SEABIRD ASSEMBLAGES ARE LINKED TO THE MAJOR WESTERN BOUNDARY CURRENT OFF EASTERN AUSTRALIA

SEABIRD ASSEMBLAGES OFF EASTERN AUSTRALIA

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Abstract

Aim: To identify assemblages of seabird species and relate them to environmental conditions off eastern Australia, a world seabird hotspot. Location: East coast of Australia. Methods: We used 15 seabird at-sea ship-based survey data sets collected over 37° of latitude, from 2016 to 2021, and fitted Region of Common Profile (RCP) mixture models for each season and type of data (presence-absence and abundance). RCP groups are defined as a set of regions where the probability of encountering a specific species profile is constant within regions, but

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different amongst them. These groups also vary according to covariates, which in our case included oceanographic, climatic, and physiographic parameters. Results: Results were based on 142,646 seabirds recorded, from 80 species, including albatrosses, petrels, prions, shearwaters, boobies, and terns, among other taxa. All models suggested two macro-scale assemblages ('northern' and 'southern'), except for the autumn presence-absence model that identified three groups. The model results consistently show a biogeographic transition at ~34°S, near the latitude at which the East Australian Current (EAC) separates from the Australian continental slope. Sea surface temperatures or sea surface salinities were selected in all final models, further indicating a close relationship between seabird assemblages and water masses. Results from both data types, presence-absence and abundance, resulted in similar spatial and species profile patterns. Main conclusions: RCP models clearly identified two seabird assemblages off the east coast of Australia, suggesting the persistence of these groups at seasonal and macro spatial scales. Given ongoing poleward intensification that the EAC is experiencing, which is projected to continue over the next century, and its importance in influencing distributions of seabirds, the methods applied in our study could be replicated to assess possible changes in seabird assemblages, and how they are affected by changing environmental conditions.

Keywords Assemblages · Biogeography · Community ecology · East Australian Current · Mixture models · Region of Common Profiles

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Biosketch

Nicholas W. Daudt (he/him) is broadly interested in spatial ecology and ecological modelling. His research uses data science and modelling to understand biogeographic patterns and ecological processes that build marine top predators' biodiversity patterns across space and time.

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Seabird assemblages are linked to the major western boundary current off eastern Australia

2 Running title: Seabird assemblages off Eastern Australia

4 Abstract

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- 6 Australia, a world seabird hotspot.
- 7 Location: East coast of Australia.
- 8 Methods: We used 15 seabird at-sea ship-based survey data sets collected over 37° of latitude, from 2016
- 9 to 2021, and fitted Region of Common Profile (RCP) mixture models for each season and type of data
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- 11 encountering a specific species profile is constant within regions, but different amongst them. These groups
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- 13 parameters.
- 14 Results: Results were based on 142,646 seabirds recorded, from 80 species, including albatrosses, petrels,
- prions, shearwaters, boobies, and terns, among other taxa. All models suggested two macro-scale assemblages
- 16 ('northern' and 'southern'), except for the autumn presence-absence model that identified three groups. The
- 17 model results consistently show a biogeographic transition at ~34°S, near the latitude at which the East
- Australian Current (EAC) separates from the Australian continental slope. Sea surface temperatures or sea
- 19 surface salinities were selected in all final models, further indicating a close relationship between seabird
- 20 assemblages and water masses. Results from both data types, presence-absence and abundance, resulted in
- 21 similar spatial and species profile patterns.
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- 23 suggesting the persistence of these groups at seasonal and macro spatial scales. Given ongoing poleward
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30 1 Introduction

Bioregionalization is the discipline that identifies, delimits, characterises, and names biogeographical regions 31 (Morrone, 2018; Vilhena & Antonelli, 2015). Although there is often no agreement on the naming systems 32 among proposed regionalization schema (but see Ebach et al., 2008), the hierarchical nature of bioregions 33 is well recognized (Kreft & Jetz, 2010; Vilhena & Antonelli, 2015). Thus, bioregions are usually described within geographically larger classifications, such as 'provinces' or 'realms' (Costello et al., 2017; Spalding et al., 2007). The delimitation of bioregions is made through identifying sets of species that are endemic or have high levels of co-occurrence, and are distinct enough from other regions. In this sense, bioregions are 37 interpreted as natural representations of common historical and/or ecological processes shaped by physical 38 and biological forces (Kreft & Jetz, 2010). In particular, the identification of bioregions allows representation 39 of the distribution of multiple species at a given time and space, and relating these regions to environmental 40 characteristics. By doing so, bioregions can be interpreted as species assemblages (N. A. Hill et al., 2017; 41 Lyons et al., 2017). Given these features, bioregions can be used as surrogates for biodiversity monitoring in data-poor regions and in regional conservation planning (Ferrier, 2002; Spalding et al., 2007), playing a fundamental role in conservation biogeography (Whittaker et al., 2005). Seabird assemblages have been widely described and related to environmental characteristics (Hyrenbach et al., 2007; Ribic et al., 1997; Woehler et al., 2003). Considering the hierarchical nature of oceanographic processes and their resulting features (Haury et al., 1978), it is not surprising that the distribution of highly mobile 47 seabirds are linked differently across temporal and spatial scales (Haury et al., 1978; Hunt & Schneider, 1987). 48 In the marine environment, large-scale oceanographic features (e.g. boundary currents and water masses) act 49 similarly to terrestrial biomes (based on vegetation), and influence seabird assemblage structure—i.e. the 50 occurrence of species and their relative numerical composition (Commins et al., 2014; Hyrenbach et al., 2007; 51 Ribic et al., 1997). In contrast, local abundance is related to meso- (10–100 km) and sub-mesoscale (1–10 km) processes that aggregate prey and therefore predators, such as fronts and eddies (Scales, Miller, Hawkes, et al., 2014; Weimerskirch, 2007). Such processes, although relatively ephemeral, are primarily driven by physical and atmospheric forces, and are thus somewhat predictable in space and time, providing consistent feeding areas for marine predators when they occur (Davoren, 2013; Hazen, Suryan, et al., 2013). Sea surface temperatures are a key driver of seabird distributions, at individual- and assemblage-levels, operating over 57 a range of spatial and temporal scales from macro (e.g. water masses) to fine (e.g. fronts) (Dunlop et al., 58 1988; Quillfeldt et al., 2015; Ribic et al., 1997). Despite extensive surveys (e.g. Ballance et al., 1997) and 59 tracking studies (e.g. Carneiro et al., 2020), many oceanic regions still lack information on the distributions 60 and abundances of seabird species, and how they assemble in space and time. 61 One area in need of better understanding of seabird distributions is Australia, considered a global seabird

hotspot (Karpouzi et al., 2007; Ramírez et al., 2017). Given the region's importance for seabird diversity,

there are surprisingly few studies on seabird distributions at sea (Bernard et al., 2021; Mott & Clarke, 2018). Off northwest Australia, at-sea surveys, multivariate analyses (Dunlop et al., 1988) and ensemble species 65 distribution models of species assemblages (Lavers et al., 2014) showed that habitat use by tropical seabird 66 species was related to sea surface temperatures and salinity. Dunlop et al. (1988) suggested that salinity was the main variable influencing species' distributions in oceanic waters while temperature was the key factor for species on the continental shelf. In contrast, there are few studies at large spatial and temporal scales off eastern Australia. T. A. Reid & Hindell (2000) and Evans et al. (2021) analysed seabird assemblages and 70 their relationships with oceanographic and prev data at mesoscale off southeast Tasmania. The remaining 71 studies are mostly descriptive accounts locally (Blaber & Milton, 1994; Norman, 1992), or focused on a single 72 species (Priddel et al., 2014). 73 The East Australian Current (EAC) is the major western boundary current in the South Pacific Ocean, and 74 the dominant boundary current off eastern Australia (Ganachaud et al., 2014). The EAC has been extensively 75 studied, although the links between the EAC's oceanography and marine predators are not well known. The dynamic features associated with the EAC, such as mesoscale eddies and upwelling, are likely to influence marine biota (Scales, Miller, Hawkes, et al., 2014; Suthers et al., 2011). While most of the biological studies 78 in this system were focused on plankton entrapment within the cores of eddies (Firme et al., 2023; Garcia et al., 2022; Hassler et al., 2011), there is a growing body of work focusing on fisheries (Brieva et al., 2015; 80 Hobday, 2010; Young et al., 2011). These studies emphasized how the probability of occurrence of a given 81 taxon, and the structure of species assemblages, were driven by the EAC. However, the relationships between 82 oceanography and marine megafauna within the EAC are poorly known (Suthers et al., 2011). Some progress 83 has been made using shark and ray movement data (Jaine et al., 2014; Niella et al., 2020; Niella et al., 2022), 84 and satellite tracking has also demonstrated that increased sea surface temperatures in the EAC decreases little penguin Eudyptula minor foraging success (Carroll et al., 2016), but information at the assemblage-level for seabirds in the EAC is absent. Marine species range shifts are more likely to occur faster than terrestrial species due to their higher sensitivity to warming (Lenoir et al., 2020; Pinsky et al., 2019). As a result, climate change is rearranging the structure of 89 species assemblages around the globe (Antão et al., 2020; Dornelas et al., 2014; Poloczanska et al., 2013). Top 90 predators, for instance, may change their core habitats (Hazen, Jorgensen, et al., 2013; Reisinger et al., 2022), 91 with unprecedented population consequences (Woehler & Hobday, 2023). For seabirds, Péron et al. (2010) and 92 Sojitra et al. (2022) showed that sea surface temperature anomalies affected the occurrence and abundances of albatross and petrel species with latitude, thus affecting assemblage structure. In eastern Australia, changing EAC conditions have affected a range of marine species. For example, changes in zooplankton assemblage structure and fish distribution patterns have been recorded and attributed to warming water temperatures, and shifts in shark distributions are expected (C. R. Johnson et al., 2011; Niella et al., 2022). A severe marine 97 heatwave driven by a poleward strengthening of the EAC, showed drastic environmental impacts including disease outbreaks in aquaculture farms and wildlife mortalities (Oliver et al., 2017). Given that the EAC is strengthening and extending farther southward and getting warmer (Cai et al., 2005; Phillips et al., 2022), biological changes are expected to occur (Hobday, 2010; Poloczanska et al., 2007).

In light of eastern Australia's importance for seabirds, and the recent attention on how predicted climate change will influence the EAC's marine biota (Poloczanska et al., 2007), an assessment of the contemporary spatial distributions of seabirds within the EAC at large temporal and spatial scales is timely. Seabirds can serve as sentinels for tracking ecological changes in the EAC in many ways (e.g. Sydeman et al., 2012). Surprisingly though, seabird distribution is under-represented in climate change studies, despite being highlighted as a priority topic (Grémillet & Boulinier, 2009; Sydeman et al., 2012).

In this study, we aim to identify assemblages of seabird species within the EAC and adjacent areas, and 108 relate these assemblages to environmental conditions, by using at-sea survey data and a state-of-the-art 109 modelling technique. We use the Region of Common Profiles – RCP (Foster et al., 2013), a mixture model 110 capable of describing bioregions (N. A. Hill et al., 2020; S. N. C. Woolley et al., 2020). RCP is a model-based 111 approach, and has a range of advantages over traditional algorithm-based statistics (N. A. Hill et al., 2020; Warton et al., 2015). Eastern Australia supports a high diversity of seabirds and offers wide environmental gradients over a near 40° latitudinal range, from the tropics to the cool temperate. We hypothesised to find 114 different seabird assemblages relating to different environmental conditions within the EAC. In addition, 115 presence-absence and abundance data were used to test if the different numerical resolutions would result in 116 the identification of similar assemblages. This study provides a repeatable framework on which future studies 117 can assess how future changes in the EAC may influence the distribution of seabird assemblages. 118

119 2 Methods

120 2.1 Study area

The EAC originates from the South Equatorial Current (SEC), a tropical nutrient-poor current. When the 121 SEC's North Caledonian Jet meets the Queensland Plateau (~18°S), it splits into a north-flowing branch 122 towards the Gulf of Papua and a south-flowing branch, which marks the start of the EAC (Ganachaud et 123 al., 2014; Kessler & Cravatte, 2013). The South Caledonian Jet completes the EAC system at ~22°S after 124 encountering the shelf break in the southern portion of the Great Barrier Reef (Figure 1a; Kessler & Cravatte 125 (2013); Ganachaud et al. (2014)). The EAC interacts with the bathymetry of the seafloor, giving rise to 126 oceanographic features and processes, such as smaller currents, upwellings, and persistent dynamic eddies 127 (Oke, Roughan, et al., 2019; Oke & Middleton, 2000; Ridgway & Dunn, 2003). The EAC flows polewards and 128 is primarily associated with the shelf break, but at ~32°S it detaches from the shelf and spawns an area of 129 eddy generation that dissipates throughout the western Tasman Sea (Mata et al., 2006; Oke, Roughan, et 130 al., 2019), the 'EAC eastern extension', often termed the 'Tasman Front' (Oke, Pilo, et al., 2019). Although most EAC energy is converted into eddies, the EAC still runs southwards following the shelf break towards
Tasmania (the 'EAC southern extension'; **Figure 1a**), carrying eddy-like features as a transient current (Oke,
Roughan, et al., 2019).

135 2.2 Seabird and environmental data sources

Seabird data were gathered aboard RV Investigator and RV Falkor on 15 voyages, between 2016/17–2020/21, 136 spanning over ~37° of latitude (Figure 1b; Table S1). One or more observers continuously recorded seabirds 137 within a 300 m forward quadrant (eye height 25 m a.s.l) from sunrise to sunset hours while the ship was 138 underway. The species, the numbers of birds, and their behaviours were logged in real time into a ship-based 139 web server. Environmental parameters for atmospheric and oceanographic conditions were logged continuously 140 by automated-ship based sensors, with seabird observations supplementing relevant environmental conditions 141 such as visibility or associations with marine mammals. Individual records received a date/time/geographic 142 coordinate label (multi-species records have the same label but are logged as separate entries for each species). Birds following the vessels were excluded from analyses. Seabird observation protocols followed Raymond et al. (2010), in use around Australia since 1980. 145 To investigate and predict species' distributions and relationships with environmental variables, a considered 146 choice of broad-scale climatic and oceanographic covariates is important (Barbet-Massin & Jetz, 2014). We 147 used three physiographic variables and seven oceanographic variables, from which a further three climatic 148 layers were calculated (Table 1). These variables are the most frequently used in seabird-environment 149 relationship studies (Tremblay et al., 2009), capturing processes at a range of spatial and temporal scales 150 from macro-scale water-mass properties to prev availability in meso- to submesoscale oceanographic features 151 (Haury et al., 1978; Scales, Miller, Hawkes, et al., 2014; Weimerskirch, 2007). 152

153 2.3 Data treatment

157

Data wrangling, modelling and visualization were done in R 4.2.0 (R Core Team, 2022); the full list of packages can be found in the Supplementary Material. Code and the modelling dataset (see Section 2.3.2) are available in (Zenodo).

2.3.1 Environmental layers

The spatial and temporal persistence of oceanographic features such as eddy kinetic energy ('eke') and thermal fronts (sea surface temperature gradient: 'sst_grad') may influence the movements and distributions of seabirds by offering a higher likelihood of finding food (Scales, Miller, Embling, et al., 2014; Weimerskirch, 2007). Thus, 'eke' and 'sst_grad' environmental layers were calculated before extracting environmental data for each seabird record. For both 'eke' and 'sst_grad', we used their original cell spatial resolution (Table 1) and calculations were done layer-by-layer to match seabird data on the finest spatial scale possible.

Eddy kinetic energy was calculated as $0.5(u'^2+v'^2)$, where $u'=u-\bar{u}$ and $v'=v-\bar{v}$ are the surface zonal 164 (latitudinal) and meridian (longitudinal) geostrophic currents, respectively, using a rolling mean of 90 days as 165 \bar{u} and \bar{v} (Sun et al., 2022). The spatial 'sst_grad' for each cell was calculated by averaging the maximum value 166 for each neighbouring cell (average neighbourhood maximum), from north-to-south (NS) and west-to-east (WE); then, the magnitude of the gradient was defined as $NS^2 + WE^2$ (Burrows et al., 2011). In addition, we calculated climatic seasonal average layers for 'eke' and 'sst $_$ grad' based on the 11-year data for 2011–2021 as 169 a proxy for persistent oceanographic features, and the 'eke' standard deviation over the same period (named 170 with the prefix 'clim' in **Table** 1). Seabed slope was calculated using 'raster::terrain()' (Hijmans, 2022a). 171 and distance from the coast was calculated through 'sf::st distance()' (Pebesma, 2018) using Australia and 172 Tasmania as the mainland (i.e. ignoring smaller islands). Then, for each seabird record, the most proximate 173 value for each environmental layer (original or calculated) was extracted using built-in functions from 'raster' 174 3.5-21, 'terra' 1.6-7, and 'sf' 1.0-8 packages (Hijmans, 2022a, 2022b; Pebesma, 2018).

176 2.3.2 Data

Seabirds are highly mobile animals, and many species migrate and/or disperse during their non-breeding 177 periods (Schreiber & Burger, 2002). Therefore, seasonal differences in species composition were expected 178 (e.g. K. Reid et al., 2023), and in the probabilities of occurrence and in total numbers. Thus, we built 179 separate modelling datasets for each season. We aggregated seabird data by 1° latitude × 1° longitude 180 grid cells (c. 110×110 km) and season (**Figure 1b**), by summing seabirds from the same species. To 181 test if the presence-absence and abundance (raw counts) data would result in similar assemblage patterns, 182 we built two species matrices, where the presence-absence matrix transformed values >1 to 1. To avoid 183 numerical instabilities in the models prompted by unusual or rare species, we kept only species occurring in 184 at least six grids within each season (summer = 12 species; autumn = 21; winter = 20; spring = 33). The 185 associated environmental data of each seabird record were averaged within the same grid cells and seasons, 186 and environmental data were standardised before modelling. 187

188 2.4 Statistical modelling

We used the mixture model Region of Common Profiles (RCP) to identify and describe seabird species assemblages. RCP is a one-stage bioregionalization approach, as it identifies bioregions by specifying a statistical model that simultaneously relates response (multi-species data) and predictor (environmental data) (N. A. Hill et al., 2020; S. N. C. Woolley et al., 2020). Given a set of environmental variables, the model can predict assemblage probabilities for sites where no biotic (observational) data exist. As a model-based approach, RCPs can be compared by information criteria, present uncertainty around the results, and check for model assumptions through assessing residuals (N. A. Hill et al., 2020; Warton et al., 2015).

The model is specified in terms of assemblages and site-specific probabilities. There are K assemblages 196 (i.e. RCPs), K = (1, ..., k), that are common across the study area. The outcome variable is a vector 197 $y_i = (y_{i1}, ..., y_{iS})$, that gives the count or presence-absence of each of the S species at site i. Depending 198 on whether we model the counts or presence-absence data, the assemblages are defined in terms of the 199 expected frequency or relative frequency of the species, respectively. An output of the model is the estimated probability that site i has each of the K possible assemblages. RCPs are a multivariate adaptation of a 201 mixture-of-experts model; as such, the mixing proportions (assemblage structure) vary according to covariates 202 (predictors) (Foster et al., 2013). RCPs can be directly interpreted as assemblages; the assemblage structure 203 is given by the species profiles. 204 Before fitting models, covariates were checked for collinearity within each seasonal dataset. If a pairwise 205 collinearity of |x| > 0.7 was found we removed one of the variables (Dormann et al., 2013), favouring direct 206 measures over the 'climatic' layers we calculated (e.g. if 'sst_grad' and 'clim_sst_grad' were correlated, we 207 chose 'sst grad'). Thus, 'full models' differed between seasons as the retained covariates varied, although 208 most covariates were common for all seasons. Due to seabird migration and dispersion, seasons could act as 'sampling artifacts' (Foster et al., 2017), 210 thus biasing results (Ponti & Sannolo, 2023). In light of the difference among seasons in the numbers of 211 species that met our occurrence threshold (see Section 2.3.2), we decided to use RCP models by season 212 and data type (presence-absence and abundance). RCP models were used to relate the response variables 213 (i.e. presence-absence or abundance of each seabird species in each grid cell) to the independent predictor 214 covariates (i.e. averaged values of environmental variables in each cell). Presence-absence data were modelled 215 based on a Bernoulli model, whereas abundance data were based on a negative binomial model. RCPs were 216 fitted using the 'ecomix' 1.0.0 package (S. Woolley et al., 2022), with the following workflow: (i) find the best 217 number of groups (K) that describes the data, (ii) re-fit the model by fixing K and selecting the optimal 218 combination of covariates, (iii) check residuals, (iv) bootstrap the best model to estimate uncertainty of 219 parameter estimates, and (v) produce results. 220 To identify the best number of RCP groups (K) that described the data, we used full models (all covariates) 221 with 100 random start numbers (see Foster et al., 2013) between k = (2, ..., 7) groups, and chose the number 222 of RCPs that minimized the Bayesian Information Criterion (BIC). Then, we fixed K at the optimal number 223 and fitted models for every possible linear, additive combination of covariates. We chose the best model as 224 the one with the lowest BIC. Visual inspection of randomized quantile residual plots modified for mixture 225 models was used to verify model assumptions (Dunstan et al., 2013). To obtain estimates of uncertainty for 226 model parameters (confidence intervals), we used the default settings of 'ecomix::regional_mix.bootstrap()', 227 which applies a Bayesian bootstrap 1000 times. A custom code was written to generate partial plots for 228 exploring the influence of covariates on RCP membership.

We built a seasonal 11-year (2011–2021) average layer for predicting model results for each environmental covariate. These variables were then averaged within 1° latitude × 1° longitude grid cells, by season, between 9–47°S and 141–161°E. Note that 'clim_eke_mean' and 'clim_sst_grad' were processed beforehand as they may characterise persistent oceanographic features (described above) and thus represent the same value as 'eke' and 'sst_grad'. Data for prediction were standardised using the same mean and standard deviation from the modelling dataset.

Finally, we ran rarefaction curves based on the presence-absence matrix and the assigned RCP group for each sample (grid cell) to assess if the amount of information in the RCP model was representative. Rarefaction curves were fit using the 'iNEXT' 3.0.0 package (Chao et al., 2014; Hsieh et al., 2016), and results are presented based on alpha diversity (i.e. species richness).

240 3 Results

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3.1 Summary of species richness and numbers

A total of 10,261 seabird records was obtained during the 15 voyages, comprising 142,646 individuals from 242 80 species, belonging to 11 families and five orders. The five most abundant species made up 83% of total 243 birds counted (43.0% short-tailed shearwater Ardenna tenuirostris, 17.2% fairy prion Pachyptila turtur, 244 14.3% wedge-tailed shearwater Ardenna pacifica, 5.2% sooty tern Onychoprion fuscatus, 3.2% shy albatross 245 Thalassarche cauta). The five most frequently encountered species were A. pacifica (frequency of occurrence 246 =35.3%), T. cauta (33.2%), brown booby Sula leucogaster (25.4%), red-footed booby S. sula (25.4%) and 247 black-browed albatross T. melanophris (25%) (Figure S1). An increase in species richness towards the south of the study region was clear in all but the winter season (Figure S2). Noticeably higher counts were made around Tasmania, possibly due to the 'colony effect' and/or the presence of Southern Ocean species 250 (Figure S2). However, in summer, one cell in the Coral Sea had counts of the same order of magnitude as 251 off Tasmania. 252

253 3.2 RCP models

For all models, two RCPs was the optimal number of seabird groups, except for the presence-absence-autumn model, which indicated three RCPs as the best fit (**Figure S3**). The best fitting models generally had one or two covariates selected, although the abundance-autumn model had five covariates (**Table 2**). The best models showed a good fit with no apparent deviations from the assumptions (**Figure S4**). All the best models selected sea surface temperature ('sst') or salinity ('sss'), as essential for characterising water masses. The thresholds for defining RCP membership were 18–21°C for 'sst' and 35.1 PSU for 'sss' (**Figure 2**, **Figure S5**), similar to the characteristics used to to define the EAC. Furthermore, most of the best models selected bathymetry ('bat'), suggesting a gradient between 'coastal' and 'oceanic' taxa (**Figure S5**). Of particular

262 note, proxies for persistent oceanographic features that may aggregate food resources (the 'climatic' variables)
263 for seabirds were selected only in the autumn and spring abundance models.

Grid point-predictions are the assignment of the RCP with the highest probability value for a given cell. 264 When predicting RCP onto space, patterns of point-predictions suggest a biogeographic transition in seabird 265 species around ~34°S, with both presence-absence and abundance models pointing to similar results (Figure 3). These patterns are further reinforced when looking at the mean probability along with the lower and 267 upper confidence intervals of predictions (Figure 4, Figure S6), showing that around ~34°S, the 'northern' 268 RCP tended to have higher probabilities in coastal regions, whereas the 'southern' RCP spread northward 269 offshore. The autumn presence-absence model (three RCPs) had a similar broad-scale pattern, although the 270 southern RCP is farther south, and a strong 'Tasmania/Bass Strait' assemblage appears (Figure 3, Figure 271 S6). 272

The broad pattern of two large-scale assemblages is reflected in their species compositions. Although there are overlaps among the seabird groups, both presence-absence and abundance models captured typical tropical species in the northern RCP (e.g. boobies *Sula* spp., lesser frigatebird *Fregata ariel*) and typical Southern Ocean species in the southern RCP (e.g. small albatrosses *Thalassarche* spp., fairy prion *P. turtur*) (**Figure** 5, **Figure S7**). The autumn presence-absence model, which had three RCPs, also showed the characteristic taxa representing the 'northern' and 'southern' RCPs. However, the intermediate 'Tasmania/Bass Strait' assemblage showed a mixture of both groups with shy albatross and Australasian gannet *Morus serrator* prominent with high probabilities of occurrence.

281 4 Discussion

This study showed that seabird assemblages off eastern Australia were correlated with oceanographic conditions, and that the East Australian Current appeared to be key in structuring seabird distributions at meso- to 283 macro scales (sensu Hunt & Schneider, 1987). Our results are based on a multivariate statistical model (N. A. Hill et al., 2020), which brings advantages over classical methods for describing species assemblages. RCP 285 models allow estimating uncertainty around the point probability of a sample (grid cell) belonging to an 286 assemblage, thus representing the fuzziness between bioregion boundaries. These transition zones between 287 bioregions are essential for monitoring biological changes due to environmental changes (Morrone, 2023). As 288 such, the boundary between the two consistently identified bioregions in this study may be a crucial region to 289 monitor in future, given noticeable anomalies in the EAC at this same region (Phillips et al., 2022). Our 290 results draw an important baseline for seabird assemblages in a world hotspot for seabirds.

292 4.1 Seabird bioregions off eastern Australia

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Hunt, 1993; Pocklington, 1979; Ribic et al., 1997). Thus, it is not surprising that two seabird assemblages 294 were identified off eastern Australia. However, the spatial consistency of these patterns throughout seasons is 295 surprising and remarkable. Resident species seem to characterise these assemblages. Shy albatross (Hedd et al., 2001; Mason et al., 2018) and fairy prions (Fromant et al., 2022) in the southern assemblage, and boobies (Bunce, 2015; Miller et al., 2018) and wedge-tailed shearwaters (McDuie et al., 2015; Weimerskirch 298 et al., 2020) in the northern assemblage, were always profiled with higher probabilities of occurrence and 299 predicted abundances compared to other species in the same groups. Given that changes in assemblages over 300 time are typically driven by a few species (Gotelli et al., 2022), and that those species are likely residents 301 (Souza & Santos, 2023), the relationships between these resident seabirds and the EAC system over the 302 broad latitudinal range detailed here should be closely monitored. Elsewhere, coastal seabirds such as little 303 penguin are signalling the adverse effects that climate change might have in changing seabird behaviours 304 and decreasing foraging success (Carroll et al., 2016; Woehler & Hobday, 2023), which ultimately influence 305 their breeding success and population dynamics (B. Johnson & Colombelli-Négrel, 2021). Thus, the species 306 profiles and the spatial predictions of assemblages from RCP models could be used as tools for monitoring 307 future seabird responses to the fast-changing EAC. 308 The seasonal models also captured the influence of migratory species, particularly in the southern assemblage. 309 The summer and autumn models reflected the high abundance of short-tailed shearwater before they depart 310 on their trans-equatorial migration (Carey et al., 2014); the presence of the Australasian gannet was picked 311 up by the autumn and winter models (Ismar et al., 2011); and the white-headed petrel Pterodroma lessonii 312 movements during chick-rearing and pre-laying exodus (Taylor et al., 2021) were detected in autumn and 313 spring models, respectively. Given the geographic location of the southern assemblage, however, we believe 314 sub-Antarctic taxa were not well represented in our models due to their low number of occurrences and 315 relative paucity of survey effort to date (Figure S1; hence they were removed before modelling). Additional surveys may reveal the seasonal northward pulse of these taxa into the southern assemblage (e.g. albatrosses 317 Diomedea spp. and Phoebetria spp.), given that more effort translates as higher likelihood of recording these 318 species. In turn, including these taxa in the models could result in a more stable 'three assemblage pattern' 319 (as revealed by the autumn presence-absence model) in all but the summer season, when the majority of 320 Antarctic species are breeding (S. Marchant & Higgins, 1990). 321 Our analyses clearly revealed a 'northern' and a 'southern' assemblage of seabirds off eastern Australia within 322 the EAC. There is a noticeable link with the EAC dynamics: the boundaries of RCPs occurred at ~34°S, near 323 the average separation latitude (~32°S; 25–38°S) of the EAC and its eastward extension into the Tasman 324 Sea (Oke, Roughan, et al., 2019). However, the 'northern' southern' assemblage pattern identified herein

The relationships between seabird assemblages and water masses has been known for a long time (Elphick &

does not agree with previously-proposed large-scale regionalizations, such as the pelagic Longhurst Provinces 326 (Longhurst, 2007), the Marine Ecoregions of the World [MEOW; Spalding et al. (2007)], or the Large Marine 327 Ecosystems [LME; Sherman (1994)]. The MEOW schema was based on coastal and shelf biota and was 328 meant to represent habitats to the 200 m isobath, which clearly does not encompass the entirety of seabird 329 habitats. The LME schema is characterised by physiographical, hydrological and productivity data but has a strong focus on practical management of areas and thus there is a political component embedded within 331 it, which does not align with seabird distributions. Longhurst (2007) designated only one biogeochemical 332 province for eastern Australia based on productivity patterns, which may work well at larger (macro) scales, 333 but appears to be oversimplified for the EAC. 334 The results do agree with the marine biogeographic units proposed by Costello et al. (2017), who identified a 335 boundary at ~34°S between their 'Coral Sea' and 'South Australia' realms. Their approach used 65,000 taxa. 336 from benthic/pelagic and coastal/offshore habitats, thus integrating more complexity of the biogeographic 337 boundaries in the marine environment. Other biological data also suggest 'northern' assemblages 338 off eastern Australia, including tuna and billfish [also highly mobile top predators; Reygondeau et al. (2012)],

the Longhurst Provinces approach was updated to include SST, SSS and bathymetry (the same variables retained by our best RCP models) and a temporal component to the classification (Reygondeau et al., 2013). 344 The updated provinces are more similar to the seabird assemblages described here, although still not a perfect 345 match, which could be partially explained because the seabird assemblage in the EAC region includes several 346 long-distance migratory species. Australia's bioregionalization has a long story of disagreements (Ebach, 2012), however, the inherent link that RCPs develop between environment and biological data can be a strong 348

mesopelagic fauna [which includes seabird prey; Sutton et al. (2017)] and deep-sea invertebrates and fish

(O'Hara, Williams, Althaus, et al., 2020; O'Hara, Williams, Woolley, et al., 2020). All these studies focused

mainly on describing biological assemblages, as opposed to classifying environments. It is worth noting that

Fitting RCP models to highly mobile animals 4.2

tool to support marine bioregionalization.

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To the best of our knowledge, this is the first study applying RCPs to highly mobile animals such as seabirds. 351 RCP models have been previously applied to plants (Lyons et al., 2017), and relatively sedentary animals, 352 such as macrobenthic organisms (Pantiukhin et al., 2021) and demersal fish (N. A. Hill et al., 2017). Oceanic 353 seabirds can have daily movements of hundreds of kilometres (e.g. McDuie et al., 2015), during which they 354 traverse a variety of habitats and environmental gradients. As such, capturing the distribution of highly 355 mobile species, and their relationships with environmental parameters is not a trivial task, and much harder 356 than for sessile organisms. 357

While remote tracking studies are advancing our understanding on how individual seabirds use their envi-358 ronment at fine scales (Tremblay et al., 2009), they fail to perceive how species assemblages interact with

the environment and each other. Therefore, using at-sea counts of seabirds is a well established method 360 to capture these relationships (Ainley et al., 2012; Hyrenbach et al., 2007; Woehler et al., 2003). Further, 361 the spatial scales on which assemblages relate to their environment is variable and species-specific (Hunt & 362 Schneider, 1987; Ribic et al., 1997; Scales et al., 2017). Understanding the biology of the studied group is 363 thus essential to set the appropriate scales of analyses. Our choice of 1° latitude \times 1° longitude was mainly determined by data availability reflecting survey effort, and the trade-off between the number of species per 365 grid/season (see also Section 4.3). Nonetheless, the chosen spatial scale is similar to other seabird distribution 366 analyses conducted elsewhere (Harris et al., 2007; Santora & Veit, 2013; Sojitra et al., 2022). Therefore, for 367 describing seabird assemblages at a macro-scale, the chosen resolution seems to have been a good fit, accords 368 with studies elsewhere, and the resultant models described patterns of seabird species' composition consistent 369 with our current understanding of species' ecologies and life history strategies. 370

RCP models based on presence-absence and abundance data are able to complement each other. Previous 371 studies using RCPs have opted to use only one type of response data. Lyons et al. (2017), N. A. Hill et al. 372 (2017), and Pantiukhin et al. (2021) all used presence-absence data, whereas Lee et al. (2019) and Receveur et al. (2020) used abundance (count) data. The added information contained in abundance data make them 374 inherently more powerful than binary presence-absence data. However, the use of different data inputs in 375 multivariate biodiversity assessments and assemblage descriptions has shown contrasting patterns at different 376 spatial scales. At larger spatial and temporal scales, presence-absence data describe similar patterns to 377 abundance data (Anderson et al., 2005), as we found. In contrast, at smaller scales, abundance data may 378 highlight local conditions and therefore reveal hidden patterns at coarse to fine scales (Anderson et al., 2005; 379 R. Marchant, 1990; Melo, 2005; Waters, 2008). If combined with ecological knowledge of a particular group, 380 the information contained in both presence-absence and abundance data could be leveraged to better interpret 381 the resulting patterns. For example, seabirds will likely occur in specific water masses or currents (Hunt 382 & Schneider, 1987; Ribic et al., 1997), whereas their abundance is related to coarse- to fine-scale processes 383 that enhance prey availability (Hunt & Schneider, 1987; Weimerskirch, 2007). Future studies assessing 384 bioregions of highly mobile animals could describe the compositions of assemblages using smaller resolution 385 grid cells and presence-absence models. Then, abundance models using higher resolution grid cells could 386 improve our understanding of how these animals relate to their environment and possibly reveal multi-species 387 aggregation areas (therefore 'new bioregions') likely nested within the bioregions described at large scales. 388 Such a hierarchical approach would require substantial concurrent observational and environmental data sets, 389 which we suspect do not exist at present.

391 4.3 Limitations

Although the results suggest consistent patterns, there are a number of caveats to our analyses. Seabird data were gathered during multi-disciplinary cruises, whose sampling designs were not specifically planned to

account for spatial ecology of seabirds. As a result, there are spatial discontinuities in the data, which might 394 have limited the ability of our models to detect the relationships between the seabird assemblages and the full 395 spectrum of environmental predictors. Even though Australia is a hotspot for seabirds (Karpouzi et al., 2007; 396 Ramírez et al., 2017), there is a paucity of information on seabird distributions in the region (Bernard et al., 2021; Mott & Clarke, 2018). Dedicated seabird surveys would help to advance seabird research in this region, and improve our understanding of the patterns described here. By increasing the amount of data, future 399 studies could set the grid cells at higher resolution, and potentially reveal assemblages at finer spatial and 400 temporal scales. In addition, ideally, we would have had more information within each grid cell by season 401 (our sampling units). Nonetheless, rarefaction curves show that the sample coverage (see Chao & Jost, 2012) 402 was high at the RCP level, although there is room for detecting more species in most of them (Figure S8). 403

404 5 Conclusions

We applied a robust, repeatable method to assess seabird assemblages off eastern Australia within the EAC 405 for the first time. The models point toward a consistent pattern of a 'northern' and a 'southern' seabird 406 assemblage, where the spatial division is attributable to the eastward separation of the EAC from the coast, 407 and that SST seems to be the key variable discriminating these groups. The EAC has penetrated poleward 408 over the past half-century (K. L. Hill et al., 2008; Li et al., 2022), a trend which is anticipated to continue 409 into the future (Yang et al., 2016), and extreme weather events such as marine heatwaves to happen more 410 often (Oliver et al., 2018). As such, seabirds will have to respond to these changes (Woehler & Hobday, 2023). 411 Southern Ocean albatrosses off southeast Australia, for instance, have shown 'early stages of a southward range shift' due to environmental variability (Sojitra et al., 2022). Resident species such as little penguins 413 and wedge-tailed shearwater change their behaviour and have decreased foraging success when SST is higher 414 (Carroll et al., 2016; Peck et al., 2004). Given that eastern Australia is a biodiversity and numerical hotspot 415 for seabirds, it is imperative to establish a monitoring program allowing the assemblages and species profiles 416 presented here to be used as the baseline for understanding possible effects of a warming ocean on seabird 417 distribution. In particular, the transition zone between assemblages (Morrone, 2023), associated with the 418 eastern extension of the EAC, may be fundamental to effectively monitoring strengthening of the EAC and 419 the subsequent impact on marine biota. 420

421 6 References

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Data Accessibility Statement

- 821 The gridded data used as the basis for all analyses are available in the archived repository along with the code
- 822 to reproduce the results. The seabird raw data are currently being used for other projects and thus can not
- be publicly available. If you are interested in the seabird raw data, please contact XXX. The environmental
- data used in this study are freely available from the source (see Table 1 in the main text).

825 Tables and Figures

- TABLE 1. Environmental data used in the Region of Common Profile models applied to seabird data off eastern Australia, their spatial/temporal resolutions, units, and sources
 - TABLE 2. Full (a) and best (b) Region of Common Profile Models applied to seabird data off eastern Australia, for both data types used (presence-absence, based on a Bernoulli model; abundance, based on a Negative Binomial model). Refer to Table 1 for the environmental data acronyms
 - FIGURE 1. Study area showing the main ocean currents (a), and seabird records made by season on top of the 1° latitude × 1° longitude grid cells (b). In (a), the East Australian Current (EAC) system is highlighted with its acronyms in bold and drawn in blue. The South Equatorial Current (SEC) and its branches, the North Caledonian Jet (NCJ) and South Caledonian Jet (SCJ), are drawn in red. The EAC 'eastern extension' is an eddy field (blue shaded area) often called the 'Tasman Front', from which the East Auckland Current originates (EAUC, in green). Figure (a) is adapted after Oke, Roughan, et al. (2019); for colour references see the online version
 - FIGURE 2. Partial plots for the retained covariates in the best spring (a) presence-absence and (b) abundance models. The plot shows the fitted probability of belonging to a Region of Common Profiles (RCP) against the environmental value. All other predictors are held at their mean values. Results for other seasons and models can be found in the Supplementary Material (Figure S5). Refer to Table 1 for the environmental data acronyms
 - FIGURE 3. Spatialized point predictions (hard-classes) for seabird assemblages (Region of Common Profile; RCP), by season, based on presence-absence (a) and abundance (b) models, off eastern Australia. Each grid was assigned to the RCP group with highest membership probability
 - FIGURE 4. Predicted probability membership of each seabird assemblage (Region of Common Profiles; RCP) and grid, off eastern Australia, from spring presence-absence (a) and abundance (b) models. The central column, 'mean', corresponds to the point prediction and Bayesian boot-strapped, lower and upper confidence intervals (CI), on its sides. Note that, in case of two RCPs, RCP2 probability equals (1 RCP1 probability). Results for other seasons and models can be found in the Supplementary Material (Figure S6)
 - FIGURE 5. Species profiles of seabird assemblages (Region of Common Profile; RCP) off eastern Australia in spring. Values are the average and 95% confidence intervals of probability of occurrence (a) and predicted mean abundance (b) for each species, based on 1000 Bayesian bootstraps. Values in (b) were log10-transformed to accommodate the high variation between species. Results for other seasons and models can be found in the Supplementary Material (Figure S7)

Table 1:

Variable	Abbreviation	Units	Spatial.resolution	Temporal.resolution	Source
Sea Surface Temperature	sst	°Celsius	$0.25^{\circ} \times 0.25^{\circ}$	daily	Huang et al. $(2021)^a$
Sea Surface Temperature gradient	sst_grad	Δ °Celsius	$0.25^{\circ} \times 0.25^{\circ}$	daily	Derived from Huang et al. $(2021)^a$
Sea Surface Salinity	SSS	PSU	$0.083^{\circ} \times 0.083^{\circ}$	daily	E.U. Copernicus Marine Service Information bc
Mixed Layer Depth	mld	m	$0.083^{\circ} \times 0.083^{\circ}$	daily	E.U. Copernicus Marine Service Information ^{bc} ERDDAP/NOAA ^d
8-day average Chlorophyll-a	chl8	$\mathrm{mg/m^3}$	$0.036^{\circ} \times 0.036^{\circ}$	8-days composite	
Log10 8-day average Chlorophyll-a	$\log 10 _{\rm chl} 8$	$log_{10}({ m chl8})$	$0.036^{\circ} \times 0.036^{\circ}$	8-days composite	Derived from ERDDAP/NOAA d
Eddy Kinetic Energy Climatic mean EKE	eke clim_eke_mean	m/s mean(m/s)/season/10 years	$0.2^{\circ} \times 0.2^{\circ} \\ 0.2^{\circ} \times 0.2^{\circ}$	daily seasonal climatology	Derived from ERDDAP/NOAA ^d IMOS/AODN ^e Derived from IMOS/AODN ^e Derived from
Climatic standard deviation EKE	${\rm clim}_{\rm eke}_{\rm sd}$	sd(m/s)/season/10 years	$0.2^{\circ} \times 0.2^{\circ}$	seasonal climatology	TA COC / A ODATA
Climatic mean SSTgrad	${\rm clim_sst_grad}$	$\mathrm{mean}(\Delta^{\circ}\mathrm{Celsius})/\mathrm{season}/10~\mathrm{years}$	$0.25^{\circ} \times 0.25^{\circ}$	seasonal climatology	Derived from Huang et al. $(2021)^a$
Bathymetry	bat	m	$0.004^{\circ} \times 0.004^{\circ}$	NA	GEBCO^f
Slope	slope	degrees (°)	$0.004^{\circ} \times 0.004^{\circ}$	NA	GEBCO ^f Derived from GEBCO ^f Calculated in R (see main text)
Distance from coast	dist_coast	km	NA	NA	Calculated in R (see main text)

^a NOAA OISST: (https://psl.noaa.gov/data/gridded/data.noaa.oisst.v2.highres.html);

^b 2011–2019: GLOBAL_MULTIYEAR_PHY_001_030 (https://doi.org/10.48670/moi-00021);

^c 2020–2021: GLOBAL ANALYSIS FORECAST PHY 001 024 (https://doi.org/10.48670/moi-00016);

^d ERDDAP/NOAA (https://coastwatch.pfeg.noaa.gov/erddap/griddap/erdMH1chla8day.html);

e IMOS/AODN (https://portal.aodn.org.au/search), 'OceanCurrent - Gridded sea level anomaly - Delayed mode' data set;

f GEBCO (https://www.gebco.net/data_and_products/gridded_bathymetry_data/)

Table 2:

b bat + sss 31	332.69 319.34 1184.29 1147.94 1089.43 1062.98
a bat + slope + eke + sst + sss + mld + sst_grad 33 Autumn a bat + slope + eke + chl + sst + sss + mld + sst_grad + clim_sst_grad 11 b bat + sst 11 Winter a bat + slope + eke + sst + sss + mld + sst_grad + clim_sst_grad 10 Spring a bat + slope + dist_coast + eke + chl + sst + sss + mld + sst_grad + 23	319.34 1184.29 1147.94 1089.43
b bat + sss 31 Autumn a bat + slope + eke + chl + sst + sss + mld + sst_grad + clim_sst_grad 11 b bat + sst 11 Winter a bat + slope + eke + sst + sss + mld + sst_grad + clim_sst_grad 10 b sst 10 Spring a bat + slope + dist_coast + eke + chl + sst + sss + mld + sst_grad + 23	319.34 1184.29 1147.94 1089.43
Autumn a bat + slope + eke + chl + sst + sss + mld + sst_grad + clim_sst_grad 11 Winter a bat + slope + eke + sst + sss + mld + sst_grad + clim_sst_grad 10 b sst 10 Spring a bat + slope + dist_coast + eke + chl + sst + sss + mld + sst_grad + 23	1184.29 1147.94 1089.43
a bat + slope + eke + chl + sst + sss + mld + sst_grad + clim_sst_grad 11 b bat + sst 11 Winter a bat + slope + eke + sst + sss + mld + sst_grad + clim_sst_grad 10 b sst 10 Spring a bat + slope + dist_coast + eke + chl + sst + sss + mld + sst_grad + 23	1147.94 1089.43
b bat + sst 11 Winter a bat + slope + eke + sst + sss + mld + sst_grad + clim_sst_grad 10 b sst 10 Spring a bat + slope + dist_coast + eke + chl + sst + sss + mld + sst_grad + 23	1147.94 1089.43
Winter a bat + slope + eke + sst + sss + mld + sst_grad + clim_sst_grad 10 b sst 10 Spring a bat + slope + dist_coast + eke + chl + sst + sss + mld + sst_grad + 23	1089.43
a bat + slope + eke + sst + sss + mld + sst_grad + clim_sst_grad 10 b sst 10 Spring a bat + slope + dist_coast + eke + chl + sst + sss + mld + sst_grad + 23	
b sst 10 Spring a bat + slope + dist_coast + eke + chl + sst + sss + mld + sst_grad + 23	
Spring a bat + slope + dist_coast + eke + chl + sst + sss + mld + sst_grad + 23	1062.98
$a bat + slope + dist_coast + eke + chl + sst + sss + mld + sst_grad + 23$	
	2345.6
b bat $+$ sst 23	2306.7
Negative Binomial	
Summer	
$a bat + slope + eke + sst + sss + mld + sst_grad 15$	1518.96
b bat $+$ sss 15	1505.13
Autumn	
$a bat + slope + eke + chl + sst + sss + mld + sst_grad + clim_sst_grad - Is$	Inf (n.f.)
$b bat + chl + sst + mld + clim_sst_grad 30$	3082.14
Winter	
$a bat + slope + eke + sst + sss + mld + sst_grad + clim_sst_grad 32$	3262.45
b sst 32	3234.21
Spring	
- •	5854.32
b sst + clim_eke_mean 58	



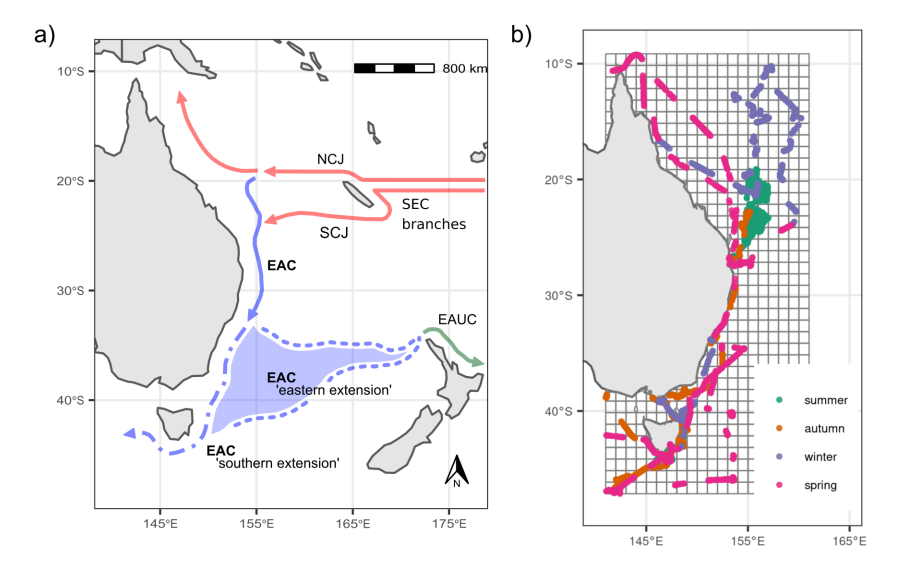


Figure 1:



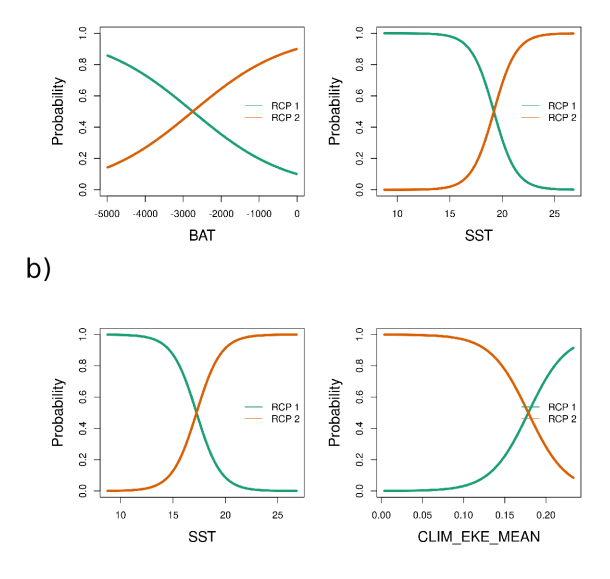
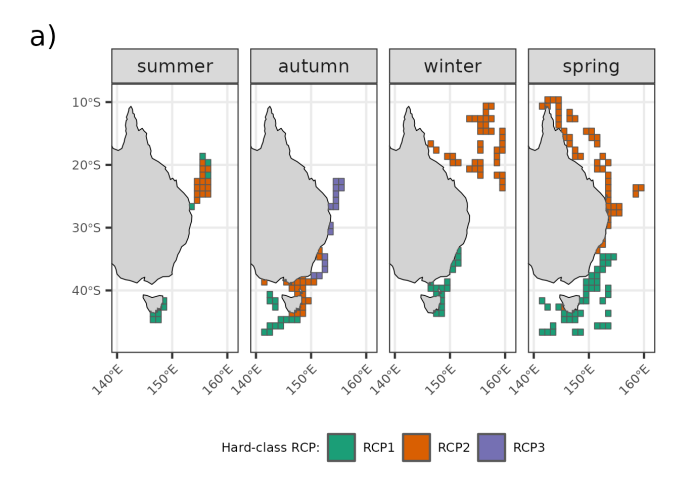
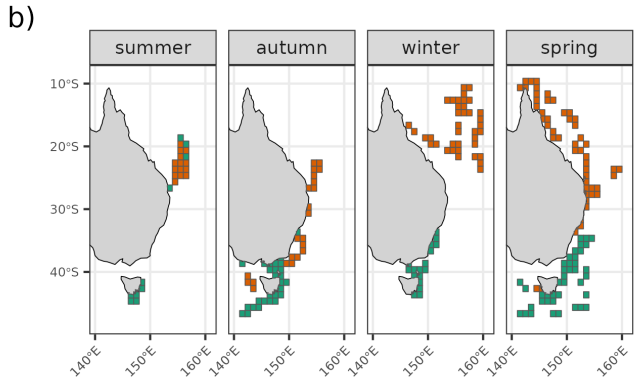


Figure 2:





 $\begin{array}{c} 35 \\ \text{Figure 3:} \end{array}$

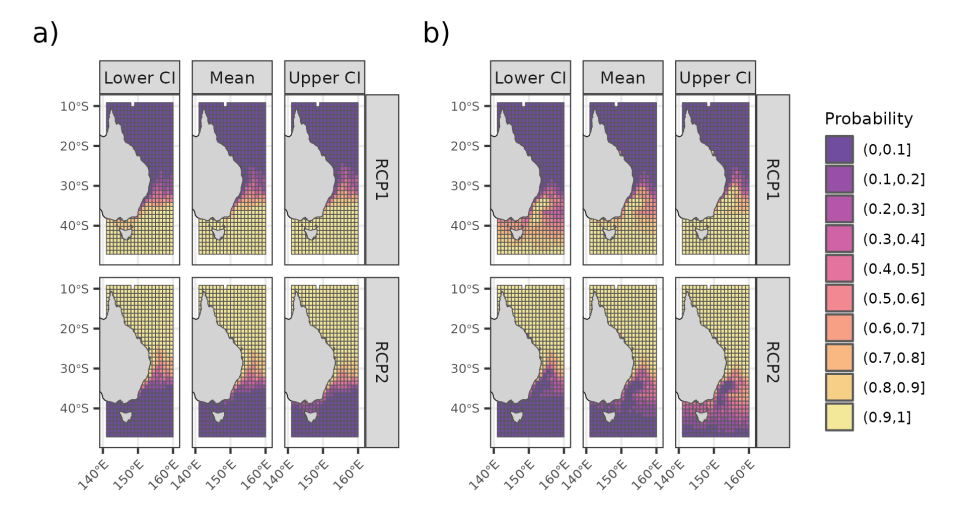
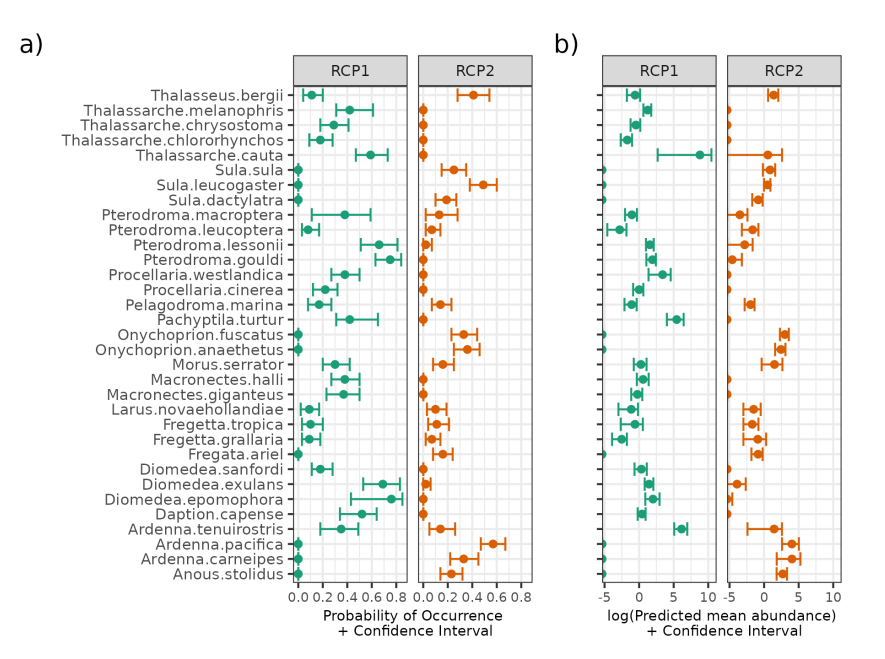


Figure 4:



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Figure 5: