

Seaweeds in Two Oceans: Beta-diversity

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2 ABSTRACT

3 Several species assembly mechanisms have been proposed to structure ecological communities. We assess
4 the biogeography of seaweeds along 2,900 km of South Africa's coastline in relation to a thermal gradient
5 produced by the Agulhas Current, and contrast this with the environmental structure created by the Benguela
6 Current. We subdivided the coastline into 'bioregions' to examine the regional patterning. To investigate the
7 assembly mechanisms, we decomposed Sørensen's β -diversity into 'turnover' (β_{sim}) and 'nestedness-resultant'
8 (β_{sne}) dissimilarities, and used distance-based redundancy analysis (db-RDA) to relate them to the Euclidian
9 thermal difference, d_E , and geographical distance. Moran's eigenvector maps (MEM) were used as an additional
10 set of spatial constraints. Variation partitioning was then used to find the relative strengths of thermal and
11 spatially-structured thermal drivers. Spatial and environmental predictors explained 97.9% of the total variation
12 in β_{sim} and the thermal gradient accounted for 84.2% of this combined pool. β_{sim} was the major component
13 of overall β -diversity in the Agulhas Current region, suggesting niche influences (environmental sorting) as
14 dominant assembly process there. The much weaker thermal gradient in the Benguela Current-influenced region
15 resulted in a high amount of β_{sne} that could indicate neutral assembly processes. The intensification of upwelling
16 during the mid-Pliocene 4.6–3.2 Ma (i.e. historical factors) were likely responsible for setting up the strong
17 disjunction between the species-poor west coast and species-rich south and east coast floras, and this separation
18 continues to maintain two systems of community structuring mechanisms in the Atlantic and Indian Ocean
19 influenced sides of South Africa.

20 Keywords: beta-diversity, species assembly, seaweed, macroalgae, turnover, nestedness-resultant, Benguela Current,
21 South Africa

22 1 Introduction

23 The assembly processes that structure biodiversity across a range of scales form the theme of macroecology
24 (Chave, 2013). This paper deals with the composition of seaweed assemblages along the ~2,900 km South
25 African coastline, including the identification, description and explanation for the spatially-structured
26 patterns at scales from 100s to 1,000s of kilometers. Inshore conditions along this coastline range from
27 cool through warm temperate to fringe tropical (Bolton and Anderson, 2004; Bolton et al., 2004), and are
28 influenced by two major ocean currents in two oceans that set up a strong thermal gradient along the shore

29 (Smit et al., 2013). We investigate how a gradient in seaweed community composition may have arisen in
30 response to this temperature gradient (e.g. Qian and Ricklefs, 2007), and the extent to which deterministic,
31 niche-based species assembly processes could have contributed towards the assembly of the seaweed flora
32 of the region.

33 Biodiversity may be viewed in terms of α -, γ - and β -diversity, with the former two referring to 'local'
34 and 'regional' diversity, respectively. β -diversity as defined by Whittaker (1960, 1972) is a measure of
35 variation in species composition from place to place and is comprised of two processes (Baselga, 2012):
36 the replacement of species independently of the difference in species richness (called 'turnover', β_{sim}) and
37 a term that considers this difference in species richness called ('nestedness-resultant', β_{sne}). The relative
38 contribution of these two processes is not immediately evident from species dissimilarities, and yet such
39 considerations should be implicit in macroecological studies. The ability to decompose β -diversity into
40 turnover and nestedness-resultant components has become possible within the last decade (Baselga, 2010),
41 but the use of this form of β -diversity partitioning has not yet widely permeated the phycological literature
42 (or even in marine studies more broadly; Anderson et al., 2013). Cognisance of spatial scaling should also
43 be deeply rooted in the study of biodiversity more generally, but such questions have only recently begun
44 to be addressed (Barton et al., 2013).

45 Since studies of β -diversity consider the variation in species assembly processes from place to place, and
46 necessitate an understanding of mechanisms that drive these processes (Davidar et al. 2007), studying β -
47 diversity of adjacent coastal marine bioregions or marine provinces (Spalding et al., 2007) could provide
48 deeper insight into how such processes operate along gradients (Qian and Ricklefs, 2007). Few studies
49 dealing specifically with β -diversity exist for marine biota, especially over spatial areas in excess of 1,000s
50 of kilometers. Recent examples include studies on fish distribution (Zintzen et al., 2011; Anderson et al.,
51 2013), the biogeography of macroalgae along 6,600 km of Australian coastline (Leaper et al., 2011), and a
52 study on deep-sea bivalves (McClain et al., 2011). All four have β -diversity as a primary interest. However,
53 macroecological studies in the terrestrial realm have yielded the most comprehensive insights into the
54 drivers responsible for assembling species into communities (Whittaker, 1960; Davidar et al., 2007; Qian
55 and Ricklefs, 2007; Soininen et al., 2007). What these studies showed is that, perhaps universally, high
56 rates of species turnover are associated with steep environmental gradients, such as those occurring along
57 mountain slopes or across latitudes. In the ocean, the biggest driver latitudinally seems to be temperature
58 (Tittensor et al., 2010; Stuart-Smith et al., 2017; Straub et al., 2016); other drivers such as nutrients, salinity,
59 turbidity, wave action and photoperiod may also structure species composition, but these are likely to
60 depend more on the regional context, and some, such as nutrients (e.g. Waldron and Probyn, 1992), are
61 often strongly correlated with a more distal variable, such as temperature.

62 Recent decades have seen temperature emerging as a readily available environmental driver that's able
63 to resolve the ocean's heat content across a multitude of temporal and spatial scales. Temperature is also
64 very useful as a predictor of species range limits due to it being one of the key output variables of coupled
65 global ocean-atmosphere circulation models used to project the physical milieu of the future Earth that
66 has experienced anthropogenic climatic change (Church et al., 2013), hence making it the focus of our
67 present study. The controlling effect of seawater temperature on the survival and reproduction of benthic
68 organisms and patterns in the evolution and ecology of biological assemblages at regional scales is well
69 known (van den Hoek, 1982b; Breeman, 1988; Blanchet et al., 2008; Broitman et al., 2008; Byrne et al.,
70 2009; Verbruggen et al., 2009; Wieters et al., 2009; Couce et al., 2012; Potts et al., 2014). It influences the
71 species composition of the biota associated with various thermal zones, which forms patterns on global,
72 regional and local scales (Tittensor et al., 2010; Spalding et al., 2012). At the global scale, ocean currents and

73 well-characterised latitudinal solar heat flux gradients maintain the ocean's thermal regime. At local scales
74 (<10 km) at the land/sea margin, however, additional less well-known physical phenomena contribute to
75 thermal patterns and dynamics that may differ markedly from those at the mesoscale (*i.e.* spatial scales
76 >50 km), and the different properties of the temperature regime which set up the biogeographical patterns
77 are less well understood there — this is especially true at regional and local scales near the land where
78 satellite data poorly reflect reality (Smit et al., 2013). Most studies that recognise temperature as a major
79 driver of species distribution take the annual mean temperature (Tittensor et al., 2010); fewer recognise the
80 importance of variability and range (*e.g.* Couce et al., 2012; Tyberghein et al., 2012). Detailed laboratory
81 culture studies with seaweed species, for example, closely link biogeographical distribution limits with
82 maximum and minimum monthly mean temperatures (van den Hoek, 1982b; Breeman, 1988). The time-
83 integrated 'thermal environment' is an amalgam of various statistical properties derived from a multitude
84 of instantaneous temperature recordings, and it is pertinent to question if one or a few of these properties
85 have an overriding imprint on the species assembly.

86 Understanding the processes influencing species turnover and geographical range is of practical
87 importance to spatial biodiversity planning. This is especially relevant in today's warming world because
88 it will allow us to project future biodiversity responses. Focusing conservation efforts on areas of high
89 β -diversity will ensure the preservation of a diversity of species and their environmental niches. Peaks
90 in β -diversity along ecoclines may indicate boundaries between bioregions, highlighting regions where
91 communities are poised at their environmental limits, thus aligning with long-term monitoring surveys
92 that aim to project ecosystem responses to climatic change.

93 Using a detailed data set of seaweed presence and absence records coupled with coastal *in situ* seawater
94 temperature climatologies that are able to resolve the coastal zone, we investigated the thermal properties
95 and species composition of 58 coastal sections spaced around the South African coastline. The coastline is
96 broadly influenced by two major ocean currents, the Benguela Current and the Agulhas Current. One is
97 an eastern boundary upwelling system and defines a cool temperate environment, and the other a western
98 boundary current driving meridional transport of sub-tropical water towards the tip of Africa. We were
99 primarily interested in establishing how these ocean currents, of which the effect at the coast can readily be
100 measured as gradients in thermal properties, may have influenced the species assembly processes operating
101 along an approximately 2,900 km long coastline. To do so, we first link matrices of species dissimilarity to
102 environmental distance matrices and Moran's eigenvector maps (MEM) using distance-based redundancy
103 analysis, and then apply variance partitioning (Peres-Neto and Legendre, 2010) to determine the relative
104 contributions of thermal metrics and other spatially-organised drivers of seaweed community composition
105 (as seen in the β -diversity components, β_{sim} and β_{sne}). Second, we examine how the thermal and spatial
106 structuring agents operate at smaller spatial scales within marine provinces of the region by undertaking
107 a more detailed analysis of the various distance matrices, and also ask which of the thermal properties
108 we examined were most influential in setting up the patterns that emerged. Our findings are congruent
109 with existing knowledge (*e.g.* Stephenson, 1948; Lombard et al., 2004; Spalding et al., 2007) and add a
110 more nuanced understanding of the processes responsible for structuring the seaweed biodiversity in the
111 sub-region.

112 2 Methods

113 2.1 Species data and explanatory variables

114 We used three sets of data in this analysis. The first comprises distribution records (presence/absence) of
115 846 macroalgal species belonging to the Divisions Ochrophyta, Rhodophyta and Chlorophyta within each
116 of 58 × 50 km-long sections of the South African coast (mentioned in **bold** font in the text and listed in
117 Appendix A, Table1). The *seaweed data* represent ca. 90% of the known seaweed flora of South Africa in
118 the intertidal and shallow subtidal, but excludes some very small and/or very rare species for which data
119 are insufficient. The data are from verifiable literature sources and our own collections, assembled from
120 information collected by teams of phycologists over three decades (Bolton and Stegenga, 2002; Bolton,
121 1996, 1986; De Clerck et al., 2005; Stegenga et al., 1997).

122 The second is a dataset of *in situ* coastal seawater temperatures (Smit et al., 2013) derived from daily
123 measurements over up to 40 years. The *thermal data* set was used as the first set of explanatory variables. The
124 following statistical properties were entered into the analysis: the means for the year (*annMean*), February
125 (*febMean*, Austral summer) and August (*augMean*, Austral winter); the annual standard deviation (SD)
126 around the mean for the year, February and August (*annSD*, *febSD* and *augSD*, respectively); and the annual
127 thermal range between the mean temperature of the warmest and coldest months (*annRange*), and the
128 mean range of February and August temperatures (*febRange* and *augRange*).

129 The third set of explanatory variables was generated to represent the spatial connectivity among
130 coastal sections. Because of the strong environmental gradients along the shore, species community
131 composition should be spatially organised (including the modelled spatially structured environmental
132 variables, exogenous spatial variables and autocorrelation), and this was accounted for in the analysis. To
133 this end we produced a section–section connectivity matrix based on a minimum spanning tree (MST)
134 topology, starting from a geographical distance matrix. This topology focuses on relationships between
135 neighbouring sections and discards connections that are further away. In the case of the coastline data,
136 sections are connected only to other sections that are below a certain truncation distance. This largely
137 resulted in a ‘string-of-beads’ series of section–section connections; for example, Section 4 is directly
138 connected to only Sections 3 and 5; sections at the termini of the coastal ‘string’ of sections are each
139 connected to only one other section. We generated Moran’s eigenvector maps (MEM; Dray et al., 2006,
140 2012) from this connectivity matrix through a principal coordinates analysis (PCoA) using the PCNM
141 function of the PCNM package in R 3.3.3 (R Core Team, 2017), and kept the MEMs with positive spatial
142 correlation. The MEMs are completely orthogonal and represent the spatial structures over the full range
143 of scales from 50 to 2,900 km. Large eigenvectors represent broad spatial scales while smaller ones cover
144 finer features. The *spatial data* were used as the second set of explanatory variables in multiple regression
145 type analyses (Dray et al., 2012). Details and code are provided in Appendix B and may be accessed online
146 at https://github.com/ajsmits/Seaweeds_in_Two_Oceans.git.

147 2.2 Distance-based redundancy analysis and variance partitioning

148 Using the distance-based redundancy analysis (db-RDA; Minchin, 1987) implemented with the **capscale**
149 function of the R package, **vegan** (Oksanen et al., 2016), we explored the role of the thermal and spatial
150 descriptors in structuring the seaweed communities across the 58 coastal sections. To represent the biotic
151 data, we decomposed Sørensen’s dissimilarity ($\beta_{\text{sør}}$) into its ‘nestedness-resultant’ (β_{sne}) and ‘turnover’
152 (β_{sim}) components (Baselga, 2010) using the **betapart** package (Baselga et al., 2013). This approach was

necessary because β -diversity is strongly coupled with α -diversity, and it allowed us to make inferences about the possible drivers of β -diversity. Turnover refers to processes that cause communities to differ due to species being lost and/or gained from section to section, *i.e.* the species composition changes between sections without corresponding changes in α -diversity. The nestedness-resultant component implies processes that cause species to be gained or lost, and the community with the lowest α -diversity is a subset of the richer community.

The assessment of the biotic ordination within the context of the underlying environmental properties proceeded using the environmental variables' z-scores. To determine the descriptors that best describe the patterns in the seaweed dissimilarity data, we first applied full (global) db-RDAs using the complete sets of thermal and spatial variables, separately for each set. Using the forward selection procedure implemented in the **packfor** package for R (Blanchet et al., 2008), we reduced the number of variables in each set and retained only those that best fit the biotic data. Forward selection prevents the inflation of the overall type I error and reduces the number of explanatory variables used in the final model, which improves parsimony. The reduced set of thermal variables retained collinear variables (Graham, 2003), which were identified and removed using variance inflation factors (VIF; Dormann et al., 2013). This was not necessary for the MEMs as they are orthogonal by definition. The remaining significant orthogonal thermal and spatial variables were then regressed with β_{sim} and β_{sne} and final db-RDA models produced. The computation of db-RDAs was followed by permutation tests of the adjusted R^2 to assess the significance of constraints (thermal and spatial descriptors). Lastly we undertook variance partitioning (Peres-Neto et al., 2006; Peres-Neto and Legendre, 2010) between the environmental (thermal) and spatial predictors using the **varpart** function in the **vegan** package. Refer to Appendix B for more information about the methodology and for the R code underlying the analysis.

2.3 Analyses of dissimilarity and distance matrices

We then explored regional patterns in β -diversity. Because connectivity between sections is constrained by their location along the shore and thus direct distances between sections do not apply, the total distance between a pair of arbitrary sections is the cumulative sum of the great circle distances between each consecutive pair of intervening sections along the coast (this is in fact encapsulated by the connectivity matrix used in the PCNM analysis, above). Plots showing the relationship of β_{sim} and β_{sne} with distance are limited because they do not provide a geographical context. To overcome this problem, we used a 'network graph' to show spatial relationships in regional species dissimilarity. See Appendix C for details and R code.

The last step of our analysis was applied to the four bioregions recognised for South Africa (Bolton and Anderson, 2004), namely the Benguela Marine Province (BMP; 1–17), the Benguela-Agulhas Transition Zone (B-ATZ; 18–22), the Agulhas Marine Province (AMP; 19–43/44) and the East Coast Transition Zone (ECTZ; 44/45–58). To this end, we calculated an Euclidian distance matrix that encapsulated all pairwise differences between coastal sections for each of the thermal metrics highlighted in the db-RDA (above) using the **vegan** package; these are called thermal differences (d_E) throughout. We then correlated β_{sim} and β_{sne} with geographical distance and d_E and provided matching plots in which the four bioregions were colour-coded to discern bioregional affiliations and differences. Together these analyses were able to capture β -diversity at two spatial scales: among sections within bioregions, and among all sections for the whole country.

194 3 Results

195 3.1 The thermal environment and species richness

196 South Africa's annual mean coastal water temperature ranged from $12.0 \pm 0.9^\circ\text{C}$ (mean \pm SD) at its north-
197 western limit near the Namibian border (1) to $24.0 \pm 1.9^\circ\text{C}$ on the east coast near the Mozambican border
198 (58) (Fig. 1). The global latitudinal gradient of diminishing temperature with increasing latitude was seen
199 only along the east coast where the annual mean temperature decreased from *ca.* 24.5°C near 58 to 17.5°C
200 around 39. The alongshore thermal gradient for this 950 km stretch of coastline was *ca.* 0.7°C per 100 km,
201 with steeper gradients near 54. The latitudinal gradient largely reversed in direction along the west coast
202 (1–16), *i.e.* temperatures became slightly cooler further north.

203 **Figure 1 near here.**

204 On average, these data indicated an increase in inshore annual mean temperatures from west to east
205 (1–58) of $12.1\text{--}24.4^\circ\text{C}$ (range: 12.3°C). In February the thermal range was 13.7°C , while in August it was
206 10.5°C . In August the west–east temperature transition was smooth whereas in February substantial warm
207 fluctuations in the mean monthly temperature were observed in embayments such as 13 and the False Bay
208 sections from 17–18, and 28 and some sections around 35/36 and Algoa Bay from 34–36. In summer the
209 mean monthly temperature gradient steepened between 19–28, and thereafter decreased eastwards along
210 the coast from 33–34.

211 The number of species within the BMP was low (Fig. 1, inset), and many northern sections had fewer
212 than 150 species, and it rose significantly in the warmer section around 12/13, and the southern sections
213 around the Cape Peninsula (16/17). Thereafter, richness increased markedly in the B-ATZ, the AMP and
214 the ECTZ. The highest number of species in any one section was 340 (Section 39 near the eastern end of
215 the AMP).

216 3.2 Environmental correlates of seaweed diversity

217 db-RDA, forward selection and the assessment of VIF retained *augMean*, *febRange*, *febSD* and *augSD* as
218 the most parsimonious descriptors of β_{sim} with an adjusted R^2 of 0.885, explaining 89.8% of the variation
219 (global permutation test on final model: d.f. = 4, $F = 110.16$, $p = 0.001$). The model consisted of two
220 significant canonical axes: CAP1 and CAP2 explained 73.3% and 14.9% of the variation, respectively. The
221 biplot scores (vectors) showed that *augMean* was heavily loaded along CAP1 and the metrics related to
222 variation around the mean, *i.e.* *febRange* and *febSD*, strongly influenced β_{sim} along CAP2 (Fig. 2). Plots
223 of the 'lc' scores on geographic axes are given in Fig. 3. The scores representing CAP1 increased gradually
224 along the shore from west to east, reflecting the pervasive influence of *augMean* as coastal sections changed
225 from cool to warm temperate through to sub-tropical thermal regimes. CAP2 site scores were lowest along
226 the southern sections. β_{sne} was only influenced by *annMean* (Fig. 2) along CAP1 that explained 20.3% of
227 the total variation ($R^2 = -0.140$, d.f. = 1, $F = -6.018$).

228 **Figure 2 near here.**

229 The db-RDA analysis procedure retained 17 significant MEMs that fully encapsulated the spatial
230 dependence within β_{sim} (d.f. = 18, $F = 84.055$, $p = 0.001$), resulting in an adjusted R^2 of 0.963 and accounting
231 for all of the variation. Fifteen canonical axes were produced of which five were significant. The first one
232 alone explained 77.1% of variance with the remaining axes accounting for *ca.* 1% or less of the inertia.
233 MEM2, MEM3 and MEM5 were most strongly loaded along CAP1 (Fig. 3). These MEMs caused the

sections belonging with the BMP to separate out from all other sections, and also for the northern sites of the ECTZ to diverge from the sections and bioregions further south. The scales of spatial dependence captured by these MEMs could all be considered to be broad-scaled. Two canonical axes comprised of two significant MEMs were able to explain 87.3% of the variation in β_{sne} (R^2 of 0.437, d.f. = 4, $F = 12.06$, $p = 0.001$; Fig. 3). CAP1 consumed 79.0% of the inertia due to MEM1 and MEM5.

Figure 3 near here.

The partitioning of the variance associated with the seaweed community along the coast (Table 1) was explained jointly by the thermal and spatial variables selected in the preceding db-RDAs. Combining the thermal and spatial predictors (fractions [E+S]) allowed the model to capture 97.9% of the total β_{sim} variance ($F = 191.56$, $p = 0.001$), with a residual variance of 2.1%. The thermal variables on their own (*i.e.* those that are spatially unstructured; fraction [E|S]) were able to account for only 1.8% of the total variation ($F = 20.506$, $p = 0.001$), but including some spatially structured thermal properties, [E], raised the proportion of explained variation to 84.2% ($F = 110.16$, $p = 0.001$). Pure spatial patterning (*i.e.* in the absence of temperature influences, perhaps with exogenous environmental influences or autocorrelation; fraction [S|E]) drove 13.7% of the species variation ($F = 23.649$, $p = 0.001$), and adding some thermal influences together with spatial descriptors, [S], increased this to 96.1% ($F = 80.731$, $p = 0.001$). Turning now to β_{sne} , we see that our explanatory variables were less successful in capturing the variation. The spatial variables, [S], and the spatial plus thermal variables, [E+S], were able to account for 67.8% ($F = 12.06$, $p = 0.001$) and 71.4% ($F = 9.077$, $p = 0.001$) of the variation, respectively.

Table 1 near here.

3.3 Pairwise dissimilarities

Network graphs show the spatial relationships of β_{sim} (Fig. 4). β_{sim} clearly highlighted the effect of the sharp change in α -diversity between the BMP and the B-ATZ. Sections in the B-ATZ retained similarities with sections as far east as 33 within the AMP. Eastwards from 30 similarities with sections within the ECTZ became apparent, but they generally did not extend past 43. Aside from a very low similarity with sections at the eastern extent of the AMP, the ECTZ sections retained similarities with sections within the same biogeographical province only over very short distances, and again it highlighted the high β -diversity in this region.

Figure 4 near here.

The overall and regional mean values for the three measures of pairwise β -diversity are presented in Table 2. The overall Sørensen β -diversity ($\beta_{\text{sør}}$, 0.496 ± 0.287) was larger than that of the bioregions, and only a small fraction of it was comprised of nestedness-resultant β -diversity. Of the four bioregions, the ECTZ had the highest $\beta_{\text{sør}}$ (0.259 ± 0.157). At the bioregional scale, it is important to note that the nestedness-resultant component was about four times larger for the BMP ($\beta_{\text{sne}}/\beta_{\text{sør}} = 0.581$) than that of the other three ($\beta_{\text{sne}}/\beta_{\text{sør}}$ ranged from 0.097 to 0.170).

Plots of β_{sim} and β_{sne} among all possible section pairs indicated clear differences among the four bioregions in their relationships with geographic and thermal distances (Fig. 5). The BMP showed a weak relationship between β_{sim} and distance ($r^2 = 0.052$; regression statistics in Table 3) since much of the compositional variation between sections in this region was due as much to nestedness-resultant β -diversity as it was to turnover (Fig. 5) as noted above (see $\beta_{\text{sne}}/\beta_{\text{sør}}$ in Table 2). Within the ECTZ (~900 km long coastline) and the B-ATZ (only 170 km long) the rates were moderate at $\beta = 0.079$ ($r^2 = 0.936$)

and high at $\beta = 0.109$ ($r^2 = 0.658$) per 100 km, respectively. In the AMP it was lower at $\beta = 0.029$ per 100 km ($r^2 = 0.834$). Higher rates (β) indicate that communities turned over more rapidly per unit distance of coastline; furthermore, this also provided strong evidence that β -diversity was structured along environmental gradients. β -diversity was less influenced by changes in species numbers between sections in the B-ATZ, AMP and the ECTZ, as these two marine provinces were characterised by a relatively even number of species and hence had low β -diversities attributed to nestedness (Fig. 5). β_{sim} expressed with respect to the *augMean* thermal distance showed similarly steep slopes for three of the bioregions (β s ranging from 0.290 to 0.350, with $r^2 > 0.605$), with that for the BMP about a two-thirds lower (Fig. 5; Table 3). With respect to *febRange*, a significant relationship with β_{sim} existed only for the two transitional areas, the B-ATZ and the ECTZ (r^2 of 0.548 and 0.583, respectively; Fig. 5). The steepness of the relationship between β_{sim} and the latter thermal metric was lower than that seen with *augMean*. Concerning *febSD*, this relationship was steepest for the B-ATZ ($\beta = 0.103$, $r^2 = 0.276$) and then the ECTZ ($\beta = 0.082$, $r^2 = 0.310$; Fig. 5); the same general trend held for *augSD*, with $r^2 = 0.310$ and $r^2 = 0.276$ for the ECTZ and B-ATZ, respectively. The pattern of β_{sne} with geographic and thermal distance was generally significant but very poor (low r^2 -values) and with weak gradients (β) (Fig. 5). The notable outcome there was that it was the BMP where β_{sne} was strongest even though its r^2 values were weak ($r^2 = 0.205$ and 0.164 for the relationship with geographical distance and thermal distance, respectively).

Figure 5 near here.

Table 2 near here.

Table 3 near here.

4 Discussion

In this paper, we considered the drivers of seaweed β -diversity at the scale of bioregions (marine provinces) nested within a ~2,900 km stretch of coastline. We examined β -diversity using a numerical ecological approach that permits assessing its spatial pattern independently of the influence of α -diversity (Baselga et al., 2013), which in our study displays major regional differences. There are fewer species of macroalgae on the Benguela Current dominated west coast, as is also shown by studies on invertebrates and fishes (Thandar, 1989; Turpie et al., 2000). There is a similar species richness per coastal section on the south coast and the east coast. This was not the case up until the early 2000s (Bolton and Stegenga, 2002), when a more restricted data set showed more species per section on the south than east coast. This difference disappeared during the mid-2000s, when major collections on the south coast filled out the species lists available for the region (De Clerck et al., 2005). Continuous sampling of over three decades has now resulted in a robust α -diversity data set, and it is still different from the animal data: fish and invertebrates become more abundant per section on the east coast (towards the tropics) but the seaweeds do not. This matches previously published global seaweed diversity patterns, which suggested that biogeographical seaweed diversity does not peak in the tropics. Globally, seaweed species diversity at a biogeographical scale is highest in warm temperate regions (Bolton, 1994; Santelices et al., 2009). Examining spatial trends in α -diversity on its own offers an incomplete understanding of the structuring of biodiversity at regional scales, and more insight may be obtained by looking at species richness (*i.e.* lists of species, such as which we use in this study) across two scales, *sensu* Whittaker (1972) — *i.e.* α -diversity (species richness at the local scale, such as per coastal section) and γ -diversity (total species richness in all coastal sections). In this broad sense, the two scales of species richness are not independent, and the loss or gain in species from section to

316 section will be seen in β -diversity. In this work, we employ Baselga (2010)'s framework to represent overall
317 β -diversity as two components, one of which excludes the effect of regional differences in α -diversity.

318 With the exception of the some Australian studies (Smale et al., 2010, 2011; Waters et al., 2010; Leaper
319 et al., 2011; Wernberg et al., 2013), one in Europe (Tuya et al., 2012) and another two in the Arabian region
320 (Schils and Wilson, 2006; Issa et al., 2016), studies of this scale and nature have so far been infrequently
321 seen for marine macroalgae. Of the above-mentioned studies, only two specifically considered β -diversity
322 (Leaper et al., 2011; Issa et al., 2016). Our data focus on the taxonomic representivity of a region by
323 aggregating species occurrence records within 50 km long sections of coastline, and are blind to the effects
324 of small-scale habitat heterogeneity (e.g. as seen in Smale et al., 2010). For this reason we cannot infer
325 influences of stochastic processes and other aspects of environmental heterogeneity at scales of <50 km; as
326 such we excluded drivers that may be only of local relevance, such as salinity, turbidity (light), and wave
327 action. Rather, our data emphasise broad biogeographic patterns (Lawton, 1999), with induced spatial
328 dependence (Peres-Neto and Legendre, 2010) emerging as the main species assembly process east of the
329 Cape Peninsula. This points to deterministic niche influences that correlate with environmental drivers that
330 function over a broad spatial scale. These environmental and biotic patterns reflect the nature of the two
331 dominant ocean currents of the region, and we see that β -diversity at the scale of the country is moderately
332 high and generally influenced by processes that cause species turnover (β_{sim}). Our selected environmental
333 drivers comprise mostly thermal properties of seawater with a strong spatial dependence across a broad
334 scale (82.4%) and a smaller amount of unknown non-thermal broad-scale spatial influences (13.7%).
335 Only in the species-poor west coast region is there evidence of neutral assembly processes (e.g. dispersal
336 limitation, and stochastic processes as one might expect within kelp forests), as shown by the higher rates
337 of nestedness-resultant β -diversity (β_{sne}) emerging there. We recognise that some environmental drivers
338 such as nutrients can and do co-vary with temperature, but they were not used in our analysis because
339 they are not as readily measured as temperature, they are not always directly produced by coupled global
340 ocean-atmosphere models and thus have less value as predictive (long-term) variables, and because of the
341 issues arising due to collinearity.

342 The most comprehensive assessment of global marine biogeography (Spalding et al., 2007) places much
343 of South Africa's coast within two realms: Temperate Southern Africa (from southern Angola to around
344 56°S) and the western-most edge of the Western Indo-Pacific Realm (across the Western Indian Ocean to
345 Sumatra). The transition into the Western Indian Ocean realm is just visible in Section 57 where our data
346 show a steady rise to tropical water temperatures that exceed 20°C, between 28.5 and 29°S. Our study region
347 is too restricted to capture the rates of β -diversity change at the transition of realms. Here we would expect
348 higher rates of turnover than we report for the smaller-scale 'Provinces' and transition zones that make up
349 the Temperate Southern African Realm (see below). We would also expect this at higher taxonomic levels,
350 consistent with the definition for a realm (Spalding et al., 2007): this is true for the temperate Southern
351 African seaweed floras, which have high species endemism but very low generic endemism (except for the
352 Fucales, Ochrophyta). We hypothesise that this is because the cool temperate region may be geologically
353 recent (4.6–3.2 Ma, Marlow et al., 2000).

354 Nested within the realms are marine provinces, areas that according to (Spalding et al., 2007) are large,
355 with distinct biotas, some level of endemism (mainly at species level), and distinctive abiotic environments.
356 In this context, the Benguela and Agulhas Currents prescribe the broadest scale hydrographic features
357 whose imprint can be seen, at that scale, on the seaweed flora: they maintain the Benguela Marine Province
358 (BMP) and Agulhas Marine Province (AMP, as per Spalding et al., 2007) of South Africa, separated
359 by an area of transition. The southward flowing Agulhas Current has an overriding effect on the east

360 coast, extending as far as the eastern portion of the Western Cape Province (58–22). Along the east
361 coast it is responsible for setting up a region that transitions from sub-tropical in the north to warm
362 temperate near Cwebe (*i.e.* the region 58–44), which is termed the East Coast Transition Zone (ECTZ).
363 From 44–22 the Agulhas Current continues to maintain an influence on the coast, but its direct effect is
364 subdued because of the widening of the continental shelf southwards of Cwebe, and the warm-temperate
365 biogeographical region, the AMP, is consequentially formed. The northward flowing Benguela Current
366 from which upwelling is maintained by prevailing south-easterly trade winds influences the remainder of
367 the Western Cape Province (west of 22), particularly from the western side of the Cape Peninsula (16/17)
368 northwards to about 16°S. The influence of the Benguela Current here defines a cool-temperate regime, the
369 BMP, with the range of monthly mean temperatures at most sections intermediate between cold-temperate
370 and warm-temperate, according to the definitions of (Lüning et al., 1990).

371 The oceanographic features of the Benguela and Agulhas Currents set up gradients in seawater
372 temperature that run predominantly from east to west (*i.e.* August mean temperatures increasing in this
373 direction), but also from north to south (*i.e.* with the southern coastline becoming more variable in its
374 thermal regime). Of particular ecological importance is the gradient in August mean temperature. It clearly
375 structures the thermal distance decay curve, which shows a steadily increasing dissimilarity in thermal
376 distance along the shore from 1. The gradient may be smooth or punctuated at irregular intervals by
377 peaks in dissimilarity. Specifically, the mean temperature for Austral winter (August) increases more-or-
378 less smoothly from the west. Within the region, species richness along the coastline reflects the general
379 global trend — for most taxa but ironically not for seaweeds (Bolton, 1994; Santelices et al., 2009) — of
380 diminishing diversity with decreasing temperature, which at that scale is seen as a latitudinal gradient:
381 the cold temperate area of the west coast has a much lower α -diversity than the warm temperate and sub-
382 tropical south and east coasts, with the latter two regions having similar values. This same pattern exists
383 for fish and invertebrates in this region, albeit in data which include coastal waters further from the shore
384 (Griffiths et al., 2010). In the previous presentation of the seaweed dataset (Bolton and Stegenga, 2002), the
385 ECTZ had far fewer species per section than the AMP, but this change in the dataset results from significant
386 research in the period 2000–2005 on the seaweeds of the ECTZ (De Clerck et al., 2005). In contrast, bringing
387 the annual mean temperature or thermal range during Austral summer into the calculation of the thermal
388 distance introduces several large punctuations in dissimilarities between some sections — most notably
389 near Cape Point into False Bay (17), and near 30 and 48. The region of the coastline where low biodiversity
390 community composition changes to highly biodiverse communities is reflected in the β -diversity, which
391 peaks around the Cape Peninsula (16/17). Correlations between β -diversity (resulting from a change in
392 α -diversity) and areas of transition have also been noted in several studies on terrestrial biota (Melo et al.,
393 2009; Tonial et al., 2012), but such investigations are less established for marine biota (Schils and Wilson,
394 2006; Anderson et al., 2013).

395 β -diversity is generally lower within individual bioregions than at the scale of the country, because the
396 coastlines are shorter and large differences in diversity cannot approach that which can occur over the entire
397 coastline. Even within bioregions, β -diversity is still *generally* dominated by species turnover rather than
398 by nestedness-resultant processes. The exception is the BMP where nestedness-resultant (β_{sne}) assembly
399 processes contribute to around 58% of the β -diversity. The generally strong influence of turnover-based
400 processes elsewhere highlights the role of environmental drivers, which here are spatially structured as
401 one would expect of the steep environmental gradients, and hence of niche influences (*i.e.* environmental
402 filtering) in determining species assembly (Fitzpatrick et al., 2013).

403 It seems plausible that factors linked with the establishment of oceanographic regimes (historical
404 factors *sensu* Baselga et al., 2012) that influenced dispersal and speciation along the coast resulted in
405 the current-day gradients in β -diversity within bioregions. This hypothesis was put forward to explain
406 the low endemism and low species richness of Ochrophyta in the upwelling dominated west coast of
407 southern Africa (Bolton, 1986). But is the underlying mechanism the selection of cold tolerant species
408 from amongst a more diverse pool in the wider region (Bolton, 1986), or is it due to a dispersal barrier
409 that prevented elements of the warmer south coast seaweed flora from occupying the west coast? Evidence
410 from seaweed phylogeography postulates different evolutionary origins for seaweeds of the BMP and AMP
411 (Hommersand, 1986; Hommersand and Fredericq, 2003). The mechanism could be the intensification of
412 upwelling during the mid-Pliocene 4.6–3.2 Ma (Marlow et al., 2000). Turnover is almost entirely absent
413 in favour of nestedness-resultant processes in the BMP, but the AMP is defined entirely by a moderate
414 rate of turnover with respect to both geographical and thermal distance. The dominance of the nestedness-
415 resultant component of β -diversity within the BMP suggests that the complex, non-linear influence of the
416 annual mean temperature selectively influences species richness from section to section in a pattern that is
417 not spatially coherent, but this climatic variable (*i.e.* fraction [E], which has *some* spatial structure) explains
418 only 39.1% of the β_{sne} patterning in the region. Unaccounted-for non-climatic variables and perhaps
419 non-environmental variables may explain some of the remaining variation. This points to habitat and/or
420 environmental heterogeneity and neutral processes such as dispersal limitation, stochastic influences or
421 habitat heterogeneity as the main structuring agents of communities within the region. Indeed, habitat
422 heterogeneity generated through stochastic influences such as patchiness due to storms denuding portions
423 of kelp beds (which is the dominant habitat type along the west coast) is well-known in kelp forests (Smale
424 et al., 2011). Furthermore, the prevalence of two kinds of assembly processes to either side of the Cape
425 Peninsula (17), coinciding with the point where a region of low α -diversity (BMP) transitions into a region
426 of higher α -diversity (the Benguela-Agulhas Transition Zone, B-ATZ), suggests that the answer to our
427 question is probably that some historical event was responsible for that disjunction, and that this barrier is
428 still maintained today due to limited mixing between the Agulhas and Benguela Currents (although some
429 of the sections along the west coast between 16/17 and 10 certainly have a reasonable probability of sharing
430 species with sections at the western-most side of the AMP, *e.g.* 23, 24; see more on the B-ATZ, below).

431 Rapid rates of species turnover with respect to geographic and thermal distance along the east coast,
432 and the steep decrease in species richness around the 16/17–22 transition region, suggest that relatively
433 more endemic species should occur within the BMP and AMP. Seawater becomes warmer along a steep
434 gradient along the east coast northward towards 54 where-after there is a further steepening of the gradient
435 towards Mozambique (north of 58). This steepening of the already strong temperature gradient in the
436 northern portion of the ECTZ supports the conclusion of (Bolton and Anderson, 2004) that it represents
437 the transition from a tropical Indian Ocean seaweed flora in Mozambique to a temperate flora in the south.
438 A similar pattern exists for the rocky intertidal biota in the region (Sink et al., 2005). The temperature of
439 the coldest month of the year sufficiently accounts for the environmental gradient in all bioregions, except
440 for in the BMP. This strong coupling between the thermal gradient and β_{sim} suggests that a niche difference
441 mechanism is the primary species compositional assembly process (Nekola et al., 1999). This implies species
442 being sorted based on their physiological tolerances along a gradient, and the particularly high rate of
443 species turnover along the east coast reflects the species' narrow thermal ranges. Species here have narrow
444 local distributions, and there is only a 2% seaweed endemism in the region (Bolton and Anderson, 2004).

445 The overlap region between 22 and the southern tip of the Cape Peninsula (16/17), which we call the
446 B-ATZ, is an area where aspects of both currents may be periodically seen (Largier et al., 1992), and it
447 is not surprising that biogeographically this region includes biotic elements from the marine provinces

on both sides of it. This is clearly visible in our analyses, which show the highest rates of species turnover (β_{sim}) with respect to geographical distance, the mean temperature for August and the range in temperature in February. This is a form of regional biogeographical structuring called the temporal/spatial constraint model (Nekola et al., 1999). Similar conclusions with regards to the B-ATZ being an transitional area have been reached by (Stephenson, 1948; Bolton, 1986; Bolton and Anderson, 2004; Mead et al., 2013). For marine macroalgae at least, this area cannot be recognised as a marine province as it does not meet the criteria for it to be classified as such (Spalding et al., 2007), *i.e.* it lacks cohesion, levels of endemism, and there is an absence of distinct abiotic features (with exception of such conditions periodically arising in False Bay due to prevailing mesoscale oceanographic conditions).

It was recently suggested that studies which associate marine macroalgal distributional patterns with broad-scale temperature gradients neglect to consider alternative or additional explanations, such as that offered by connectivity due to ocean currents (Wernberg et al., 2013). It is possible that seaweeds are similarly influenced by ocean currents around South Africa, and it is tempting to suggest that the *direction* of turnover along the east coast is from north to south, and along the south coast from east to west, as this would coincide with the direction of the Agulhas Current. From a purely thermal point of view, the imprints of those ocean currents are certainly present at the coast in the zone that our seaweed data represent, but the physical processes that operate in that coastal zone are considerably more complex and decoupled from mesoscale influences (Schlegel et al., 2017). Personal observations by one of us (AS) recount dislodged individuals of the kelp, *Ecklonia maxima*, having washed ashore some 400 km east of the eastern-most distributional limit for the species in South Africa — this is *against* the direction of the current, which in this region is situated considerably far from the shore along the edge of the Agulhas Bank. Our data on the rate of species turnover as a function of the thermal difference between coastal sections (β_{sim} vs. d_E) within bioregions show an almost precise relationship between thermal difference and change in species composition (even though the relationship between d_E and geographical distance differs between the east and south coasts), and provide very strong support for temperature, rather than connectivity due to currents, as an overriding driver of seaweed community composition over large spatial scales. Within the context of Beijerinck's 'Law' that "everything is everywhere but the environment selects" (De Wit and Bouvier, 2006), the only role that ocean currents can have in setting up biogeographical patterns is to cause *everything not to be everywhere*. At these scales, with or without the influence of ocean currents, temperature still selects. In other words, at broad spatial scales within the region encompassed by the ECTZ and the AMP, the seaweed flora is structured predominantly by niche-assembly processes driven by a thermal gradient.

However, we should point out that *ca.* 13.7% of the seaweed flora's turnover is also explained by some other broad scale spatial influences (MEM2, MEM3 and MEM5), which represent exogenous (unmeasured) environmental variables and autocorrelation. The importance of these relative to each other is unknown. Future work may find that some of these environmental influences may indeed be that of ocean currents, which operate at those broad scales, but our current interpretation does not support this notion. Rather, we suggest that the unknown influence is more likely to be neutral influences (Hubbell, 2001) manifesting at a reasonably broad scale. These may include stochastic events (*e.g.* Smale et al., 2011) or biotic processes caused by differences in species demography, dispersal or the influence of grazers, etc. Such processes are usually finer scaled, but, having said that, the entire west coast region that makes up the BMP is dominated by nestedness-resultant β -diversity, which is also symptomatic of neutral assembly processes.

491 4.1 Conclusions

492 Our approach throughout this paper hinges upon the biogeographical provinces and overlap regions
493 defined using both seaweed and thermal data. Together, thermal and species gradients/patterns provide
494 the most parsimonious explanation for the processes assembling the seaweed flora within the study region.
495 The explanation is that thermal gradients set up β -diversity based on species turnover as a result of
496 environmental filtering within the southern and eastern coastal sections, while along the west coast a
497 very different kind of assembly process, nestedness-resultant β -diversity, emerges. We suggest that this
498 disjunction results from a historical event that is currently still limiting connectivity between the Indian
499 and Atlantic Oceans around the Cape Peninsula.

500 We also show how thermal metrics other than the annual mean temperatures can aid in the delineation
501 of macroecological patterns. Some species are limited by cold and some by warm temperatures, and
502 the constraining factor may differ at a species' southern and northern limits (e.g. van den Hoek, 1982a;
503 Breeman, 1988). If these are important determinants, as we indeed show there are, other measures of
504 climate should be used in our repertoire of explanatory variables. For example, the mean temperature for
505 the coldest month, standard deviation, variation in annual temperature extremes, or temperature ranges
506 (e.g. as used by Qian and Ricklefs, 2007; Leaper et al., 2011) may readily be extracted from time series of
507 daily temperatures. These approaches, as we apply them here, may greatly improve macroecological and
508 biogeographic studies and aid our ability to analyse broad-scale communities patterns and the processes
509 that assemble them.

510 Conflict of Interest Statement

511 The authors declare that the research was conducted in the absence of any commercial or financial
512 relationships that could be construed as a potential conflict of interest.

513 Author Contributions

514 AS conceptualised the scope of the research reported in this paper, undertook all the numerical and
515 statistical analyses, made the first round of interpretation, and did the bulk of the writing. JB and RA
516 collected all the samples over the last 30 years, compiled the database of species distribution records for the
517 region included in this analysis, contributed in equal part to the conceptualisation of the research and the
518 interpretation of the findings, and provided significant editorial input into penultimate and final drafts of
519 the document.

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Tables

Table 1. Variance partitioning of β_{sim} and β_{sne} by the thermal and spatial predictors. F and p -values are shown for the testable fractions only.

Fraction		d.f.	Adj. r^2	F	P
turnover, β_{sim}					
[E], thermal variables		4	0.842	110.160	0.001
[S], spatial variables		16	0.961	80.731	0.001
[E+S], all thermal and spatial		20	0.979	191.560	0.001
[E S], thermal, non-spatial		4	0.018	20.506	0.001
[E with S], spatially structured thermal			0.824		
[S E], spatial, non-thermal		16	0.137	23.649	0.001
1-[E+S], residual variance			0.021		
nestedness-resultant, β_{sne}					
[E], thermal variables		1	0.391	-6.017	—
[S], spatial variables		4	0.678	12.060	0.001
[E+S], all thermal and spatial		5	0.714	9.077	0.001
[E S], thermal, non-spatial		1	0.040	-1.018	0.997
[E with S], spatially structured thermal			0.355		
[S E], spatial, non-thermal		4	0.323	14.277	0.001
1-[E+S], residual variance			0.286		

Table 2. β -diversity (β_{sor}) as well as its partitioning into β_{sim} and β_{sne} components. The partitioning represents the contribution of turnover and nestedness-resultant processes. Mean \pm SD values are presented for the whole South African coast, as well as for each of the four bioregions.

Region	β_{sor}	β_{sim}	β_{sne}	$\beta_{\text{sne}}/\beta_{\text{sor}}$
BMP	0.105 ± 0.086	0.044 ± 0.053	0.061 ± 0.059	0.581
B-ATZ	0.098 ± 0.068	0.083 ± 0.071	0.014 ± 0.009	0.143
AMP	0.117 ± 0.076	0.087 ± 0.067	0.030 ± 0.020	0.170
ECTZ	0.259 ± 0.157	0.234 ± 0.162	0.025 ± 0.018	0.097
overall	0.496 ± 0.287	0.433 ± 0.277	0.063 ± 0.061	0.127

Table 3. Linear regressions of pairwise β_{sim} and β_{sne} as a function of the geographical or thermal distance between the section pairs. The thermal distances that best explain β_{sim} and β_{sne} were selected during the db-RDA procedure. Here, β is the slope of the regression line (per 100 km in the case of the relationship with distance), and the P -value test the hypothesis that the slope is significantly different from 0 using a t -test. Refer to Fig. 5a–g.

bioregion	β	t -value	P	Adj. r^2
β_{sim} vs. distance				
BMP	0.007 ± 0.002	3.886	<0.001	0.052
B-ATZ	0.109 ± 0.016	6.873	<0.001	0.658
AMP	0.029 ± 0.001	44.751	<0.001	0.834
ECTZ	0.079 ± 0.001	65.003	<0.001	0.936
β_{sim} vs. augMean d_E				
BMP	0.009 ± 0.012	0.765	0.445	-0.001
B-ATZ	0.342 ± 0.060	6.139	<0.001	0.605
AMP	0.290 ± 0.010	29.209	<0.001	0.681
ECTZ	0.350 ± 0.011	31.200	<0.001	0.772
β_{sim} vs. febRange d_E				
BMP	0.008 ± 0.011	0.710	0.479	-0.002
B-ATZ	0.259 ± 0.047	5.491	<0.001	0.548
AMP	-0.006 ± 0.004	-1.422	0.156	0.003
ECTZ	0.183 ± 0.009	20.102	<0.001	0.583
β_{sim} vs. febSD d_E				
BMP	0.018 ± 0.011	1.583	0.115	0.006
B-ATZ	0.161 ± 0.080	2.029	0.054	0.115
AMP	-0.007 ± 0.002	-2.758	0.006	0.016
ECTZ	0.171 ± 0.013	13.401	<0.001	0.383
β_{sim} vs. augSD d_E				
BMP	0.038 ± 0.008	4.590	<0.001	0.073
B-ATZ	0.103 ± 0.032	3.183	0.004	0.276
AMP	-0.003 ± 0.003	-0.936	0.350	0.000
ECTZ	0.082 ± 0.008	11.411	<0.001	0.310
β_{sne} vs. distance				
BMP	0.015 ± 0.002	8.178	<0.001	0.205
B-ATZ	0.006 ± 0.004	1.541	0.137	0.054
AMP	0.006 ± 0.000	15.255	<0.001	0.367
ECTZ	-0.001 ± 0.001	-2.200	0.029	0.013
β_{sne} vs. annMean d_E				
BMP	0.067 ± 0.009	7.141	<0.001	0.164
B-ATZ	0.037 ± 0.013	2.929	0.008	0.240
AMP	-0.009 ± 0.004	-2.499	0.013	0.013
ECTZ	-0.002 ± 0.003	-0.845	0.399	-0.001

Figures and captions

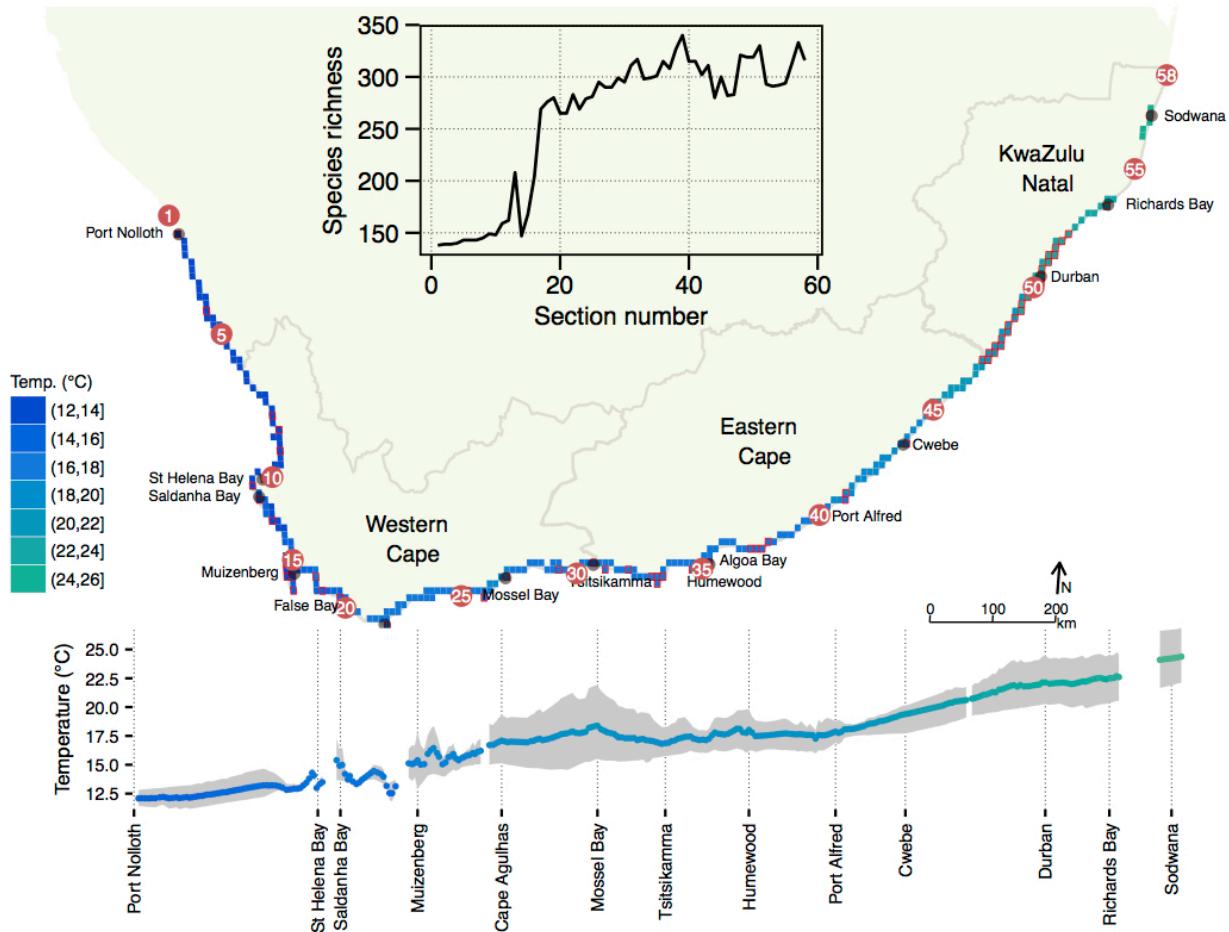


Figure 1. Profile of mean annual temperature along the coast of South Africa. Profiles are indicated as a function of geographical position on a coastal map (top) and as a function of distance away from Section 1 (bottom). The latter visualisation also indicates the long-term minimum (mean August) and maximum (mean February) temperatures as a grey shaded area around the annual mean temperature. The inset shows the species richness of macroalgae along the coast. Note that although a detailed temperature profile is displayed here, further analyses in this paper proceed with temperature data interpolated to the 58 sections for which seaweed diversity data are available.

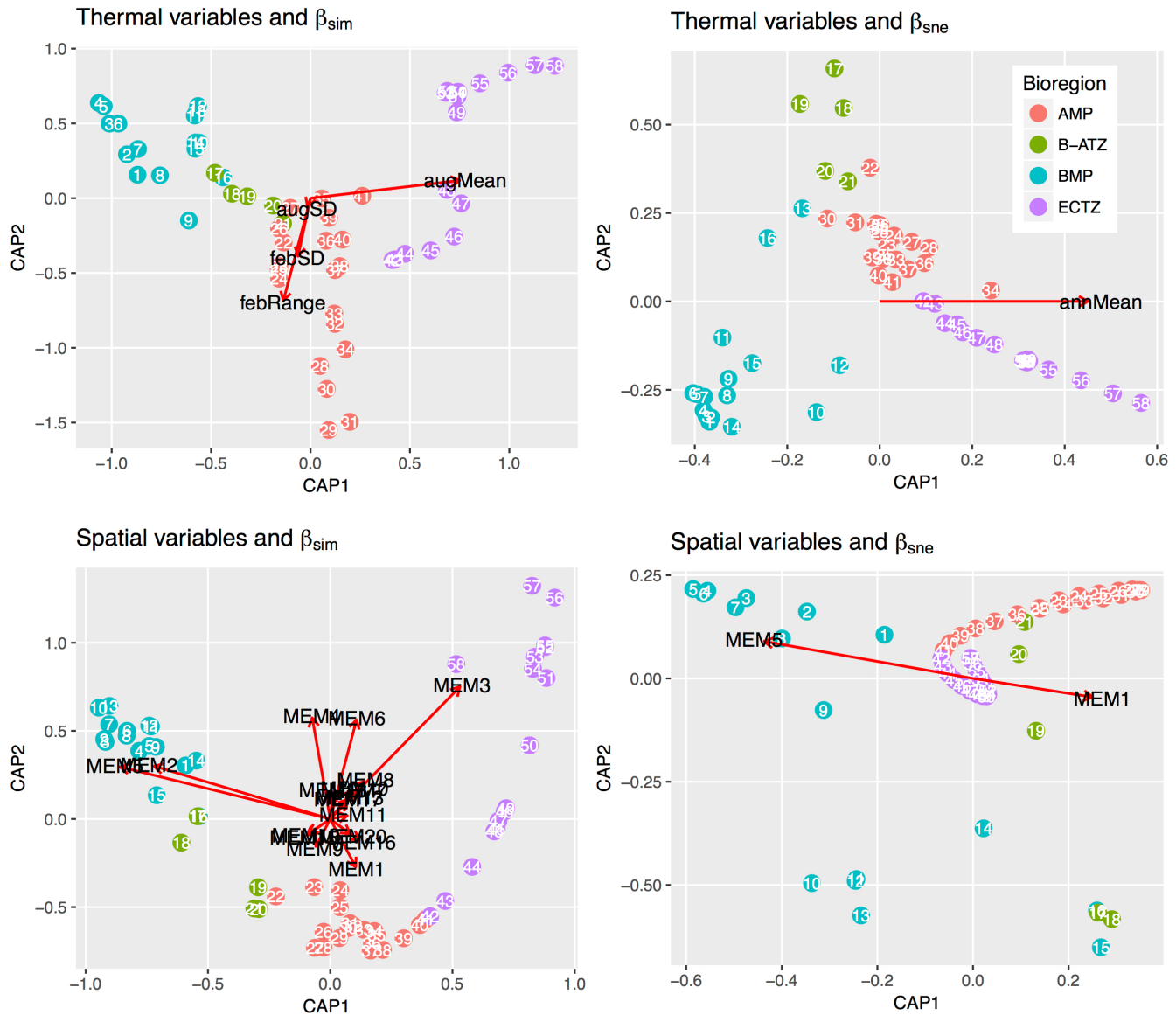


Figure 2. db-RDA biplots of lc scores for β_{sim} and β_{sne} constrained by the thermal and spatial MEM variables. The only two canonical axes shown are CAP1 and CAP2 since these capture the bulk of the inertia present in the ordinations. The constraining vectors were selected during the db-RDA steps.

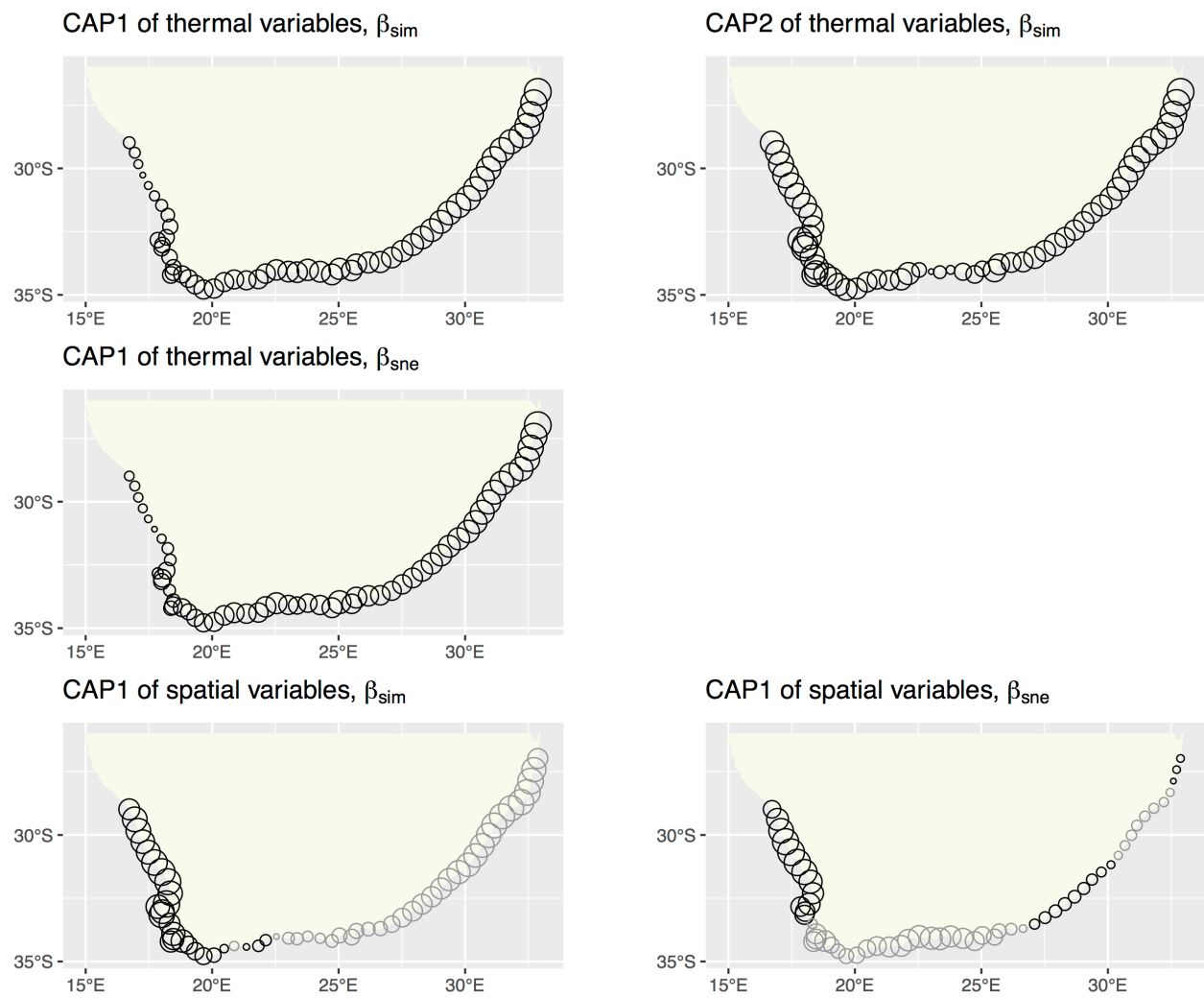


Figure 3. db-RDA site scores for β_{sim} and β_{sne} on geographic coordinates. The site scores indicate the major gradients captured by the environmental and spatial (MEM) constraints. Gray circles indicate negative site scores.

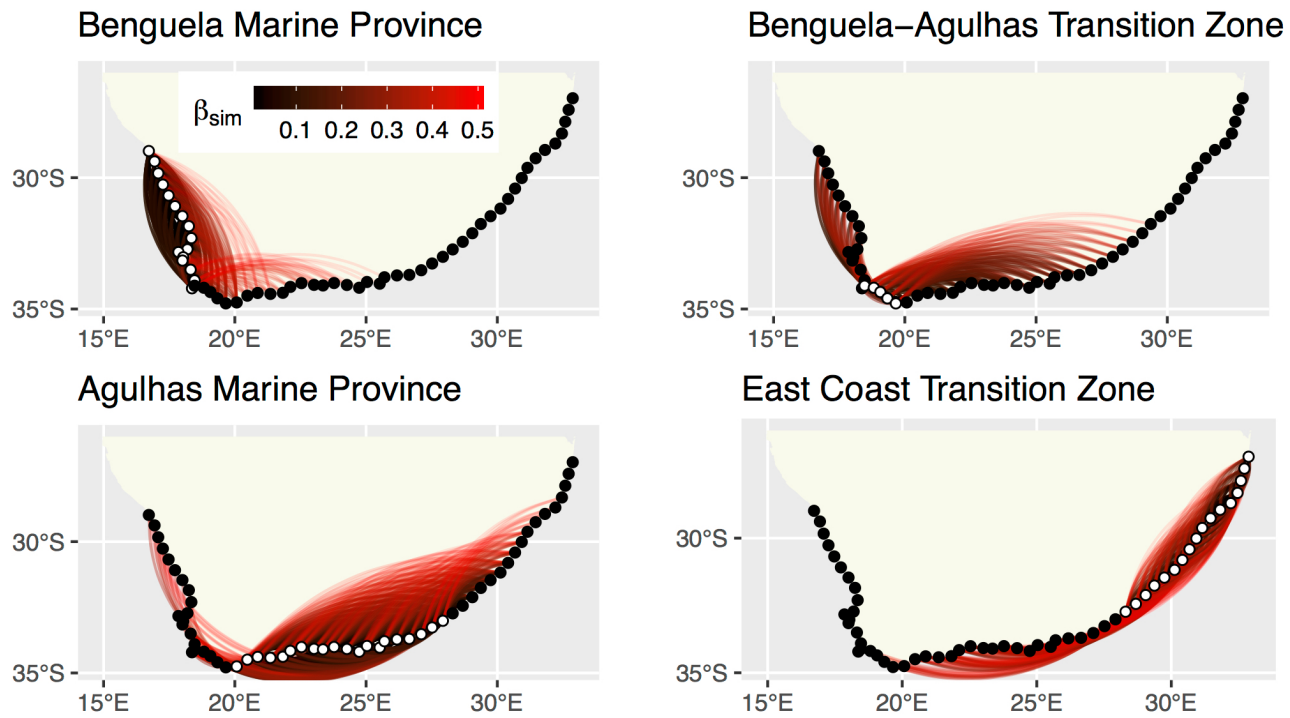


Figure 4. Network graphs of pairwise β_{sim} showing sections that are similar to one-another in species composition. Dissimilarity indices range from >0.0–0.5 (possible dissimilarities range from 0.0–1.0). The 58 sections in Appendix A appear as dots at the vertices of the network graph, with those belonging with the ‘active’ bioregion shown in white. The pairwise dissimilarities are shown by the coloured lines, with blacker lines indicating lower dissimilarity indices (species composition more similar) and redder ones higher dissimilarity indices (species composition more dissimilar).

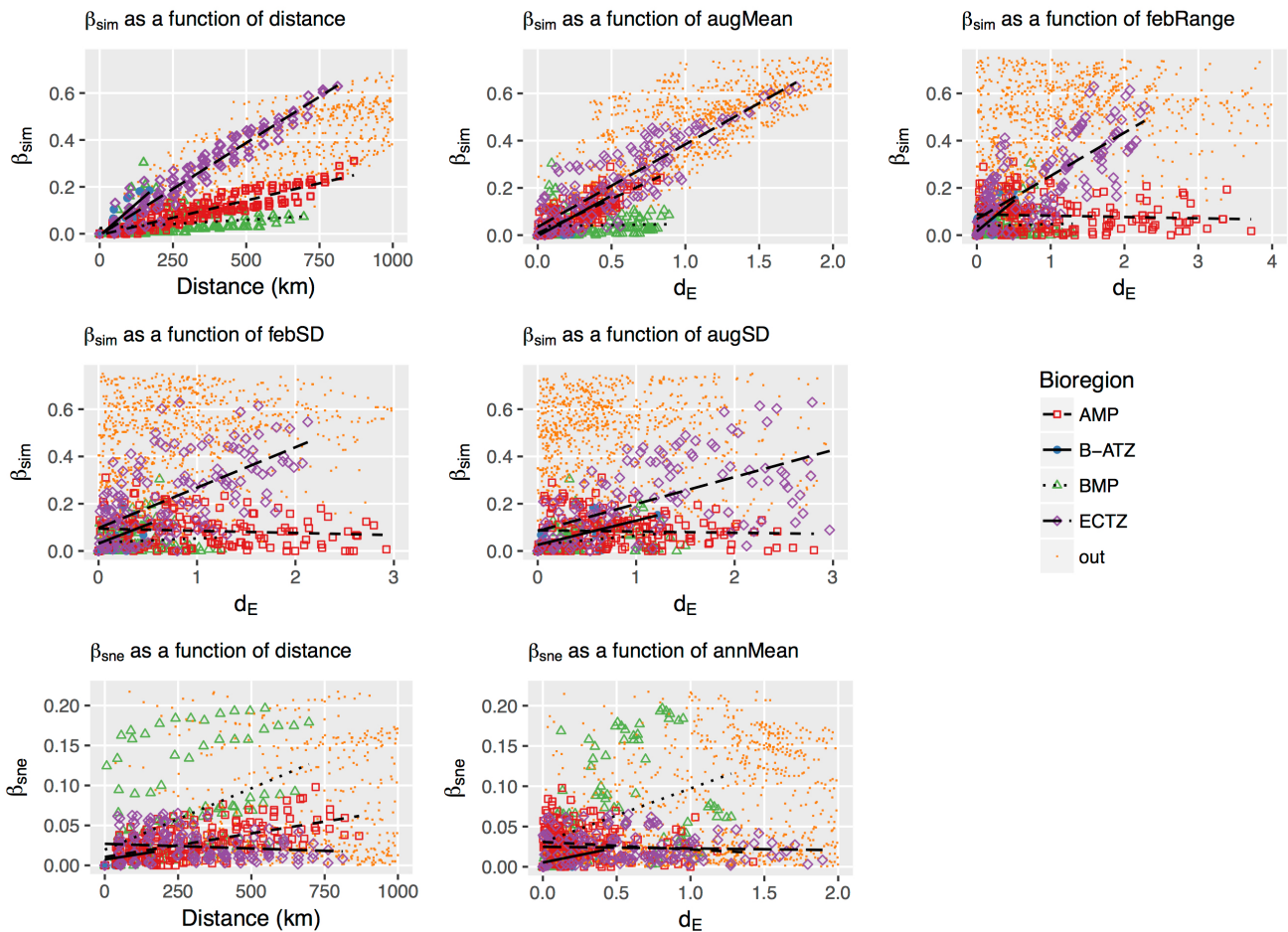


Figure 5. Plots of the turnover (β_{sim}) and nestedness-resultant (β_{sne}) forms of β -diversity as a function of geographical or thermal distance between the coastal sections. The influential d_E variables (*augMean*, *febRange*, *febSD* and *augSD* for β_{sim} , and *annMean* for β_{sne}) were determined in the db-RDA procedure. β_{sim} , β_{sne} , d_E and geographical distance were calculated as differences between coastal section pairs, and data points representing section pairs falling between bioregions are coloured yellow and labelled 'out'. The strengths and gradients of the regression lines indicated in the panels are presented in Table 3.