Environmental heterogeneity patterns plant species richness and turnover in two hyperdiverse floras

Running title: Environmental heterogeneity and plant species richness

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Abstract

- 2 Aim:
- 3 Location: The Greater Cape Floristic Region in southwest Africa (the Cape), and the Southwest Australia
- 4 Floristic Region (SWA)
- 5 **Taxon:** Vascular plants
- 6 Methods: Geospatially explicit floral and environmental data, non-parametric statistics, boosted regression
- 7 tree modelling
- 8 **Results:** The Cape is more environmentally heterogeneous and has higher levels of floristic turnover than
- 9 SWA. We find that environmental heterogeneity is the main predictor of species richness in the Cape, and
- 10 somewhat less so for SWA. Edaphic conditions are found to be of more biologically important in the Cape,
- though this is contingent on the quality of the data modelled.
- 12 Main conclusions:
- 13 Keywords: biodiversity, environmental heterogeneity, fynbos, Greater Cape Floristic Region, kwongan,

14 macroecology, species richness, species turnover, vascular plants, Southwest Australia Floristic Region

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22 1 Introduction

- 23 Biodiversity represents the variety of species and the ecological and evolutionary processes that bring about
- 24 those species (???; Bøhn & Amundsen, 2004). Studying the distribution of biodiversity in space is a major
- 25 avenue of biological research (???; Kreft & Jetz, 2007). Regional-scale geographic patterns in species richness
- 26 have long been studied, particularly in biodiversity hotspots (???; Cook et al., 2015). The spatial distribution of
- 27 species richness can be explained in terms of the physical environment. Properties of the environment have
- 28 been suggested to influence species richness in three ways: (i) resources and energy, which can determine the
- 29 number of species able to co-exist in an area (Gaston, 2000; Kreft & Jetz, 2007; Mouchet et al., 2015); (ii)
- 30 stability through time, which enables species' persistence; and (iii) spatial heterogeneity, which can stimulate
- 31 ecological speciation and possible barriers to gene flow and can facilitate greater levels of species' co-existence
- 32 (Thuiller et al., 2006; Mouchet et al., 2015; Cramer & Verboom, 2016). The physical environment, then, can be
- used to explain species richness in both a local-deterministic and historical sense (Ricklefs, 1987).
- 34 The maintenance of species richness, particularly the coexistence of high numbers of species in biodiversity
- 35 hotspots, is often regarded as "paradoxical" (Hart et al., 2017), and is a central problem in ecology (Ricklefs,
- 36 1987; Kreft & Jetz, 2007; Hart et al., 2017). Species richness is constrained by the ability of habitats to support
- 37 a variety of species—its ecological carrying capacity (Mateo et al., 2017). This is exemplified in modelling
- 38 approches, wherein species richness is a function of environmental predictors in a correlative framework
- 39 ("macro-ecological models"; Mateo et al., 2017). Macroecological models of species richness implicitly

- 40 assume that communities are saturated, following species-area and species-energy relationships, and at
- 41 equilibrium with the environment (Mateo et al., 2017).
- 42 A solution to the "paradox" of species co-existence is environmental heterogeneity (EH): a more heterogeneous
- 43 environment gives rise to a larger environmental space, and can thus facilitate co-existence between more
- 44 species, generating more diverse species assemblages. Heterogeneity in the physical environment is known to
- 45 be positively associated with species richness (Rensburg et al., 2002; Hart et al., 2017), and has been
- demonstrated to do so across many taxa—e.g. Canadian butterflies (???), European vertebrates (Mouchet et al.,
- 47 2015), South African birds (Rensburg et al., 2002), in communities along marine continental margins (Levin et
- 48 al., 2010), French scarab beetles (Lobo et al., 2004), and for global terrestrial plants (Kreft & Jetz, 2007). The
- 49 spatial scale of heterogeneity, or "grain" of the environment, is also important to consider (Hart et al., 2017), as
- 50 spatial scale in absolute environmental conditions has also been explored (???; Baudena et al., 2015; Mouchet
- et al., 2015). Species co-existence and biodiversity maintenance is indeed suggested to be scale-dependent
- 52 (Hart et al., 2017).
- 53 EH is often under-represented in macro-ecological models of species richness, and has recently been found to
- 54 explain up to ca. 95% of biome level species richness across South Africa (Cramer & Verboom, 2016). Indeed,
- 55 models that include EH yield better estimates of the richness of the Cape flora (Thuiller et al., 2006; Cramer &
- 56 Verboom, 2016). Mediterranean-type terrestrial biodiversity hotspots, such as the Cape flora included in the
- 57 models by Cramer & Verboom (2016), present interesting study systems in which to investigate the relationship
- 58 between the environment and species richness. These systems exhibit far greater species richness than
- 59 predicted by their areas, productivities and latitudes (Cowling et al., 1996; Kreft & Jetz, 2007). There are five
- 60 Mediterranean biodiversity hotspots on Earth: the California Floristic Province, the Mediterranean Basin, the
- 61 Chilean Winter Rainfall-Valdivian Forests, the Greater Cape Floristic Region, and the Southwest Australia
- 62 Floristic Region (Cowling et al., 1996; Hopper & Gioia, 2004; Cook et al., 2015). These ecosystems have
- 63 regular fire-cycles (Cowling et al., 1996), climatic buffering, and long term stability (Kreft & Jetz, 2007),
- 64 shrubby, sclerophyllous flora (Hopper & Gioia, 2004). Together, they account for ca. 20% of global vascular
- 65 plant species, yet only ca. 5% of global land surface areas (Cowling et al., 1996). Various hypotheses have
- been proposed to explain the high levels of plant species richness in these regions (Cook et al., 2015). The
- 67 species accumulation hypothesis states that the stability of these regions has allowed many species to accrue.
- 68 The species co-existence hypothesis states that these hotspots may facilitate greater degrees of species
- 69 co-existence in smaller spatial areas, due to fine-scale heterogeneity in their environments. Indeed, EH has
- 70 evolutionary implications too, stimulating ecological speciation across sharp environmental gradients.

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Both the Southwest Australia Floristic Region (SWA) and the Greater Cape Floristic Region (Cape) are
     Mediterranean-type biodiversity hotspots, particularly in terms of plant species. Where the Cape (with an area
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     of ca. 189,000 km<sup>2</sup>) is known to contain about 11,400 plant species (about 0.060 species per km<sup>2</sup>), SWA (area
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     of ca. 270,000 km<sup>2</sup>) has about 3,700 species (0.014 species per km<sup>2</sup>) (???). So, the Cape has ca. 4.3 times as
     many species per km<sup>2</sup> as SWA. The Cape and SWA are appropriately often compared, due to the similarities
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     between their environments (e.g. oligotrophic soils, an oceanically buffered moderate climate) and their plants'
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     ecologies (Hopper & Gioia, 2004). These two regions present unique flora out of the five Mediterranean
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     systems, with high levels of endemism (Cowling et al., 1996), and many obligate fire-adapted species (Cowling
     et al., 1996). Similarities withstanding, SWA is topographically and edaphically distinct from the Cape. The
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     former is topographically rather uniform (i.e. flat)—uniquely so among the world's five Mediterranean-climate
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     regions (Hopper & Gioia, 2004)). SWA possesses a mesoscale chronosequence dune system (Laliberte et al.,
     2014; Cook et al., 2015), while the Cape is mountainous, topographically heterogeneous, and therefore
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     associated with a large degree of spatial climatic variability, with a fine-scale mosaic of geologies and soils
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     (Cowling et al., 1996; Cramer et al., 2014; Verboom et al., 2017).
     Both regions have sources of edaphic heterogeneity, but at different scales. This edaphic variability may aid in
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     explaining the species richness in these regions (Beard et al., 2000; Verboom et al., 2017). EH of many forms
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     will likely be important in macro-ecological models in both regions, as both regions have been relatively
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     environmentally stable over evolutionary time-scales (Wardell-Johnson & Horwitz, 1996; Hopper & Gioia,
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     2004; Lambers et al., 2010; Cramer et al., 2014; Laliberte et al., 2014; Cook et al., 2015). For the Cape, high
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     levels of species richness are thought to result from long term climatic stability, and fine grain variation in
     geology and soils (Cramer et al., 2014). The question thus arises whether heterogeneity is a significant
     contributor to SWA species richness. In the absence of topographic variability in SWA, it is proposed that the
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     heterogeneity of that region is due to the juxtaposition of soil types (Laliberte et al., 2014; Cook et al., 2015),
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     creating extreme edaphic variation.
     Our hypotheses concern the Cape and SWA's environments and floras. Our main hypothesis is that the Cape
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     possesses greater abiotic heterogeneity, and at finer grain, compared to SWA, such as to explain the Cape's
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     greater species richness per unit area, and proposed greater levels of species turnover between areas. We also
     conjecture that the heterogeneity that predicts species richness in SWA will be more pronounced in terms of
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     edaphic variables. Here we attempt to assess five key predictions of this hypothesis, additionally investigating
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     a seventh prediction to test the conjectured role of edaphic heterogeneity in SWA. Dealing with the two
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     regions' environments, we assess (i) whether the Cape environment is more heterogeneous than that of SWA
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and (ii) whether the Cape environment has more pronounced heterogeneity at finer scales than that of SWA.

Dealing with the distribution of species in the two regions, we assess (iii) whether the Cape exhibits greater

levels of species turnover between areas. Relating each regions' environment and flora, we finally assess (iv)

whether species richness and species turnover are adequately predicted by EH in both regions and whether (v)

species richness and species turnover are better predicted by different forms of EH in either region (e.g. the

importance of edaphic heterogeneity in SWA).

108 2 Materials and methods

109 **2.1 Overview**

- Our analyses defined boundaries for each region, those regions' environmental data and geospatially-explicit
- vascular plant occurrence records, all based on publicly available data. The environmental variables chosen
- 112 (Table 1) for this study were intended to cover a reasonable spread of climatic, edaphic, and ecologically
- 113 relevant environmental axes, and are not intended to be exhaustive. We selected variables describing
- topography (elevation), productivity (NDVI), soil status and climate and climatic seasonality.
- We carried out this investigation at four principal spatial scales: 0.05° x 0.05° squares (the finest common
- resolution among the environmental data sources used), quarter degree squares (QDS) (Larsen et al., 2009),
- 117 half degree squares (HDS) (Larsen et al., 2009) and three-quarter degree squares (3QDS). For the Cape, most
- 118 plant occurrence records are only accurate to QDS level. Thus, analyses involving species occurence data were
- 119 necessary limited to scales including and above QDS.
- 120 Analyses were performed in R v3.4.0–3.5.1 (R Core Team, 2018). Version-numbers of specific R packages
- used are presented in the bibliography.

122 2.2 Environmental data sources

- 123 The GCFR was treated as the areas occupied by the Succulent Karoo and Fynbos biomes in the current
- delineation of South Africa's biome boundaries (Mucina & Rutherford, 2006). The SWAFR was treated as the
- 125 areas occupied by the Southwest Australia savanna, Swan Coastal Plain Scrub and Woodlands, Jarrah-Karri
- 126 forest and shrublands, Southwest Australia woodlands, Esperance mallee, and Coolgardie woodlands in the

- 127 World Wildlife Fund Terrestrial Ecoregions dataset (Olson et al., 2001) in order to closely match the currently
- delineated SWAFR (Gioia & Hopper, 2017, Hopper & Gioia (2004)). For the sake of readability, we shall refer
- to the GCFR and SWAFR simply as the Cape and SWA from hereon.
- 130 Geospatially-explicit raster layers were acquired for a selection of environmental variables (Table 1), for the
- regions of interest. Raster data were re-projected to a common coordinate reference: WGS84 (NIMA, 2000),
- using the "rgdal" (???) package in R (R Core Team, 2018). All data were re-sampled to 0.05° resolution using
- the "resample" function in the R package "raster" (???), with the "bilinear" method.
- 134 An emphasis was made on using satellite-derived environmental data in this work, in order to minimise
- differences in data quality and methodologies between the Cape and SWA. Additionally, satellite-derived data
- have been shown to benefit regional-scale species distribution models (Deblauwe et al., 2016), thus motivating
- their use in this regional-scale study. The environmental data used in this study were derived from NASA's
- 138 SRTM digital elevation model (Farr et al., 2007), NASA's MODIS/Terra spectroradiometric data for land
- surface temperature and NDVI, the Climate Hazards Group's CHIRPS rainfall dataset (Funk et al., 2015), and
- the International Soil Reference and Information Centre's SoilGrids250m edaphic dataset (Hengl et al., 2017)
- (Table 1). SRTM and MODIS are entirely derived from satellite measurements, whereas CHIRPS is
- interpolated from weather station data with satellite-derived radiometric measurements. SoilGrids250m is a
- machine-learning derived product, based on soil measurements as a function of many covariates, including
- MODIS and STRM sources (see Hengl et al., 2017), using random-forests and other classification-tree-based
- methods, including gradient-boosting. For the soil data considered here (Table 1), we used depth-interval
- weighted average values as the value for a particular soil variable in a given place.
- 147 Climatic and spectral data arise from satellites monitoring properties of the Earth's surface through time. We
- therefore use the mean annual values for rainfall, surface temperature, and NDVI in each pixel in our analyses.
- Pronounced seasonality of rainfall is a known feature of mediterranean systems (???). We describe this
- seasonality by computing computing the precipitation in the driest quarter (PDQ), using methods based on the
- "biovars" function in the R package "dismo".

152 2.3 Plant occurrence data

- 153 Geospatially-explicit records of vascular plant occurrences were downloaded from the Global Biodiversity
- 154 Information Facility (GBIF, Table 1). Queries were made for tracheophyte records from within the borders of

the Cape and SWA as treated here (GBIF, 24 July 2017, GBIF (24 July 2017)). Only records with defined 155 species and intra-specific ranks were kept. Intra-specific occurrences were treated as simply being 156 representative of their species. This resulted in FIXME unique species names in the Cape, and FIXME in SWA. 157 We cleaned these data using the R package "taxise" (???, (???)) to check that these species names had 158 accepted-status among taxonomic databases. We queried two major taxonomic databases: the Global Name 159 160 Resolver (GNR), and the Taxonomic Name Resolution Service (TNRS). Should either one of these services return at least one match for a given name, then that name was accepted. Those names for which no full 161 binomial matches were found in either database were excluded from the final list of species. The number of 162 species names excluded totalled at FIXME and FIXME for the Cape and SWA respectively. Especially for 163 SWA, these numbers may be deemed appreciably high. But, the occurrence records that would be dropped, as a consequence of these names' removals, appeared randomly distributed in geographic space in both regions. As 165 such, any effect of the loss of these records in this analysis is likely uniform within the two regions. 166 After the unaccepted names were removed, it was important to ensure that a species was not listed under 167 multiple synonyms. Such cases would skew estimates of species richness and turnover in this study. In light of 168 169 this, the remaining names were queried in the Tropicos and Integrated Taxonomic Information System (ITIS) 170 databases for their known synonyms, again using "taxize". These were collated to produce a nomenclatural "thesaurus" for the Cape and SWA species. This consisted of a list of the accepted species names in a region, 171 each associated with a list of known synonyms. We amended species' names in the GBIF occurrence data, in 172 order ensure species were listed under only one of these synonyms, replacing all appearances of a species' 173 synonyms with the first synonym used in the list. 174 Lastly, We removed any species from both regions that are invasive aliens or non-indigenous. Alien species 175 lists for plants in South Africa and Australia were acquired from the IUCN's Global Invasive Species Database 176 (http://www.iucngisd.org/gisd/). 177 The final total plant species richness in each region was FIXME and FIXME for the Cape and SWA 178 respectively. These final collections of species occurrence records were converted to raster-layers, wherein 179

pixel-values represented the species richness of vascular plants within that pixel. These rasters were produced

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at QDS, HDS, and 3QDS resolutions.

182 2.4 Analyses

183 2.4.1 Quantifying environmental heterogeneity

In order to assess predictions (i) and (ii), we needed to describe the EH in both regions. Using the R package "raster" (???), we used a modified version of the "roughness" index in the "terrain" function. For a three by three neighbourhood $\bf N$ of cells, our index of roughness R is the average square-root of the squared difference between each of the n neighbour cells' values x_i and the central focal cell's value $x_{\rm focal}$:

$$R(\mathbf{N}) = \frac{1}{n} \sqrt{\sum_{i=1}^{n} (x_{\text{focal}} - x_i)^2}$$
 (1)

This value, notionally equivalent to the standard deviation of values relative to the focal value, is ascribed to the focal cell. Note, in order to use as much data from within regions' borders as possible, roughness was computed if a focal cell had at least one neighbour cell. Using this index, we produced raster layers of each of our nine environmental variable's heterogeneity. We compared the distributions of "roughness" values in each variable in each region with non-parametric Mann-Whitney U-tests, as almost all variables were highly non-normal, and could not be normalised by log-transformations. We also compare the effect size of the Cape vs SWA using the "common language effect size" (CLES), using the R package "canprot". The CLES is the proportion of all pairwise comparisons between two sample groups' observations where one group's value is greater than the other's. We calculated the CLES as the proportion of pairs where Cape roughness values were greater than that of SWA. This allowed us to assess prediction (i). To compare the spatial scales of heterogeneity (prediction (ii)) between each region, we repeated this analysis at all four spatial scales. This entailed recalculating the roughness layer for each variable after the original layer (0.05 degrees resolution) had been rescaled to each of the coarser resolutions.

2.4.2 Quantifying species turnover

Regarding prediction (iii), we wished to compare the general degree of species turnover in each region. To compare the extent of species turnover between the Cape and SWA, we determined two metrics of species turnover. The first, computes the mean species turnover as Jaccard distances (???) between each pair of QDS within each HDS (\overline{J}_{QDS} , based on HDS with $2 \le n \le 4$ QDS) in both regions. The second is defined in terms of Whittaker's additive definition of β -diversity (???), as follows:

$$\gamma = \alpha + \beta \tag{2}$$

Here, we treat species richness at the HDS-scale (S_{HDS}) as γ -diversity and at the QDS-scale (\overline{S}_{QDS}) as α -diversity. Intuitively, the species richness of an area is the result of some combination of the richness of sites within that area and the difference in species complements between those sites. Thus, we partition γ -diversity as in Equation (2), such that β -diversity is the difference between γ - and α -diversity. We compare the distributions of \overline{J}_{QDS} and T_{HDS} using non-parametric Mann-Whitney U-tests, in order to guard against non-normality.

213 2.4.3 Predicting richness and turnover with environmental heterogeneity

214 Regarding prediction (iii), we wished to compare the general degree of species turnover in each region. For (iv) and (v) we modelled species richness (S) and turnover as a function of various combinations of environmental 215 and environmental heterogeneity variables in both regions using boosted regression-tree (BRT) modelling 216 techniques. This allowed us to explore which axes of environmental heterogeneity are most influential on 217 vascular plant species richness and turnover, and the differences in the importance of such axes between the 218 219 Cape and SWA. BRTs are a flexible machine learning-based model of response variables and do so without involving normal 220 null-hypothesis significance testing (Elith et al., 2008), and have been employed previously to model species 221 richness (Thuiller et al., 2006; see Mouchet et al., 2015; Cramer & Verboom, 2016) as macro-ecological 222 models. BRTs are developed through the iterative generation of non-linear regression trees. BRTs are an 223 ensemble-approach, in which a prediction \hat{y}_i is based on the weighted sum of the predictions of progressively 224 "less important" regression trees (t_k) , as opposed to the predictions of one tree (Elith et al., 2008). For $k \to nt$ 225 number of trees, where each tree is itself a function of the matrix \mathbf{X} of j predictor variables $(t_k = f(x_{ij}))$, a 226 BRT-model can be represented as follows: 227

$$\widehat{y}_i = \sum_{k=1}^{nt} w_k t_k \tag{3}$$

BRTs have two major meta-parameters over which users have control (???): the learning rate (lr, the rate at which iterative trees reduce predictive deviance during model-training, controlling the contribution of each tree

- to the final model) and tree complexity (tc, the number of nodes on a given regression-tree, i.e. the maximum
- 231 interaction depth the model is permitted to fit).
- BRTs were implemented here to predict both vascular plant species richness and turnover in each HDS, as a
- 233 function of environmental variables and environmental roughness values in those cells, as Gaussian responses,
- thus resulting in two BRT-models for each region. We treated richness as S_{HDS} and turnover as \overline{J}_{QDS} . The
- 235 natural logarithm of species richness was used, in order to satisfy the assumptions of a Gaussian response.
- Note, this is not strictly because BRTs have any parametric assumptions concerning the distribution of the
- 237 response variable, but rather to aid in applying the Gaussian-family of BRT algorithms to the richness data
- available. Additionally, BRTs were implemented to predict vascular plant species richness at the QDS-scale
- 239 (S_{QDS}) , thus resulting in a total of six BRT-models presented here.
- As recommended by Elith et al. (2008), BRT models were trained on a set of non-collinear predictor variables
- using "gbm.step" in "dismo" (???) and "gbm" (???). Collinear predictor variables can skew the interpretation
- of results, as the relatively influence of mutually collinear variables is reduced. Collinearity among the nine
- 243 environmental predictor variables and their respective nine roughness-equivalents was assessed using
- "removeCollinearity" in the R package "virtualspecies" (???) separately for each region, such that variables
- 245 were no more than 80% collinear (Pearson's $r \ge 0.80$). When faced with a cluster of collinear variables, one
- variable was chosen manually therefrom. Where possible, the roughness-equivalent of an environmental
- variable was included if its absolute-equivalent could also be included. When interpreting the results of BRTs,
- 248 it is important to consider the effects of the variables included as representative of the effect of the excluded
- variables with which it was found to be collinear.
- In order to select ideal lr and tc all models (described below) were trained on the final non-collinear predictor
- sets iteratively for 25 combinations of a range of tc values (1 to 5) and a range of lr values (0.01, 0.005, 0.001,
- 5×10^{-4} , 1×10^{-4}). The function "gbm.step" optimises the number of trees (nt) using cross-validation during
- 253 model training (Elith et al., 2008) by halting iteration when predictions begin to overfit. For all models, we
- used 10 cross-validation folds (i.e. use 10 different randomly selected training data sets), a tolerance-threshold
- of 0.001, a bagging-fraction of 0.75 (proportion of training data randomly chosen to generate each tree), and
- trained models starting with 50 trees, with each iterative step adding 50 trees at a time, up to a maximum of
- 257 10,000 trees. Following this iterative parameter optimisation, Gaussian BRT models were constructed with
- 258 tc = 3 and lr = 0.001, along with the other settings described.
- The optimum configuration of lr and tc for the final model is a trade-off between model fit (e.g. pseudo- R^2 ;

Equation (4)) and complexity (nt). A tc of 5 was chosen for the final model. This follows the recommendations of Elith et al. (2008), where lr and tc are advised to be adjusted inversely. This was chosen in order to account for the complex interactions determining species richness. To avoid overfitting, an intermediate lr of 0.001 was chosen.

264 2.4.4 Assessing BRT-predictions' fit

BRT-model performance can be described by measuring the variance in a dataset a BRT-model has explained, quantified here by R_{pseudo}^2 , which is the proportion of null deviance D_{null} explained by some model i. Formally, it is defined as follows:

$$R_{\text{pseudo}}^2 = 1 - \frac{D_i}{D_{\text{null}}} \tag{4}$$

The derivation of this metric is not easy to interpret, as it is not immediately clear what model deviance is. 268 Alternatively, comparing expected (i.e. model-predicted) and observed data has more heuristic appeal. We 269 employed this metric of BRT-model performance too. We regressed expected against observed richness and 270 turnover, and calculated the R^2 -value for those regressions (hereafter $R^2_{\rm E-O}$). 271 The BRT-model fitting algorithm contains intrinsic stochasticity, due to the random partitions made in a dataset 272 during cross-validation. Though this randomness is usually negligible (e.g. variables' contributions vary from 273 run-to-run by a few decimal places), we reran each of the six BRT-models (see above) 1000 times in order to 274 account for this stochasticity. Where indicated, we either present the averages of these replicate-models' results 275 or the results of a representative model from each set of replicates. 276 277 In order to assess the reliability of the conclusions drawn from these models, we randomly permuted the response data (S_{QDS}, S_{HDS}) and \overline{J}_{QDS} with respect to the environmental and heterogeneity data, and reran all 278 six BRT-models 999 times (with the final non-collinear predictor sets and preconfigurations above). This also 279 allows us to remove any effect of spatial autocorrelation in generating the observed correlations between 280 patterns of species occurrence and environment (???), and to allow us to assess the significance of our results 281 relative to a random null. Notably, as the predictor variables themselves are likely spatially autocorrelated, 282 correlation structure in model residuals is accounted for by the correlation structure in the environmental data. 283 Nonetheless, we wished to demonstrate our results more robustly and thus carried out these permutation tests. 284 For all six models, the majority of the 999 permuted models failed to find associations between the response 285

and predictor variables. The results of those that succeeded to fit a model to completion (usually ca. 200 out of 999) are presented. The replicate and permuted BRT-models were compared using various measures of model performance (above; nt, $R_{\rm pseudo}^2$ (Equation (4)), $R_{\rm E-O}^2$) and the ranks of these values for each replicate BRT-model relative to the 999 permuted models for that region/scope.

290 3 Results

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3.1 Describing environmental heterogeneity across scales

Across all variables considered, the Cape is more environmentally heterogeneous in the majority of pairwise 292 comparisons of grid-cells (CLES > 0.50, Mann-Whitney U-test: P < 0.05, Figure 1). The degree to which 293 the Cape is more heterogeneous varies between environmental variables. These effects also vary somewhat 294 across spatial scales. In some variables, the differentiation between Cape and SWA heterogeneity lessens at 295 coarser scales (Figure 1b). Indeed, when comparing the overall ranking and medians of Cape vs SWA 296 roughness values for each variable, we only find non-significant differences at the 3QDS scale (Mann-Whitney 297 U tests, P > 0.05, Figure 1b). 298 Most obviously, and as expected, topographic heterogeneity is generally greater in the Cape than in SWA 299 (Figure 1). Though SWA has a slightly wider distribution of elevational roughness values at coarse scales 300 compared to fine scales than the Cape, the relative difference in heterogeneity between the two regions seems 301 invariant with spatial scale ($CLES \approx 0.95$, Figure 1b). This concurs with our expectations, as the Cape is 302 mountainous and known to have steep elevational gradients (???), while SWA is much more topographically 303 uniform. Elevational roughness differs between the two regions in the manner we expected, giving us more 304 confidence in the environmental reality of our other results here. 305 Climatic heterogeneity is less differentiated between the Cape and SWA than with topographic heterogeneity 306 307 (Figure 1a), though the Cape is indeed more climatically heterogeneous (Figure ??b). Notably, the difference between roughness in mean annual rainfall (R MAP) and land surface temperature (R Surface T) in the Cape 308 and SWA is less pronounced when considered at coarse spatial scales (Figure ??b). At all spatial scales 309 considered, roughness in rainfall seasonality (R PDQ), however, is equally more heterogeneous in the Cape 310 311 than SWA. Biological productivity, as measured by NDVI, is fairly similarly heterogeneous in the Cape and SWA (CLES < 0.60, Figure 1). Concerning edaphic variables, the Cape and SWA are similarly 312

heterogeneous at coarser scales, particularly in terms of CEC and Soil C ($CLES \approx 0.50$, Figure 1b).

3.4 3.2 Comparing species turnover in the two regions

- Following calculations of \overline{J}_{QDS} and T_{HDS} for each HDS-cell in each region, we also used non-parametric
- Mann-Whitney U-tests to compare the distributions of values in the Cape and SWA. The Cape possesses
- generally greater floristic turnover than SWA, for both measures of turnover defined here (P < 0.0001, Figure
- 2a,b). \overline{J}_{QDS} measures the average pairwise Jaccard distance between QDS-cells in each HDS-cell. T_{HDS} ,
- however, represents the inferred β component of γ -diversity. As γ -diversity (= S_{HDS}) in the Cape has a
- greater β -diversity component (= T_{HDS}) than SWA, the complement is necessarily true: γ -diversity in the
- 321 Cape has a lesser α -diversity component (= \overline{S}_{QDS}) than SWA.

322 3.3 Predicting richness and turnover with environmental heterogeneity

- 323 We found vascular plant species richness and turnover both to be predicted primarily by environmental
- heterogeneity in the Cape (Figure 3a-c) and at least in-part by environmental heterogeneity in SWA (Figure
- 325 3d-f). Our six BRT-models performed adequately, and detected relationships between patterns of species
- occurrence and the environment that do not occur in the permuted datasets (Figures 4 and 3, Table 2).
- 327 BRT-models of species richness at the QDS-scale in each region generally performed best, as these models had
- fit greater number of trees (nt, Figure 4a), and possess higher R^2 -values (Figure 4b,c). SWA models of species
- 329 richness and turnover at the HDS-scale out-performed Cape models. At the QDS-scale, the Cape and SWA
- models performed equally well (Figure 4, Table 3).
- 331 Across our BRT-models of species richness and turnover, the importance of different environmental variables
- in predictions differed substantially between the Cape and SWA. Additionally, the relative importance of
- absolute and heterogeneity variables also differs between the Cape and SWA (Figure 3). Most obviously,
- 334 species richness and turnover in the Cape are predicted mostly by environmental heterogeneity, which is not the
- case in SWA (Figure 3). Species richness and turnover in the Cape are predicted by a broad suite of
- environmental variables, with no individual variable contributing more than ca. 20% to any model prediction
- 337 (Figure 3a-c). The SWA models' predictions, however, are largely determined by MAP (Figure 3d-f).
- Species richness at QDS-scales (= \overline{S}_{QDS}), and to a lesser extent at HDS-scales (= S_{HDS}), in the Cape is

predicted largely edaphic conditions (Figure 3a,b). Species richness in SWA, at both scales, is mostly predicted 339 by MAP and other climatic variables (Figure 3d,e). Interestingly, topographhic heterogeneity did not feature as 340 341 highly in contributing to Cape predictions as we expected (Figure 3a–c).

It is important to consider variables not included formally in these BRT-models that were found to be collinear with some of the variables included (see SI). Here, we interpret the effects of variables excluded from the 343 analyses as well as those included, as the forms and importances of these relationships are likely similar. In the 344 Cape (concerning clusters of collinear variables relevant to relevant to those retained during BRT-model 345 fitting), MAP was included in the BRT-analyses as representative of a cluster of collinear variables consisting 346 of itself, NDVI, surface T and soil C at the. Roughness in soil clay content represented itself, roughness in soil 347 pH and roughness in NDVI. In SWA, MAP was select as representative of itself, NDVI and soil C. 348

349 In the equations below, collinear groups of variables are listed as predictors enclosed within braces.

350 Cape:

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$$\overline{S}_{\text{QDS}} \sim pH + RSurfT + \begin{cases} MAP \\ NDVI \\ SurfT \\ SoilC \end{cases} + \begin{cases} RClay \\ RpH \end{cases} + RElev$$

$$S_{\text{HDS}} \sim RSurfT + RElev + RCEC + \begin{cases} RMAP \\ RPDQ \\ RNDVI \end{cases} + \begin{cases} MAP \\ NDVI \\ SurfT \\ SoilC \end{cases}$$

$$T_{\text{HDS}} \sim RElev + RSurfT + \begin{cases} RMAP \\ RPDQ \\ RNDVI \end{cases} + Elev + RCEC$$

$$\begin{cases} RMAP \\ RPDQ \\ RNDVI \end{cases} + Elev + RCEC$$

SWA: 351

$$\overline{S}_{ ext{QDS}} \sim egin{cases} MAP \\ NDVI \\ SoilC \end{pmatrix} + PDQ + RMAP \\ SoilC \end{pmatrix} + CEC + RElev \\ SoilC \end{pmatrix}$$
 $T_{ ext{HDS}} \sim egin{cases} MAP \\ NDVI \\ SoilC \end{pmatrix} + RElev + CEC \\ SoilC \end{pmatrix}$

Our BRT-models of species richness in both regions rank environmental variables somewhat differently at

353 QDS- and HDS-scales (Figure 3a,b,d,e). These differences in rankings are similar to those between two

unrelated lists ($P_{1-2} > 0.05$, Figure 5). This suggests some scale-dependence of different environmental

355 variables' associations with species richness.

It is noteworthy that BRT-models of species turnover (= \overline{J}_{QDS} , at HDS-scales) (Figure 3c,f) rank variables

similarly to models of richness at HDS-scales ($P_{2-3} \le 0.005$, Figure 5). This is likely due to the fact that

358 proportional floristic turnover covaries with species richness. As such, though the signs of relationships

359 determining turnover may differ from those determining richness, the importances of different variables would

360 be similar.

361 In addition to different variables being more strongly associated with species richness and turnover in the Cape

compared to SWA (Figure 3), the forms of those relationships vary (Figure 6). We found MAP, and roughness

therein, to relate positively with species richness in both regions at both scales (Figure 6a,b,d,e). As MAP is

364 collinear with NDVI and soil C in both regions (and surface T in the Cape), this can be interpreted as the signal

of a biological productivity and resource availability associating with high levels of species richness.

The positive association of heterogeneity variables in the Cape as opposed to SWA (Figure 6a,b vs d,e) concurs

with their greater importance in BRT-model predictions (Figure 3).

The fact that species turnover $(T_{HDS} = \overline{J}_{QDS})$ in the Cape and SWA is largely predicted by the same

variables as species richness, but with opposite signs to its relationships (Figure 6c,f), is indicative of the

370 richness-dependence of the measure of floristic turnover used here (Jaccard distances) to quantify turnover at

371 the HDS-scale.

372 4 Discussion

I have provided support for the hypothesis that the difference in plant species richness between the GCFR and 373 SWAFR is accounted for by the fact that the GCFR is more abiotically heterogeneous than the SWAFR. As 374 expected, the GCFR is shown to possess (i) a quantifiably more heterogeneous environment, and (ii) is 375 heterogeneous at a finer spatial scale than the SWAFR. I have shown that vascular plant species richness (iii) 376 can be explained in terms of environmental conditions, including environmental heterogeneity, in both the 377 GCFR and SWAFR. Also, I have shown that (iv) the set of environmental axes that explain plant species 378 379 richness, both absolute and as heterogeneity, differs between the GCFR and SWAFR. These findings contribute towards an understanding of the ecological conditions that facilitate species coexistence (and likely stimulate 380 ecological speciation) in these two regions. 381 382 These two regions present differentiable environmental spaces, each with heterogeneity varying across spatial scales. The clear separation of the regions' topographic features is as expected (Figures ??A, ??). Indeed, 383 topography seems to be the most striking distinction between the regions. The Cape region has been found 384 previously to have the second highest median topographic heterogeneity of the five Mediterranean-climate 385 regions (Bradshaw & Cowling, 2014). The GCFR has a much wider range of scales exhibited in the 386 heterogeneity across its environmental axes. Notably, each region has finer scale heterogeneity in some 387 variables, and coarser scale in others—neither region is necessarily more fine or coarse than the other, as it 388 389 depends on the variable concerned. BRT-models of species richness in both regions reveal species richness to depend on those environmental axes that differentiate the two regions (Figures ??), ??). The importance of 390 variables is also shown to vary with spatial scale (Figure ??), as previously suggested may be the case when 391 392 modelling geographic patterns of biodiversity (Baudena et al., 2015). Indeed, as Cowling et al. (1996) describes differing patterns of species richness across spatial scales, so do the predictors of those patterns vary 393 with scale (Hart et al., 2017). 394 The fact that a combination of absolute and roughness variables is also as predicted by the hypothesis in this 395 study. In the models developed by Cramer & Verboom (2016) for South Africa, roughness in topography was 396 largely superseded as an important predictor of species richness by other roughness variables. My models, 397 however, did not show this. Similar to the study by Rensburg et al. (2002), my models revealed roughness in 398

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topography and other variables to be important. Although, Rensburg et al. (2002) considered differences
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     within pixels, as opposed to this study, which considered differences between pixels. My models, those of
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     Cramer & Verboom (2016), and those of Rensburg et al. (2002), do not all concur as to the role of roughness in
     elevation vs. more biologically meaningful variables in explaining species richness. The source of these
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     discrepancies is unclear, though no doubt complex. The complements of environmental variables and
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     methodologies used in these studies do differ, limiting extensive comparison between these analyses.
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     The determinants of vascular plant species are shown to be region specific (Figures ??, ??, ??). The importance
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     of MAP and roughness in rainfall seasonality (PCV) in predicting richness in the SWAFR (Figure ??I, ??J),
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     aligns with the steep climatic gradients observed there (Cook et al., 2015). The soil variables that determine
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     plant species richness in the model for the SWAFR (Figures ??K, ??L) differ to those that determine richness in
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     the GCFR (Figures ??G, ??H), further highlighting the edaphic differences between these two regions.
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     Although both are nutrient leached systems, the SWAFR is flat, with soil-chronosequences (Laliberte et al.,
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411
     2014; Cook et al., 2015), while the GCFR is mountainous (Cowling et al., 1996; Cramer et al., 2014; Verboom
     et al., 2017). The importance of roughness in soil density, and absolute texture, in the SWAFR (Figures ??K,
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     ??L) highlights the changes in soil that are associable with age of the substrate (e.g. particle size) as being
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     biologically relevant to species richness. The positive effect of soil clay content on species richness in the
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     SWAFR aligns with the findings of Laliberte et al. (2014) that richness in the SWAFR increases with soil age.
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     NDVI is more heterogeneous across the GCFR than the SWAFR (Figures ??A). The fact that thermal variables
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     tend to be more rough in the GCFR (Figure ??A) is likely due to possible covariance of the MODIS/Terra
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     products with topography, as MODIS data used here describes land surface temperature. As the GCFR is
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     topographically rugged, the roughness of NDVI may arise from this. Despite this, NDVI is an integrating
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     variable, which captures information about productivity, light availability, and soil nutrients (Power et al.,
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421
     2017). The fact that absolute NDVI contributes to predicting species richness in the GCFR, especially at finer
     spatial scales (Figure ??E) demonstrates the role of ecological productivity in facilitating the coexistence
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     diverse species assemblages. Environmental heterogeneity, then, is integral to explaining patterns of species
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     richness, but must be considered along with resource- and energy-availability axes. In so much as a diverse
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     environmental space supports more species, the materials and productivity required for biota to thrive are also
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     needed to support species (???; Gaston, 2000; Bøhn & Amundsen, 2004; Kreft & Jetz, 2007). As such, my
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     findings, along with those of previous studies (Rensburg et al., 2002; Thuiller et al., 2006; Kreft & Jetz, 2007;
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     Cramer & Verboom, 2016), suggest that there is ecological and evolutionary consequence to resource
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     availability and environmental heterogeneity, in that they tend to be positively associated with species richness.
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predictor, at coarse spatial scales, despite this variable not being particularly important within each region 431 432 separately (Figure ??). Though this model does not strictly consider the regions as separate, this finding may indicate that the relationship between clay content and species richness differs between the regions. So far as 433 clay content can be used to predict species richness, it matters more to those predictions when applied to large 434 sections (i.e. coarse scales) of each regions. 435 Kreft & Jetz (2007) modelled global terrestrial vascular plant species richness, which focussed on primarily 436 absolute environmental values, underestimated the richness of the Cape flora. Though Kreft & Jetz (2007) did 437 include topographic heterogeneity in their predictor set, topography is often a proxy for more biologically 438 meaningful variables (Cramer & Verboom, 2016). This explains why the inclusion of these variables 439 (e.g. roughness in mean annual precipitation) yields more accurate predictions of species richness. Indeed, 440 Thuiller et al. (2006) also included topographic heterogeneity. @Cramer 2016 described 68% of species 441 442 richness at the QDS scale across South Africa. Regarding the GCFR, depending on whether one consults pseudo- R^2 (Table 3), the ratio of mean predicted to observed richness per grid-cell (Table 5), or the 443 distributions of predicted vs. observed richness values per grid-cell (Figure ??), I have achieved a similarly 444 suitable level of predictive accuracy. There is, though, still unexplained species richness in light of my models. 445 As Cramer & Verboom (2016), Rensburg et al. (2002), Thuiller et al. (2006), and Mouchet et al. (2015) have 446 done, these macro-ecological models are a-historical. Evolutionary considerations of species richness in 447 geographic space are worthwhile, especially in regions with environments stable over evolutionary time. 448 The findings here are correlative. There are, however, many proposed mechanisms to explain the correlative 449 signals demonstrated here. My findings support the hypothesis that Mediterranean systems' plant species 450 richness is a function of spatial variability in environmental conditions. This can stimulate diversification, and 451 452 maintain that diversity by providing a range of habitats for species co-existence. Oligtrophic soils can stimulate an increase in functional diversity, through the evolution of diverse nutrient acquisition strategies (Lambers et 453 al., 2010; Verboom et al., 2017)—e.g. sclerophylly (Cramer et al., 2014; Cook et al., 2015). An aspect of the 454 environment I have neglected to consider is fire, shown to also contribute to predictions here in the GCFR 455 (Cramer & Verboom, 2016). Cardillo (2012) have shown the structuring forces behind species co-occurrence 456 patterns, and thus likely species richness, differ between species-pairs with different post-fire responses and 457 those with similar post-fire responses. 458

The combined BRT-model of species richness for both regions reveals soil clay content as an important

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Though the GCFR was correctly predicted to have, on average, more species per grid-cell at HDS and 3QDS

scales than the SWAFR, this was not the case for ODS grid-cells (Table 5). This demonstrates that the GCFR is 460 indeed overall more rich in plant species than the SWAFR, but a given HDS in the SWAFR contains fewer 461 462 species than a given GCFR HDS. Thus, the greater richness in the GCFR is a product of greater turnover in species at spatial scales no more coarse than the HDS. Species turnover is an interesting aspect to species 463 richness studies, as it species turnover is implicit to species-area and co-existence-area relationships (Hart et 464 al., 2017). One could expect patterns of endemism and species turnover to concur with patterns in 465 environmental heterogeneity to some degree. 466 Following from the understanding that functionally diverse assemblages, which are more likely to be more 467 species rich, are likely to arise and/or occur in areas with diverse ecological pressures (Molina-Venegas et al., 468 2015), one would expect, then, heterogeneous habitats such as those in Mediterranean-type biodiversity 469 hotspots to exhibit high levels functional beta diversity along steep environmental gradients (Molina-Venegas 470 et al., 2015). If the niches concerning these functions are phylogenetically conserved among those biota, then 471 472 one would also expect high levels of species and phylogenetic beta diversity along these gradients (Molina-Venegas et al., 2015). This concurs with the notion put forward by Power et al. (2017), wherein 473 megadiverse systems such as these represent the results of "phylogenetic niche conservatism on a 474 heterogeneous landscape". Thus, species and phylogenetic turnover should covary with environmental 475 heterogeneity in some way. Indeed, endemism, at certain scales, could also follow this pattern. Thuiller et al. 476 (2006) demonstrated that there is phylogenetic and biome related determinants of species richness. This makes 477 sense, in light of the difficulty of crossing biome boundaries in Mediterranean systems (Power et al., 2017). 478 NDVI and light availability, and the heterogeneity therein, are associated with high levels of floristic turnover 479 (Power et al., 2017). This may be indicative of ecological specialisation precluding species from crossing these 480 boundaries, thus increasing the level of endemism within a region, while also increasing the level of turnover, 481 and thus likely species richness, along environmental gradients. Although, this may be debated. Beard et al. 482 (2000) state that the high levels of endemism in SWAFR are function of habitat specialisation to soil mosaics. 483 Cf. Laliberte et al. (2014), who say that this endemism is likely due to environmental filtering along these soil 484 turnover sequences, as opposed to the juxtaposition of specialised species along soil gradients. 485 I have demonstrated support for the idea that environmental heterogeneity is positively associated with species 486 richness, particularly Mediterranean systems. In the SWAFR and the GCFR, high levels of endemism and 487 biodiversity are also likely the results of long-term landscape and climatic stability (Hopper, 1979). Thus, the 488 roles of environmental variability through space, and stability through time, are the two main ways in which the 489 environment relates to biodiversity in these regions. 490

491 4.1 Future studies (a.k.a. "to do after first review")

- 3QDS scale BRTs
- $S_{\rm HDS}\sim \overline{S}_{
 m QDS}+\overline{\delta_{ij}}_{??}$ (= " $\gamma=lpha+eta$ "-analysis) (see explore-turnover-metrics.pdf note,
- where $\overline{\delta_{ij}}(\mathbf{N})_{??} = \overline{\alpha_i}(\mathbf{N}) \times \overline{\beta_{ij}}(\mathbf{N}) = \overline{S}_{\mathrm{QDS}} \times \overline{J}_{\mathrm{QDS}}$

95 Table captions

- 496 Captions are also repeated alongside their respective tables for readability.
- 497 Table 1: Georeferenced vascular plant species occurence and environmental data sources used in this study.
- 498 Data were acquired for the Cape and SWA regions, with the temporal extent of data products used described
- 499 where applicable. Abbreviations are as follows: MAP, mean annual precipitation; PDQ, precipitation in the
- 500 driest quarter; CEC, cation exchange capacity.
- Table 2: Average percentile-ranks for BRT-model performance measures (nt, $R_{\rm pseudo}^2$ (Equation (4)), $R_{\rm E-O}^2$)
- of 1000 replicate BRT-models relative to 999 BRT-models fit to permuted datasets. Ranks approaching one
- 503 indicate that a set of replicate BRT-models had greater values than the permuted models.
- Table 3: Estimated differences between replicate Cape and SWA BRT-models' performance measures (nt.
- 505 $R_{\rm pseudo}^2$ (Equation (4)), $R_{\rm E-O}^2$) following t-tests. Positive values indicate that the Cape models had greater
- values. In all cases, the Cape and SWA had highly significantly different values for these quality measures
- 507 (P < 0.0001).

508 Figure captions

- 509 Captions are also repeated alongside their respective figures for readability.
- Figure 1: Comparisons of different types of environmental heterogeneity between the the Greater Cape
- 511 Floristic Region (Cape) and the Southwest Australia Floristic Region (SWA). We present (a) distributions of
- 512 roughness values (Equation (1)) for example variables from each broad category of the environment concerned.
- 513 (b) The common language effect size (CLES; see text) of Cape versus SWA roughness values is shown for all
- variables, grouped by broad categories of the environment, describing differences in the distributions of Cape

- and SWA roughness values. We used Mann-Whitney U-tests to assess differences in these distributions.
- Non-significant differences ($P_U > 0.05$) are denoted as such ("NS"). Note, U-tests were performed using only
- a random set of 5000 cells at the 0.05-degree-scale, as the U-test as implemented in R cannot handle more than
- 518 that many values to compare.
- 519 Figure 2: Species turnover, described in two forms ((a) mean Jaccard distance between QDS in each HDS
- 520 (\overline{J}_{QDS}) , (b) additively defined turnover $(T_{HDS}, \text{ Equation (2)})$ as a proportion of HDS richness (S_{HDS})),
- compared between the Cape and SWA. Mann-Whitney U-tests between the Cape and SWA distributions of
- 522 $\overline{J}_{\mathrm{QDS}}$ and T_{HDS} yielded significant differences.
- 523 Figure 3: Relative influence of environmental variables (including heterogeneity variables—prefixed with "R")
- 524 in boosted regression tree (BRT) model predictions in the Greater Cape Floristic Region (Cape, a-c) and the
- 525 Southwest Australia Floristic Region (SWA, b-d) of vascular plant species richness at the (b,e) QDS-scale
- 526 $(=\overline{S}_{QDS})$, (a,d) HDS-scale $(=S_{HDS})$ and (c,f) turnover $(=\overline{J}_{QDS})$. All BRT-models were permitted to fit
- 527 three-way interactions between environmental variables. Points denote the mean contribution of an
- environmental variable to model-predictions across the 1000 replicate BRT-models for that region/scope.
- Horizontal ticks denote the mean for the 999 permuted BRT-models. Standard deviations above and below
- these means are shown with vertical lines. Note, in the case of the replicate, standard deviations are so small
- 531 such that the vertical lines are obscured by the points. Colours represent the general category of the
- environment to which a variable belongs (keyed), as in Figure 1b. Left-most piecharts inset in each panel
- display the same information. Right-most piecharts group contributions according to whether a variable was
- absolute or roughness-transformed (keyed). F-statistics inset are for one-way ANOVAs of differences in
- variables' relative influences—for both the replicate ($F_{\rm rep.}$) and permuted ($F_{\rm prm.}$) BRT-models.
- Figure 4: Distributions of three measures of boosted regression tree (BRT) model performance: (a) the number
- of trees in the model nt, (b) $R_{\rm pseudo}^2$ (Equation (4)), (c) $R_{\rm E-O}^2$ (see text). These measures are presented for the
- 538 six sets of permuted (pale bars) and six sets of replicate BRT-models (dark bars) as in Figure 3, coloured
- 539 according to the region of interest as in Figures 1a and 2. In all cases, replicate BRT-models almost entirely
- out-rank the permuted models in terms of performance (Table 2) and the Greater Cape Floristic Region (Cape)
- and Southwest Australia Floristic Region (SWA) models had significantly different values for each metric
- 542 (Table 3). Note, the actual differences between Cape and the SWA models' values is not realistically significant
- 543 in some cases (e.g. the difference in nt between the Cape and SWA QDS richness models is statistically
- significant, but are observedly so similar as not to affect interpretation).

Figure 5: Differences in the rankings of environmental variables' (including heterogeneity variables) relative 545 influences on boosted regression tree (BRT) model predictions of vascular plant species richness and turnover 546 547 in (a) the Greater Cape Floristic Region (Cape) and (b) Southwest Australia Floristic Region (SWA) (as in Figure 3). Each point represents an environmental variable's rank in BRT-model importance, decreasing in 548 importance from left to right. Rankings used here are the same as that of the average relative influence for 549 variables across replicate BRT-models, presented in Figure 3. Coloured lines connect points representing the 550 same environmental variable. Points' outlines are coloured according to the general category of the 551 environment (keyed) to which a variable belongs, as in Figuress 1b and 3, while points' centres are coloured 552 according to whether a variable was roughness-transformed or not. The comparisons of variables' rankings of 553 interest are between QDS- and HDS-scale richness (= \overline{S}_{QDS} and S_{HDS} respectively; rows nos. 1 and 2) and 554 between HDS-scale richness and turnover (= \overline{J}_{QDS}) (rows nos. 2 and 3). Statistics (Δ - and P-values) inset at 555 the top and bottom of each panel refer to these comparisons respectively. Δ -values represent the average 556 absolute difference in ranks across variables between two models' rankings. The associate P-value results from 557 ranking the observed Δ -values against 999 Δ -values based on random permutations of variables' rankings 558 (SI1), such that more significant P-values denote rankings more similar than would be expected by chance. 559 Figure 6: Marginal effects of environmental conditions and heterogeneity on vascular plant species richness at 560 the QDS-scale (= \overline{S}_{QDS} ; a,d), HDS-scale (= S_{HDS} ; b,e) and turnover (= \overline{J}_{QDS} ; c,f) in response variables in 561 the Greater Cape Floristic Region (Cape; a-c) and Southwest Australia Floristic Region (SWA; d-f) following 562 boosted regression tree (BRT) modelling. Marginal effect functions presented are derived from a representative 563 BRT-model from the set of replicate BRT-models (for each of the six modelling cases) (see SI regarding how 564 representative BRT-models were selected). Marginal effects represent the effect of a predictor variable when all 565 other predictors are set at their means. Marginal effect functions are shown for environmental variables that 566 contributed $\geq 10\%$ to a model's predictions. Functions are coloured as keyed, with solid lines representing 567 absolute environmental variables and dotted representing heterogeneity variables ("rough"). Environmental 568 variables were all rescaled here such as to be centred on zero (i.e. Z-transformed), facilitating comparison of 569 functions' forms. 570

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654 Biosketches

- Ruan van Mazijk is a Masters student interested in phylogenetic systematics, macro-ecology, comparative
- work and plant functional ecology.
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- 658 G. Anthony Verboom

659 Author contributions

- 660 MDC and GAV conceived the study question, which RVM investigated under their supervision for his BSc
- 661 Hons project. The analyses and programming work were largely devised by RVM, with input from the other
- authors, and was carried out by RVM. RVM wrote the first draft of the manuscript and all authors contributed
- 663 equally thereafter.

664 Tables

Table 1: Georeferenced vascular plant species occurrence and environmental data sources used in this study. Data were acquired for the Cape and SWA regions, with the temporal extent of data products used described where applicable. Abbreviations are as follows: MAP, mean annual precipitation; PDQ, precipitation in the driest quarter; CEC, cation exchange capacity.

Variable	Source	Temporal extent	Citation
Plant species occurrences	GBIF	TODO	??, ??
Elevation	SRTM v2.0		??
NDVI	MODIS (MOD13C2)	Feb. 2000 to Apr. 2017	??
Climatic variables			
Surface temperature	MODIS (MOD11C3)	Feb. 2000 to Apr. 2017	??
MAP	CHIRPS v2.0	Jan. 1981 to Feb. 2017	??
PDQ	CHIRPS v2.0	Jan. 1981 to Feb. 2017	??
Soil variables			
CEC	SoilGrids250m (CECSOL M 250m)		??
Clay	SoilGrids250m (CLYPPT M 250m)		
Soil C	SoilGrids250m (OCDENS M 250m)		
рН	SoilGrids250m (PHIKCL M 250m)		

Table 2: Average percentile-ranks for BRT-model performance measures (nt, $R_{\rm pseudo}^2$ (Equation (4)), $R_{\rm E-O}^2$) of 1000 replicate BRT-models relative to 999 BRT-models fit to permuted datasets. Ranks approaching one indicate that a set of replicate BRT-models had greater values than the permuted models.

Model	nt	R_{pseudo}^2	R_{E-O}^2		
QDS-richness					
GCFR	1.000	1.000	1.000		
SWAFR	1.000	1.000	1.000		
HDS-richness					
GCFR	0.987	1.000	0.988		
SWAFR	1.000	1.000	1.000		
HDS-turnover					
GCFR	0.977	0.992	0.979		
SWAFR	0.997	1.000	1.000		

Table 3: Estimated differences between replicate Cape and SWA BRT-models' performance measures (nt, $R_{\rm pseudo}^2$ (Equation (4)), $R_{\rm E-O}^2$) following t-tests. Positive values indicate that the Cape models had greater values. In all cases, the Cape and SWA had highly significantly different values for these quality measures (P < 0.0001).

Model	nt	$R_{\rm pseudo}^2$	$R_{\mathrm{E-O}}^2$
QDS-richness	542.938	0.063	-0.005
HDS-richness	-808.994	-0.064	-0.233
HDS-turnover	-997.045	-0.052	-0.296

665 Figures

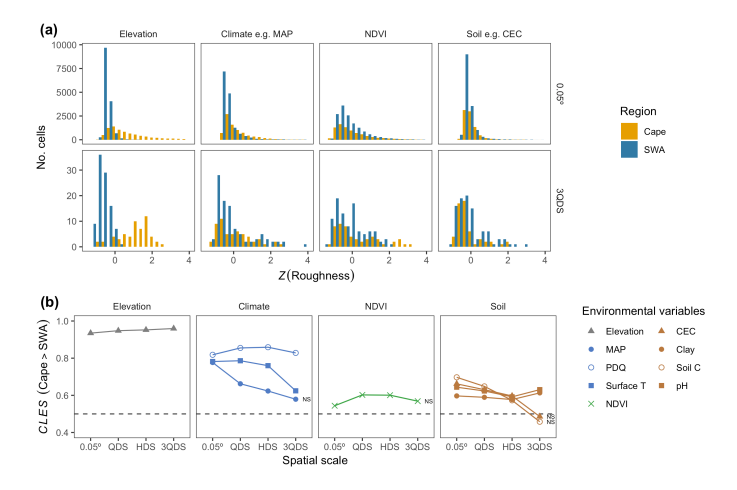


Figure 1: Comparisons of different types of environmental heterogeneity between the the Greater Cape Floristic Region (Cape) and the Southwest Australia Floristic Region (SWA). We present (a) distributions of roughness values (Equation (1)) for example variables from each broad category of the environment concerned. (b) The common language effect size (CLES; see text) of Cape versus SWA roughness values is shown for all variables, grouped by broad categories of the environment, describing differences in the distributions of Cape and SWA roughness values. We used Mann-Whitney U-tests to assess differences in these distributions. Non-significant differences ($P_U > 0.05$) are denoted as such ("NS"). Note, U-tests were performed using only a random set of 5000 cells at the 0.05-degree-scale, as the U-test as implemented in R cannot handle more than that many values to compare.

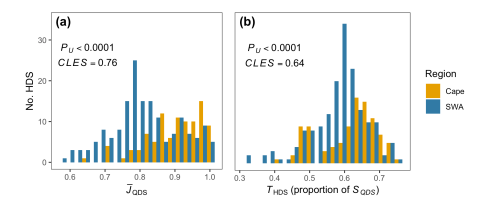


Figure 2: Species turnover, described in two forms ((a) mean Jaccard distance between QDS in each HDS $(\overline{J}_{\text{QDS}})$, (b) additively defined turnover (T_{HDS} , Equation (2)) as a proportion of HDS richness (S_{HDS})), compared between the Cape and SWA. Mann-Whitney U-tests between the Cape and SWA distributions of $\overline{J}_{\text{QDS}}$ and T_{HDS} yielded significant differences.

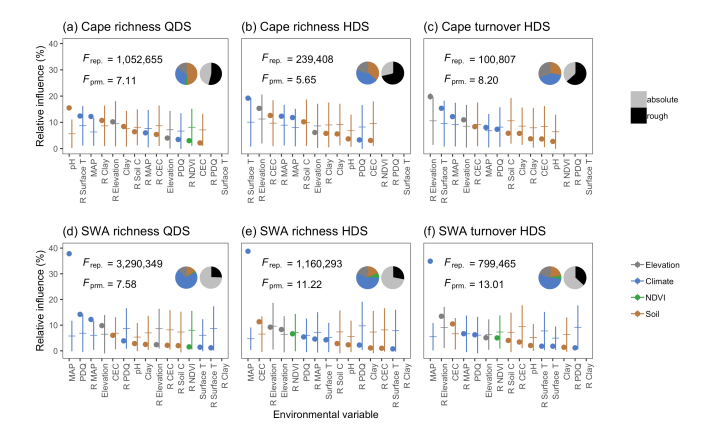


Figure 3: Relative influence of environmental variables (including heterogeneity variables—prefixed with "R") in boosted regression tree (BRT) model predictions in the Greater Cape Floristic Region (Cape, a–c) and the Southwest Australia Floristic Region (SWA, b–d) of vascular plant species richness at the (b,e) QDS-scale (= $\overline{S}_{\text{QDS}}$), (a,d) HDS-scale (= S_{HDS}) and (c,f) turnover (= $\overline{J}_{\text{QDS}}$). All BRT-models were permitted to fit three-way interactions between environmental variables. Points denote the mean contribution of an environmental variable to model-predictions across the 1000 replicate BRT-models for that region/scope. Horizontal ticks denote the mean for the 999 permuted BRT-models. Standard deviations above and below these means are shown with vertical lines. Note, in the case of the replicate, standard deviations are so small such that the vertical lines are obscured by the points. Colours represent the general category of the environment to which a variable belongs (keyed), as in Figure 1b. Left-most piecharts inset in each panel display the same information. Right-most piecharts group contributions according to whether a variable was absolute or roughness-transformed (keyed). F-statistics inset are for one-way ANOVAs of differences in variables' relative influences—for both the replicate ($F_{\text{rep.}}$) and permuted ($F_{\text{prm.}}$) BRT-models.

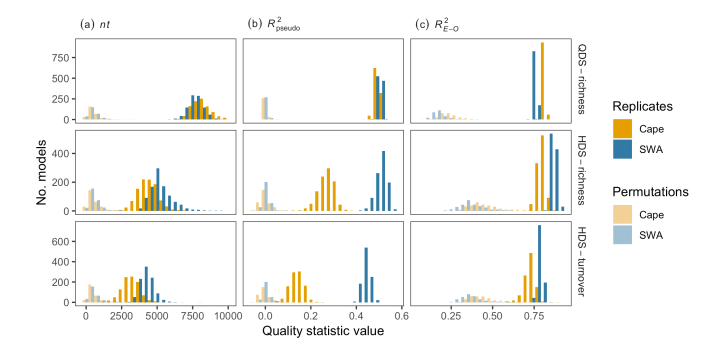


Figure 4: Distributions of three measures of boosted regression tree (BRT) model performance: (a) the number of trees in the model nt, (b) $R_{\rm pseudo}^2$ (Equation (4)), (c) $R_{\rm E-O}^2$ (see text). These measures are presented for the six sets of permuted (pale bars) and six sets of replicate BRT-models (dark bars) as in Figure 3, coloured according to the region of interest as in Figures 1a and 2. In all cases, replicate BRT-models almost entirely out-rank the permuted models in terms of performance (Table 2) and the Greater Cape Floristic Region (Cape) and Southwest Australia Floristic Region (SWA) models had significantly different values for each metric (Table 3). Note, the actual differences between Cape and the SWA models' values is not realistically significant in some cases (e.g. the difference in nt between the Cape and SWA QDS richness models is statistically significant, but are observedly so similar as not to affect interpretation).

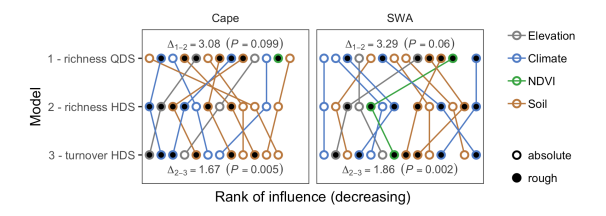


Figure 5: Differences in the rankings of environmental variables' (including heterogeneity variables) relative influences on boosted regression tree (BRT) model predictions of vascular plant species richness and turnover in (a) the Greater Cape Floristic Region (Cape) and (b) Southwest Australia Floristic Region (SWA) (as in Figure 3). Each point represents an environmental variable's rank in BRT-model importance, decreasing in importance from left to right. Rankings used here are the same as that of the average relative influence for variables across replicate BRT-models, presented in Figure 3. Coloured lines connect points representing the same environmental variable. Points' outlines are coloured according to the general category of the environment (keyed) to which a variable belongs, as in Figuress 1b and 3, while points' centres are coloured according to whether a variable was roughness-transformed or not. The comparisons of variables' rankings of interest are between QDS- and HDS-scale richness (= $\overline{S}_{\text{QDS}}$ and S_{HDS} respectively; rows nos. 1 and 2) and between HDS-scale richness and turnover (= $\overline{J}_{\text{QDS}}$) (rows nos. 2 and 3). Statistics (Δ - and P-values) inset at the top and bottom of each panel refer to these comparisons respectively. Δ -values represent the average absolute difference in ranks across variables between two models' rankings. The associate P-value results from ranking the observed Δ -values against 999 Δ -values based on random permutations of variables' rankings (SI1), such that more significant P-values denote rankings more similar than would be expected by chance.

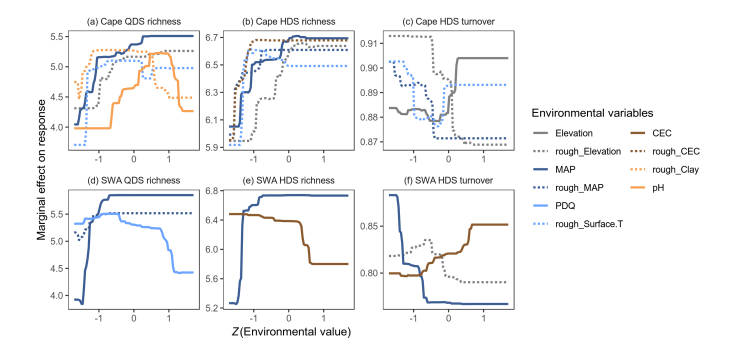


Figure 6: Marginal effects of environmental conditions and heterogeneity on vascular plant species richness at the QDS-scale (= $\overline{S}_{\text{QDS}}$; a,d), HDS-scale (= S_{HDS} ; b,e) and turnover (= $\overline{J}_{\text{QDS}}$; c,f) in response variables in the Greater Cape Floristic Region (Cape; a–c) and Southwest Australia Floristic Region (SWA; d–f) following boosted regression tree (BRT) modelling. Marginal effect functions presented are derived from a representative BRT-model from the set of replicate BRT-models (for each of the six modelling cases) (see SI regarding how representative BRT-models were selected). Marginal effects represent the effect of a predictor variable when all other predictors are set at their means. Marginal effect functions are shown for environmental variables that contributed $\geq 10\%$ to a model's predictions. Functions are coloured as keyed, with solid lines representing absolute environmental variables and dotted representing heterogeneity variables ("rough"). Environmental variables were all rescaled here such as to be centred on zero (i.e. Z-transformed), facilitating comparison of functions' forms.