specific models could not be reliably fitted for this more complex model specification with environmental covariates, due to sample size issues and because some vegetation clusters were narrow in their climate space leading to a lack of variance particularly in moisture availability.

# 346 6 Results

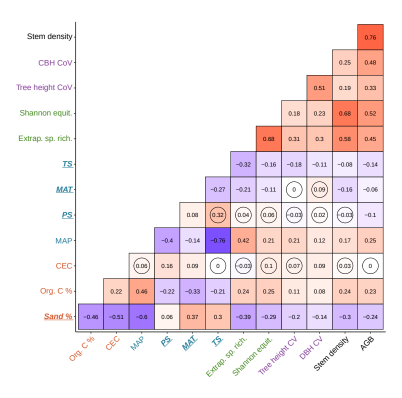


Figure 3: Correlogram of standardised observed variables used in the SEMs, with Pearson correlation coefficients (r) coloured according to sign (+ve red, -ve blue) and shaded by strength of correlation. Variables in bold and underlined on the axis labels were later reversed for SEMs to maintain positive correlations for all observed variables within each latent variable. Correlation coefficients marked by a circle indicate that the 95% confidence interval of this correlation overlapped zero. Colours of variable names group them into latent variables used in the SEMs: red = soil fertility, blue = moisture availability, green = tree species diversity, purple = tree structural diversity. See Appendix D for a full assessment of correlation fit statistics.

Pairwise correlations between all observed variables used in the Structural Equation Models (SEMs) 347 showed that all tree species diversity and structural diversity variables had moderate positive correlations with AGB. Stem density had the strongest correlation with AGB of all variables (r =0.77, p <0.01). Environmental variables had weaker correlations with AGB than diversity vari-350 ables, with all environmental variables having significant correlations with AGB, except CEC and 351 MAT. 352 The direction of these correlations was used as a test of our assumptions of the direction of in-353 fluence of latent variables later used in the SEMs. As expected, there was a positive correlation 354 between MAP and AGB (r = 0.22, p < 0.01), and a weak negative correlation between the sea-355 sonality of precipitation and AGB (r = -0.11, p < 0.01). MAT and temperature seasonality (TS) 356 negatively correlated weakly with AGB (MAT: r = -0.09, p < 0.01; TS: r = -0.14, p < 0.01). As ex-357 pected, there was a negative correlation between soil sand content and AGB (r = -0.27, p < 0.01), 358 and a positive correlation between soil organic carbon and AGB (r = 0.25, p < 0.01). 359 MAP had positive correlations with tree species richness (r = 0.39, p < 0.01), abundance evenness (r = 0.1, p < 0.01), tree height diversity (r = 0.21, p < 0.01) and tree stem density (r = 0.09, p < 0.01)361 p < 0.01). MAT had weak correlations with tree species and structural diversity variables. Tree 362 species diversity variables had clear positive correlations with stem density (Species richness: r =363 0.57, p <0.01; Shannon equitability: r = 0.48, p <0.01). 364

# 6.1 Structural and species diversity models

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In an SEM describing the effect of tree species diversity on AGB via the mediating effects of stand structural diversity and stem density (Figure 4), species diversity had a small positive direct effect on AGB ( $\beta = 0.11\pm0.039$ , p <0.01), and indirectly via structural diversity ( $\beta = 0.15\pm0.023$ , p <0.01) (Figure 4). Tree species diversity had a positive correlation with stem density. Model fit was good with high factor loadings for all observed variables, all path coefficients were significant (p <0.01) (Table 2). The R<sup>2</sup> of AGB was 0.69. The strongest direct effect on AGB was from stem density ( $\beta = 0.74\pm0.033$ , p <0.01).

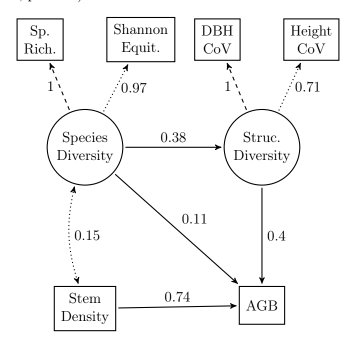


Figure 4: Path diagram with regression coefficients for the tree diversity SEM, including plots from all vegetation clusters. Latent variables are circles while observed variables are rectangles. Standardised path coefficients are solid arrows pointing from predictor to response with the effect size of the path coefficient expressed in terms of standard deviations on the latent variable response scale. The observed variables which inform the latent variables are connected by dotted arrows, observed variables with loading set to 1 are connected by dashed arrows. Correlations between variables are depicted as dotted curved arrows. Measurement errors of exogenous variables are omitted for clarity.

#### 6.2 Variation among vegetation types

When the tree species and structural diversity model (Figure 4) was refitted separately using data 374 from each of the 4 vegetation types the strengths of unstandardised path coefficients varied. The 375 direct effect of tree species diversity on AGB was positive in Baikiaea and Mopane, but negative 376 in Marginal and Core miombo (Figure 5). Relationships between structural diversity and AGB re-377 mained generally similar with the same sign and significant overlap between the 95% confidence 378 intervals of path coefficients. The total effect of species diversity on AGB remained strongly pos-379 itive for all vegetation types. All vegetation types except Mopane exhibited a positive effect of 380 species diversity on structural diversity. All models had adequate goodness-of-fit (Table 2), though 381 confidence intervals around the unstandardised path coefficients were wide particularly for Mopane 382 and Baikiaea.  $\chi^2$  statistics were high for some vegetation types, but this appears to be highly cor-383 related with sample size for each vegetation type (Hooper et al., 2008). 384 The strongest total effect of tree species diversity on AGB was in Baikiaea woodland ( $\beta = 0.19 \pm 0.154$ , 385 386

p = 0.22), which was species rich but highly variable in species diversity compared to other vegetation types (Table 1). The  $R^2$  of AGB was highest in Marginal miombo ( $R^2 = 0.71$ ) and lowest

in the Core miombo ( $R^2 = \frac{1}{389}$  strucbrsqstrucbrsq).

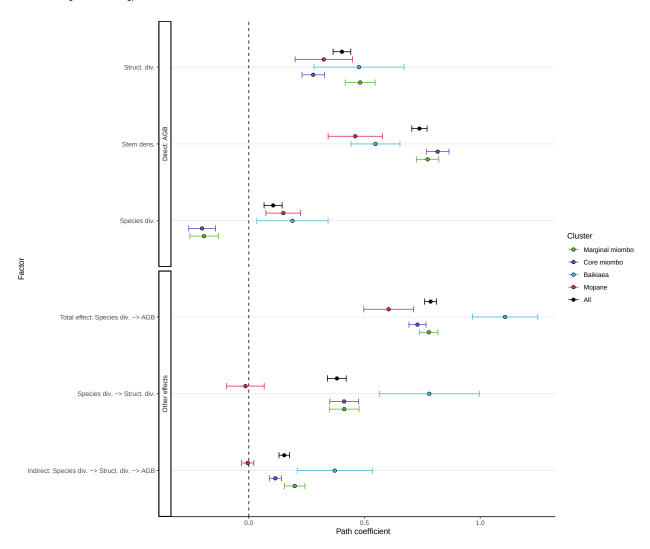


Figure 5: Unstandardised path coefficients for the effects of tree diversity on AGB, mediated by the effect of stand structural diversity. Path coefficients are  $\pm 1$  standard error. Path coefficients where the standard error does not overlap zero are considered to be significant effects.

Table 2: Model fit statistics for SEMs investigating the effects of tree diversity and stem density on AGB (Figure 4).

Cluster	n	$\chi^2$	DoF	CFI	TLI	LogLik	RMSEA	$R^2$ AGB
Marginal miombo	525	44.750	6	0.966	0.916	-3714.000	0.110	0.710
Core miombo	668	57.210	6	0.962	0.904	-4224.000	0.100	0.680
Baikiaea	47	5.860	6	0.998	0.994	-324.600	0.030	0.720
Mopane	84	9.420	6	0.971	0.927	-591.600	0.080	0.450
All	1324	78.430	6	0.975	0.936	-9119.000	0.090	0.690

# 6.3 Moderation of Diversity-AGB relationship by stem density

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We repeatedly sub-sampled the plot dataset to build 50 datasets of varying mean stem density in order to test how the relationship between species diversity, structural diversity and biomass

varied with stem density. Each dataset consisted of approximately 892 plots with overlap of plot identity between subsampled datasets. Figure 6 shows a positive effect of tree species diversity on AGB as stem density increases. In our sub-sampling of the plot dataset by mean stem density, we found a positive effect of tree species diversity on AGB as stem density increases (Figure 6). There appears to be a minimum stem density threshold at ~180 stems ha<sup>-1</sup> below which there appears to be a reasonably constant low baseline effect of tree diversity on biomass. The effect of structural diversity on AGB appears to remain constant with increasing stem density. The indirect effect of species diversity on AGB via structural diversity climbs slightly as stem density increases.

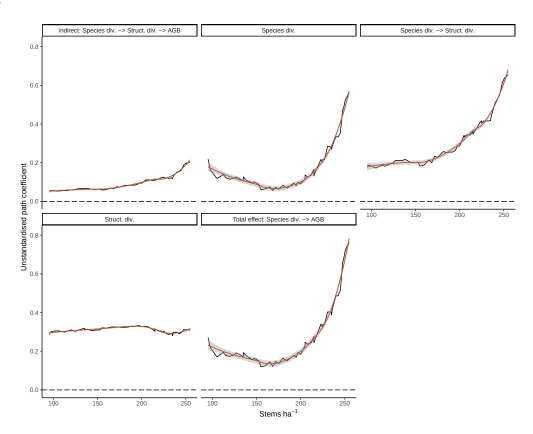


Figure 6: Line plots showing the variation in path coefficients in the SEM, using datasets with different mean stem density. Smoothed lines are loss curves with standard error shaded bars.

## 6.4 Environmental covariates and tree diversity

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A model incorporating the latent variables of moisture availability and soil fertility showed that the total effect of tree species diversity on biomass was greater than that of both moisture avail-404 ability and soil fertility (Figure 7). Surprisingly, the direct effects of moisture availability and soil 405 fertility on biomass were negligible, with nearly all of their observed effect on AGB coming from 406 the indirect path via species diversity (moisture:  $\beta = -0.01 \pm 0.005$ , p <0.01, soil:  $\beta = -0.04 \pm 0.012$ , 407 p <0.01). MAP and temperature seasonality (TS) had the greatest contributions to the latent 408 variable of moisture availability. Moisture availability and soil fertility also had negligible direct ef-409 fects on stem density. Model fit was acceptable: CFI = 0.924, TLI = 0.905, and RMSEA = 0.163, 410  $R^2 \text{ of AGB} = 0.7.$ 411 Similar to the model which only considered tree species and structural diversity (Figure 4), the 412 direct effect of species diversity on structural diversity was positive, while structural diversity itself had a positive effect on AGB, leading to a strong positive indirect effect of species diversity on 414 AGB via structural diversity ( $\beta = 0.16\pm0.023$ , p <0.01). The total effect of species diversity on 415 AGB was positive ( $\beta = 0.64 \pm 0.041$ , p < 0.01). 416

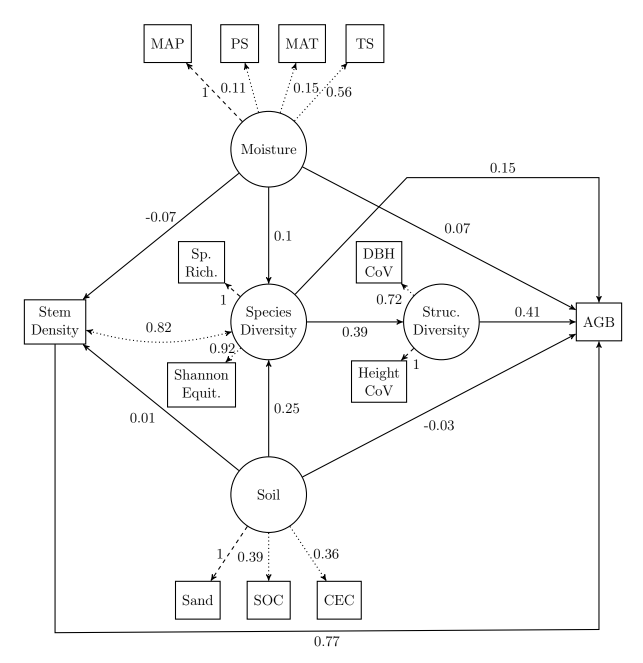


Figure 7: Path diagram with regression coefficients for the SEM incorporating environmental covariates and tree species and structural diversity across all five vegetation types. Latent variables are shown as circles while observed variables are shown as rectangles. Standardised path coefficients are solid arrows pointing from predictor to response with the effect size of the path coefficient expressed in terms of standard deviations on the latent variable response scale. Observed variables which inform the latent variables are connected by dotted arrows, observed variables with loading set to 1 are connected by dashed arrows. Correlations between variables are depicted as dotted curved arrows. Measurement errors of exogenous variables are omitted for clarity.

# <sup>417</sup> 7 Discussion

In this study, we assessed the importance of [a] tree species diversity, [b] tree structural diversity, [c] resource availability, i.e. moisture availability and soil fertility, [d] stem density and their interactions on above ground woody biomass (AGB) across SAWs, using a network of 1767 woodland survey plots. Using latent variables and Structural Equation Modelling (SEM), we found support for a general positive relationship between tree species diversity and AGB, with an indirect influence of tree species diversity on AGB via structural diversity ( $H_1$ ). We found that the effect size

of tree species diversity on AGB increased with stem density (H<sub>2</sub>). Tree diversity, structural diversity and stem density accounted for 69% of the variation in AGB across the region, while models for specific vegetation types showed even greater explanatory power in some cases (Table 2). The strongest effect on AGB was that of stem density. Interestingly, when the effects of tree species diversity, structural diversity and stem density were controlled for, we found little evidence of a direct effect of resource availability, in the form of moisture or soil fertility, on AGB (H<sub>3</sub>).

#### 7.1 Inter-related effects of tree species and structural diversity on AGB

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We found a consistent positive effect of tree species diversity on AGB across all models in this 431 study. Within SAWs we therefore find support that higher tree species richness and evenness causes 432 higher woody AGB. This finding is in agreement with many other studies across different ecosystems and biomes, supporting that there is a generalisable positive association between biodiversity 434 and ecosystem function (Liang et al., 2016; Cardinale et al., 2009). Our study provides a novel 435 dissection of the mechanisms underlying this relationship, particularly in the context of SAWs, a 436 disturbance-structured and poorly studied ecological system. 437 Much of the total variation in AGB was driven by variation in stem density. Stem density also 438 correlated with species diversity in our SEMs. It is possible that within SAWs a higher species 439 diversity allows for a greater density of tree stems, leading to an increase in total AGB. The oppo-440 site is also plausible however, with increased stem density causing higher species richness through 441 an increased probability of encountering new species. We suggest that an increase in tree species diversity through species richness and evenness produces an assemblage of species which can oc-443 cupy a greater proportion of the total woodland canopy volume with leaf area, utilising more of 444 the available light resulting in greater total AGB at the plot level (). This is supported by the 445 moderately strong indirect positive effect of tree species diversity on AGB via structural diversity. 446 We found evidence that tree species diversity led to an increase in AGB indirectly via tree struc-447 tural diversity and we therefore find support for our hypothesis (H<sub>2</sub>). A higher tree species di-448 versity allows for a greater structural diversity of trees, i.e. greater variation in DBH and height. 449 This may act as a mechanism of niche complementarity, with a highly diverse canopy being able 450 to take advantage of a greater proportion of the available light. Although we did not measure it here, we would also expect that tree species diversity allows for a greater range of tree func-452 tional forms (), i.e. wider variation in canopy shape and overall growth form; bushy understorey 453 vs. emergent canopy, for example. Variation in structural diversity may be a joint result of dis-454 turbance history and tree species diversity, with highly disturbed plots generally having a less 455 structurally diverse canopy (LaRue et al., 2019). In forests, where the tree canopy is effectively closed, as the stand matures a more diverse canopy emerges via competition and tree mortality 457 events which open canopy gaps (Muscolo et al., 2014). Indeed, our finding that the strength of 458 the effect of tree diversity on AGB increases with stem density supports this. In frequently dis-459 turbed woodlands such as those studied here however, a woodland canopy similar to that of a for-460 est is frequently not reached, Instead, a simple open canopy is maintained that can be made more 461 complex and productive via an increase in species diversity. While we did not have access to ade-462 quate data on disturbance history in our plots, previous studies have found that SAWs with higher species diversity tend to be less disturbed and tend to form a more closed canopy (Chidumayo, 464 2013: Mutowo and Murwira, 2012). 465 We found a non linear positive effect of stem density on the relationship between tree species diversity and AGB (Figure 6). At low stem densities competition between trees may not occur, 467 meaning that the niche complementarity provided by an increase in tree species richness might not 468 make any difference to plot level AGB, accounting for the low and constant effect of tree species 469 diversity on AGB below ~180 stems ha<sup>-1</sup>.

## 7.2 Effects of moisture availability and soil fertility

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Surprisingly, moisture availability and soil fertility had only small effects on AGB compared to 472 that of tree species diversity. We expected that higher moisture availability and soil fertility would lead to higher AGB under the assumption that higher resource availability would allow for a greater stem density per unit area, greater productivity per unit area and additionally greater tree species 475 diversity due to niche partitioning (Kraaij and Ward, 2006; Shirima et al., 2015). 476 Previous studies in tropical forests have shown that moisture availability increases AGB both directly and indirectly via increasing tree species diversity and via increasing stand structural 478 diversity (Ali et al., 2019a,b; Poorter et al., 2017). In this study, while we observed weak indi-479 rect effects via species diversity, we saw no evidence for a direct effect of moisture availability on 480 AGB. Compared to moist tropical forests, moisture availability is more of a limiting factor to tree 481 growth in SAWs, which are frequently droughted. It is possible that the range of observed mois-482 ture availability in this study (~460-1700 mm y<sup>-1</sup>) may not have been able to capture variation 483 in AGB. Due to the high levels of adaptation of tree species to drought conditions in southern 484 Africa, at the large scale we conducted our experiment turnover in species composition along the 485 moisture gradient may have obscured a direct relationship being observed between moisture avail-486 ability and AGB. 487 In SAWs moisture availability is closely linked with the intensity of disturbance from seasonal 488 fires. The growth of C4 grasses in wetter woodlands leads to more intense seasonal fires which 489 limit tree growth (Charles-Dominique et al., 2018), and may also limit species diversity (Linder, 490 2014). It is possible therefore that the effect of moisture availability, which is expected to increase 491 AGB, is confounded in its effect on AGB with the unmeasured variable of fire regime intensity, 492 which is expected to decrease AGB. The direct effect of moisture availability on stem density may 493 also be confounded in this way. This may also have caused us to not observe a stronger effect be-494 tween moisture availability and AGB. 495 We expected a positive effect of soil fertility on AGB, but found no evidence of this in our models. 496 We measured soil fertility using the observed variables of soil organic carbon content, sand parti-497 cle content and Cation Exchange Capacity (CEC). In wet tropical forests a clear relationship has 498

#### 7.3 Vegetation type specific responses

been observed between these variables and AGB (??).

Core miombo and marginal miombo woodland vegetation exhibited a small negative direct effect 501 of tree species diversity on AGB, while the total effect, incorporating the indirect effect via struc-502 tural diversity, remained positive in these vegetation types. Compared to Baikiaea and Mopane 503 woodlands, miombo woodlands have higher median tree species richness. Baikiaea and Mopane 504 woodlands are also dominated by fewer tree species, notably Baikiaea plurijuga in Baikiaea wood-505 lands and Colophospermum mopane in Mopane woodlands which often produce large canopy dom-506 inating trees. We postulate that this negative effect of tree species richness on AGB in miombo 507 woodlands may be due to an increase in interspecific competition through canopy crowding, but 508 that this effect is not present in Baikiaea and Mopane woodlands, where the woodland canopy is 509 dominated often by a single species. Higher functional redundancy among tree species in miombo 510 woodlands may lead to smaller trees with lower AGB in the most diverse plots, more resembling 511 thicket vegetation. Again, these highly diverse plots in miombo woodlands may be the result of 512 disturbance which can promote a mosaic of woodland of different successional stages and stem densities. Alternatively, this small negative direct effect may be an artefact of particularly noisy 514 data, especially given that the overall effect of tree diversity on AGB is positive. 515 Despite Mopane woodland having very low species diversity generally, with often monospecific stands (Timberlake et al., 2010), a positive effect of tree species diversity on AGB was observed. 517 In previous studies across ecosystem types it has been found often that the effect on ecosystem 518 function of adding species is stronger in low diversity assemblages (Hector and Bagchi, 2007). This

has been attributed to an increase in functional redundancy as species diversity increases. I.e. 520 with more species, it is more likely that the addition of a new species will occupy the same ecolog-521 ical niche space as an existing species, meaning niche complementarity will not occur and competi-522 tion will lead to niche partitioning, while making little difference to overall ecosystem functioning. Mopane woodlands also have a negligible effect of species diversity on structural diversity. This 524 may be due to the species which tend to co-exist with C. mopane, many of which are small shrub-525 like trees which do not grow into large canopy trees (Timberlake et al., 2010). Larger canopy trees 526 tend to have greater variation in physical structure (Seidel et al., 2019). 527 Baikiaea woodland had the strongest total effect of species diversity on AGB. Baikiaea also has 528 relatively low median species richness compared to miombo, but the addition of new species ap-529 pears to make a larger difference to the AGB of these plots than in mopane woodlands. We sug-530 gest that this is due mostly to the particular identity of species found in Baikiaea woodlands and 531 their contribution to ecosystem functioning. Unlike mopane woodlands, Baikiaea woodlands do sometimes contain species other than B. plurijuqa which grow to be high biomass canopy trees.

#### 534 7.4 Conclusion

In this study we found that across southern African woodlands (SAWs), there is a generalisable 535 positive association between tree species diversity and woody biomass as a measure of ecosystem 536 function. Additionally, we found that much of this effect of species diversity on biomass exists as 537 an indirect effect by increasing the structural diversity of woodland tree canopies. We found that the multiple vegetation types which comprise SAWs exhibit variation in the strength of the rela-539 tionship between species diversity and woody biomass, inferring that models of regional and global 540 biodiversity-ecosystem function relationships could benefit from including vegetation type terms 541 and the structural properties of those vegetation types, such as structural diversity and stem den-542 sity. In contrast to previous studies, we found that across the region, the direct effects of mois-543 ture availability and soil fertility on woody biomass were negligible, with most of their effect being indirectly through species and structural diversity. A gap in available data means that we could 545 not incorporate disturbance history into our models adequately, but this factor likely plays a large 546 part in the association between species diversity and woody biomass in SAWs. 547 SAWs are relied heavily upon for their ecosystem service provision, which is itself affect by ecosys-548 tem function. Resource extraction by humans in southern Africa is directly influencing biodiver-549 sity via selective tree-felling for timber, among other forest products. Our study shows that biodi-550 versity change through human actions will have the greatest negative impact on ecosystem func-551 tion in areas of high stem density and Baikiaea woodlands, which are predominantly targeted for tree felling. This raises concerns about the robustness of these ecosystems to further resource ex-553 traction and biodiversity loss.

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- 818 A Data accessibility statement
- 819 B Tables
- 820 C Figure legends and embedded figures
- B21 D Appendix 1 Data cleaning process

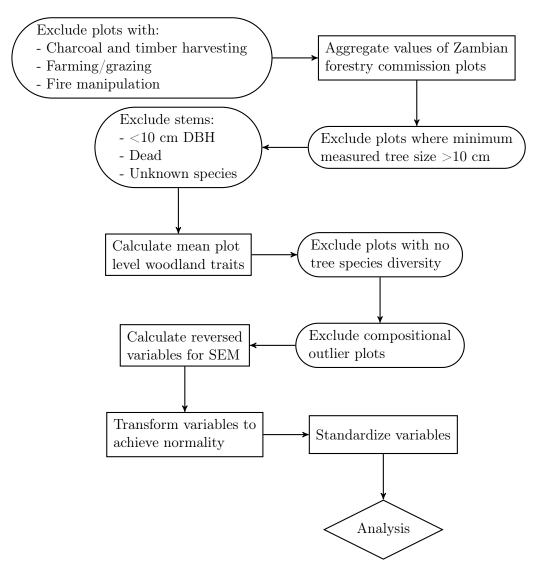


Figure 8: Flow diagram of the data filtering and cleaning process prior to analysis. Rounded boxes indicate filtering events while regular boxes indicate calculation events.

# $_{822}$ E Appendix 2 - Estimation of DBH via tree taper

```
##' @title Stem diameter Point Of Measurement (POM) adjustment

##' @description Function to estimate stem diameter at 1.3 given measurements

##' at other POMs.

##' @author Casey M. Ryan

##' @return d130, the estimated diameter at a POM of 1.3 m (in cm).

##' @param d_in the diameter measured at the POM (in cm)

##' @param POM the height of the POM (in m)

##' @details The adjustment is based on a tree taper model developed as part of
```

```
8329 ##'
           the ACES project (Abrupt Changes in Ecosystem Services
833.0
          https://miomboaces.wordpress.com/), using data from the miombo of Niassa.
    ##,
834.1
          The model is a cubic polynomial, with three equations for different sized
835 2 ## '
8363 ##' @section Warning: The model should not be used for POMs above 1.7 m.
837.4 ##'
         Extrapolating beyond the training data will give nonsense.
   ##'
         Thus, POMs >1.7 m are not adjusted.
838.5
    ##' @examples
839.6
    ##' POMadj(10, 0.3)
840 7
    ##' POMadj(1, 0.3) # d130 is negative, i.e. the stem probably wasn't 1.3 m tall
841/8
    \#\#, POMadj(50, 1.9) \# generates warning, as outside calibration data range
842.9
    ##' \dontrun{
84320
    ##'
          POMadj(50, 0) # zero or -ve POM is outside range, or nonsense
84421
8452
    ##' }
8463
    POMadj <- function(d_in, POM) {</pre>
      stopifnot(is.numeric(d_in),
848
        is.numeric(POM),
        POM >= 0,
849
        sum(is.na(POM)) == 0,
850
        length(POM) == length(d_in))
85128
      if (any(POM > 1.7))
8529
        warning("POMs >1.7 m are outside the calibration data, no correction applied")
85330
854
      NAS <- is.na(d_in)
85532
      d_in_clean <- d_in[!NAS]</pre>
8563
      POM_clean <- POM[!NAS]
8573
      # define the size class edges:
      edges \leftarrow c(5.0, 15.8, 26.6, 37.4)
8598
86087
      sm <- d_in_clean < edges[2]
      med <- d_in_clean >= edges[2] & d_in_clean < edges[3]</pre>
8618
8628
      lg <- d_in_clean >= edges[3]
863
      # compute apredictions for delta_d, for all size classes
864
      delta_d <- data.frame(</pre>
865
        # if small:
866
        small = 3.4678 + -5.2428 *
867
          POM_clean + 2.9401 *
868
          POM_clean^2+-0.7141 *
869
          POM_clean^3,
870
        # if med
871/8
        med = 4.918 + -8.819 *
872
          POM_clean + 6.367
873
          POM_clean^2+-1.871 *
874
          POM_clean^3,
875 2
        # if large
876
        large = 9.474 + -18.257 *
87754
          POM_clean + 12.873 *
          POM_clean^2+-3.325 *
8796
          POM_clean^3
8807
      )
88158
      # index into the right size class
88259
      dd <- NA_real_
88360
      dd[sm] <- delta_d$small[sm]</pre>
88461
      dd[med] <- delta_d$med[med]</pre>
885
      dd[lg] <- delta_d$large[lg]</pre>
8863
      dd[POM\_clean > 1.7] \leftarrow 0 # to avoid extrapolation mess
88764
      # add NAs back in
      d130 <- NA
      d130[NAS] <- NA
      d130[!NAS] <- d_in_clean - dd
8926
8937
      if (any(d130[!NAS] < 0))</pre>
894
        warning("Negative d130 estimated, repaced with NA")
89572
      d130[d130 <= 0 & !is.na(d130)] <- NA
89673
```

8974 return(d130)
88875 }

# <sub>900</sub> F Appendix 3 - Frequency distribution of observed variables

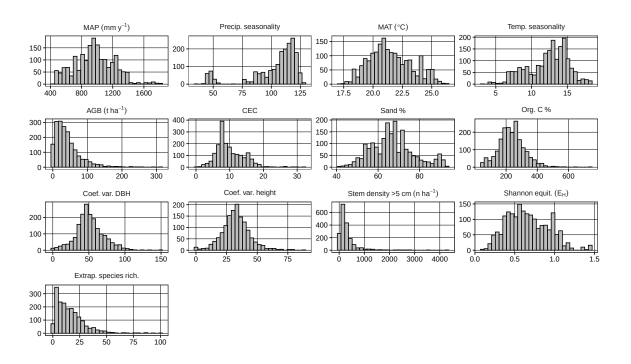


Figure 9: Histograms of raw untransformed observed variables used in final analyses.

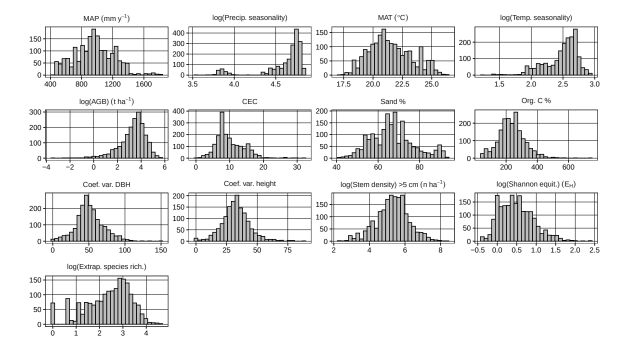


Figure 10: Histograms of observed variables transformed to achieve a normal frequency distribution.

# 901 G Appendix 4 - Table of correlation fit statistics

Table 3: Table of correlation fit statistics for each pairwise Pearson correlation test of observed variables used in Structural Equation Models.

	1					
X	Y	r	lower 95% CI	upper $95\%$ CI	n	Prob.
Sand $\%$	Org. C $\%$	-0.510	-0.540	-0.470	1767	p < 0.01
Sand %	CEC	-0.560	-0.600	-0.530	1767	p < 0.01
Sand %	MAP	-0.500	-0.530	-0.460	1767	p < 0.01
Sand %	PS	0.320	0.280	0.360	1767	p < 0.01
Sand $\%$	MAT	0.290	0.240	0.330	1767	p < 0.01
Sand $\%$	TS	0.230	0.180	0.270	1767	p < 0.01
Sand $\%$	Sp. rich.	-0.350	-0.390	-0.300	1767	p < 0.01
Sand $\%$	Shannon equit.	-0.240	-0.280	-0.190	1767	p < 0.01
Sand %	Tree height CV	-0.200	-0.250	-0.150	1324	p < 0.01
Sand %	DBH CV	-0.160	-0.200	-0.110	1742	p < 0.01
Sand %	Stems ha	-0.270	-0.310	-0.220	1767	p < 0.01
Sand %	AGB	-0.270	-0.310	-0.220	1767	p < 0.01
Org. C %	CEC	0.300	0.250	0.340	1767	p < 0.01
Org. C %	MAP	0.450	0.410	0.490	1767	p < 0.01
Org. C %	PS	-0.300	-0.340	-0.260	1767	p < 0.01
Org. C %	MAT	-0.310	-0.350	-0.270	1767	p < 0.01
Org. C %	TS	-0.150	-0.190	-0.100	1767	p < 0.01
Org. C %	Sp. rich.	0.240	0.190	0.280	1767	p < 0.01
Org. C %	Shannon equit.	0.210	0.170	0.260	1767	p < 0.01
Org. C %	Tree height CV	0.110	0.060	0.160	1324	p < 0.01
Org. C %	DBH CV	0.100	0.050	0.140	1742	p < 0.01
Org. C %	Stems ha	0.200	0.160	0.250	1767	p < 0.01
Org. C %	AGB	0.250	0.210	0.290	1767	p < 0.01
CEC	MAP	0	-0.040	0.050	1767	p = 0.85
CEC	PS	-0.510	-0.540	-0.470	1767	p < 0.01
CEC	MAT	0.190	0.150	0.240	1767	p < 0.01
CEC	TS	0.020	-0.020	0.070	1767	p = 0.35
CEC	Sp. rich.	-0.070	-0.110	-0.020	1767	p < 0.01
CEC	Shannon equit.	0.090	0.040	0.130	1767	p < 0.01
CEC	Tree height CV	0.070	0.020	0.130	1324	p < 0.01
CEC	DBH CV	0.110	0.070	0.160	1742	p < 0.01
CEC	Stems ha	0.060	0.020	0.110	1767	p < 0.01
CEC	AGB	0.060	0.020	0.110	1767	p < 0.01
MAP	PS	-0.120	-0.170	-0.080	1767	p < 0.01
MAP	MAT	-0.100	-0.150	-0.060	1767	p < 0.01
MAP	TS	-0.690	-0.720	-0.670	1767	p < 0.01
MAP	Sp. rich.	0.390	0.350	0.430	1767	p < 0.01 p < 0.01
MAP	Shannon equit.	0.390 $0.100$	0.050	0.450 $0.150$	1767	p < 0.01 p < 0.01
MAP	Tree height CV	0.100 $0.210$	0.160	0.150 $0.260$	1324	p < 0.01 p < 0.01
MAP	DBH CV	0.210 $0.110$	0.060	0.160	1742	p < 0.01 p < 0.01
MAP	Stems ha	0.110 $0.090$	0.050	0.140	1742 $1767$	p < 0.01 p < 0.01
MAP	AGB	0.090 $0.220$	0.030 $0.180$	0.140 $0.270$	1767 $1767$	p < 0.01 p < 0.01
PS	MAT		-0.210			-
		-0.170		-0.120 0.110	1767	p < 0.01
PS DC	TS	0.070	0.020	0.110	1767	p < 0.01
PS DC	Sp. rich.	0.080	0.030	0.120	1767	p < 0.01
PS DC	Shannon equit.	-0.050	-0.100	-0.010	1767	p < 0.05
PS	Tree height CV	-0.030	-0.090	0.020	1324	p = 0.25

PS	DBH CV	-0.070	-0.120	-0.020	1742	p < 0.01
PS	Stems ha	-0.030	-0.080	0.020	1767	p = 0.2
PS	AGB	-0.110	-0.160	-0.070	1767	p < 0.01
MAT	TS	-0.320	-0.360	-0.280	1767	p < 0.01
MAT	Sp. rich.	-0.220	-0.260	-0.180	1767	p < 0.01
MAT	Shannon equit.	-0.110	-0.150	-0.060	1767	p < 0.01
MAT	Tree height CV	0	-0.050	0.050	1324	p = 1
MAT	DBH CV	0.090	0.040	0.130	1742	p < 0.01
MAT	Stems ha	-0.120	-0.160	-0.070	1767	p < 0.01
MAT	AGB	-0.090	-0.130	-0.040	1767	p < 0.01
TS	Sp. rich.	-0.300	-0.350	-0.260	1767	p < 0.01
TS	Shannon equit.	-0.090	-0.130	-0.040	1767	p < 0.01
TS	Tree height CV	-0.180	-0.230	-0.130	1324	p < 0.01
TS	DBH CV	-0.110	-0.160	-0.070	1742	p < 0.01
TS	Stems ha	-0.090	-0.140	-0.050	1767	p < 0.01
TS	AGB	-0.140	-0.180	-0.090	1767	p < 0.01
Sp. rich.	Shannon equit.	0.540	0.500	0.570	1767	p < 0.01
Sp. rich.	Tree height CV	0.310	0.260	0.360	1324	p < 0.01
Sp. rich.	DBH CV	0.300	0.260	0.340	1742	p < 0.01
Sp. rich.	Stems ha	0.570	0.540	0.600	1767	p < 0.01
Sp. rich.	AGB	0.500	0.460	0.530	1767	p < 0.01
Shannon equit.	Tree height CV	0.180	0.130	0.230	1324	p < 0.01
Shannon equit.	DBH CV	0.210	0.170	0.260	1742	p < 0.01
Shannon equit.	Stems ha	0.480	0.440	0.520	1767	p < 0.01
Shannon equit.	AGB	0.390	0.350	0.420	1767	p < 0.01
Tree height CV	DBH CV	0.510	0.470	0.550	1324	p < 0.01
Tree height CV	Stems ha	0.190	0.140	0.250	1324	p < 0.01
Tree height CV	AGB	0.330	0.280	0.370	1324	p < 0.01
DBH CV	Stems ha	0.240	0.200	0.280	1742	p < 0.01
DBH CV	AGB	0.490	0.460	0.530	1742	p < 0.01
Stems ha	AGB	0.770	0.750	0.790	1767	p < 0.01