Environmental correlates of taxonomic and phylogenetic diversity in the Atlantic Forest Running title (40 characters): Correlates of diversity in the Atlantic Forest

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1 Abstract

2 **Aim**

3 We compare patterns of diversity and their environmental correlates across nine clades of

4 ecologically distinct groups of animals and plants co-existing in a single rainforest domain. We

5 ask whether there are common correlates of diversity patterns, despite ecological differences

6 across clades, to enable a unified platform to predict changes in the distribution of biodiversity in

7 these groups. We focus on predictions of species richness, phylogenetic diversity, and

8 phylogenetic endemism.

9 Location

10 Brazilian Atlantic Forest

11 Methods

12 Using carefully curated occurrence localities and phylogenetic data, we generated maps of (i)

13 species richness, (ii) phylogenetic diversity, (iii) residuals of phylogenetic diversity regressed on

14 species richness, and (iv) phylogenetic endemism for nine groups of plants and animals in the

15 Atlantic forest. We also compiled a set of 30 potential environmental descriptors including

16 records of current temperature and precipitation, climatic stability over time, and topography.

17 Through a machine learning framework, we then explored the environmental correlates of each

18 of these diversity measures for each group.

19 **Results**

20 The environmental variables used in this study were strong predictors of diversity for all studied 21 groups. However, models for phylogenetic endemism had a lower predictive power. Although 22 patterns of diversity are different among groups, correlates of diversity are very much consistent 23 across taxa. For both species richness and phylogenetic diversity, current precipitation and 24 precipitation stability over time were constantly ranked among the variables that most strongly correlate with diversity patterns. Differently from species richness and phylogenetic diversity, 25 26 the correlates of phylogenetic endemism were less homogenous across groups. The results also 27 suggest that the inclusion of climate stability over time, along with current climatic descriptors, 28 is important when predicting diversity measures that reflect historical components, such as 29 phylogenetic diversity and endemism.

30 Main conclusions

31 Investigating environmental correlates of diversity for multiple co-existing clades and diversity

- 32 measures in a single geographic area allows for a better understanding of common patterns
- 33 across taxa. In this study, we identified common environmental correlates of the patterns of
- 34 species richness and phylogenetic diversity, but not of phylogenetic endemism, across different
- 35 Atlantic Forest groups. This information can now be used to improve predictions of biodiversity
- 36 changes at broad taxonomic and geographical scales in the Atlantic Forest.
- 37

38 Keywords

- 39 Biodiversity correlates, precipitation, climate stability, phylogenetic endemism, phylogenetic
- 40 diversity, species richness

42 Introduction

43 Over the last decade, stakeholders from governmental, academic, and conservation organizations 44 have shown a growing interest in the creation of systems that remotely monitor biodiversity over 45 broad spatial scales (Scholes et al., 2008, 2012). A call for the establishment of a Global 46 Observation Network (GEOBON) followed, along with a proposal to establish standardized 47 measurements of essential biodiversity variables (Scholes et al., 2012; Pereira et al., 2013). With 48 the increasing amount of satellite data being now freely available to the public, direct near-real 49 time monitoring of some components of diversity has become a reality (Turner, 2014). Examples 50 include global estimates of forest cover change (Hansen et al., 2013), plant functional diversity 51 (Jetz et al., 2016), and penguin population locations and sizes in Antarctica (Fretwell & Trathan, 52 2009). However, many of the world's diverse groups of organisms cannot be directly observed 53 through satellites, including most animals and non-canopy plants (Turner, 2014). Indirect 54 estimates of diversity would be valuable for more inaccessible organisms. For instance, 55 environmental variables that can be obtained from remote sensing sources (e.g., temperature, 56 precipitation) and correlate with diversity patterns, can be used as proxies to predict the patterns 57 of diversity themselves (Paz et al., 2020). Despite that, the utility of remote sensing tools for 58 indirect biodiversity monitoring remains underexplored.

59 One potential caveat of indirect sensing of biodiversity is that little consensus exists 60 regarding the environmental variables that represent good predictors of the many different 61 dimensions of biodiversity, especially in megadiverse and threatened tropical ecosystems. 62 Furthermore, the selection of input variables is important and often different between studies 63 (Williams et al., 2012). While species richness (SR) and endemism have been widely used as 64 biodiversity metrics, and broadly employed to describe its spatial patterns, phylogenetic diversity 65 (PD) and phylogenetic endemism (PE) are increasingly used to explicitly quantify the amount of 66 evolutionary uniqueness of a region (Vane-Wright et al., 1991; Faith, 1992; Rosauer et al., 67 2009). Both PD and PE reflect how combinations of more distantly related species will 68 encompass higher percentages of the overall evolutionary history than combinations of closely 69 related species (Forest et al., 2007; Devictor et al., 2010). Yet, measures of phylogenetic 70 diversity are often positively correlated with species richness. In other words, areas where these 71 two measures of diversity (taxonomic and phylogenetic) are decoupled, have been shown to 72 include more or less evolutionary history than expected given their species richness (Forest et al., 2007; Devictor et al., 2010; Safi et al., 2011; Fritz & Rahbek, 2012; Tucker & Cadotte, 2013).
Importantly, however, both species richness and evolutionarily informed measures of diversity
appear to be highly correlated to environmental variation - particularly temperature and
precipitation. Still, the specific contribution of the individual climatic or landscape descriptors
appears idiosyncratic when different taxa or different measures of diversity are compared across
regions (Rompré et al., 2007; Laurencio & Fitzgerald, 2010; Peters et al., 2016; Zellweger et al.,
2016).

80 One possible reason for the observed mismatch across systems and taxa is that few 81 studies have evaluated the correlates of species richness for multiple taxa occupying the same 82 ecosystem. For example, while local temperature has been flagged as a main predictor of species 83 richness in different groups of plants and animals along elevational gradients on Mount 84 Kilimanjaro (Peters et al., 2016), a similar transect study in Switzerland identified precipitation, 85 temperature, and topography as better predictors of the bird, plant, and butterfly species 86 diversity, respectively (Zellweger et al., 2016). In the tropics, herpetological surveys throughout 87 Costa Rica (Laurencio & Fitzgerald, 2010) indicated that topography is an important predictor of 88 richness, while a study of terrestrial vertebrates in Papua New Guinea recovered the same pattern 89 for all organisms studied, except for reptiles (Tallowin et al., 2017). The latter exemplifies the 90 importance of considering several groups when exploring environmental correlates of diversity.

91 Studies of evolutionarily-informed measures of diversity also result in contrasting 92 inferences. Humidity and precipitation are related to phylogenetic diversity in amphibian 93 communities in Brazil (da Silva et al., 2012). However, patterns of phylogenetic diversity of 94 northern Europe beetles are related to maximum temperature (Heino et al., 2015). Furthermore, 95 evolutionarily informed measures might be influenced by historical climates and, in particular, 96 by how much climatic variation an area has experienced. For example, long-term climatic 97 stability has been flagged as an important predictor of avian phylogenetic diversity at a global 98 scale (Voskamp et al., 2017). On the other hand, phylogenetic endemism of African frogs 99 appears to be related to Quaternary climatic stability (Barratt et al., 2017). Consensus is even 100 harder to achieve when including different dimensions of diversity such as richness and 101 phylogenetic diversity in the equation.

102 To provide a controlled comparison in the same area and inform the ability to indirectly 103 monitor tropical biodiversity as a function of climatic conditions in a biological hotspot, we

104 compare patterns of diversity and their environmental correlates across nine clades of 105 ecologically distinct groups of animals and plants in the Brazilian Atlantic Forest (AF). This 106 domain is a known biodiversity hotspot, harboring one of the highest levels of diversity and 107 endemism in the world (Ribeiro et al., 2009). It spans a region of complex topography and 108 environments, making it an excellent location to study the potential drivers of diversity in 109 dissimilar groups. We specifically ask if there is a set of environmental predictors that work 110 sufficiently well across different dimensions of diversity and taxa in the Atlantic Forest, which 111 could be useful in community-level prediction and indirect biodiversity monitoring.

112 We combine geo-referenced locality data with phylogenetic information for nine target 113 clades: five groups of plants and four groups of animals. We used these data to map species 114 richness, phylogenetic diversity, phylogenetic endemism, and the mismatch between 115 phylogenetic diversity and species richness (i.e., the residuals of their regression analysis). Using 116 a machine learning framework, we then investigated how well these patterns are predicted by 117 each one of 30 abiotic correlates obtained from weather-station data and remote sensing sources 118 (Vermote et al., 2015; Karger et al., 2017; Title & Bemmels, 2018), which describe spatial shifts 119 in temperature, precipitation, humidity, and topography, as well as climatic stability over the last 120 120,000 years.

121

122 Methods

123

124 Phylogenetic information

125 To generate maps of species richness (SR), phylogenetic diversity (PD), and phylogenetic 126 endemism (PE) for each target clade, we first obtained phylogenetic information and species 127 distribution data for nine biological groups (five plant clades and four animal clades). Two of the 128 plant datasets and all animal data were downloaded from Brown et al. (2020), including vetted 129 occurrence points and maximum likelihood phylogenies based on mitochondrial DNA for 130 animals and chloroplast DNA for plants. This dataset included a clade with 18 species (3,774 131 occurrence points; ca. 67% of the AF species included in this lineage) of tank-forming plants 132 belonging to the Bromelioideae subfamily (Aguirre-Santoro et al., 2016; Aguirre-Santoro, 2017), 133 a clade with 177 species (25,645 sampling points; ca. 70% of the AF species included in this 134 lineage) of shrubs and small trees from the Miconieae tribe in the Melastomataceae (Goldenberg

135 et al., 2008; Michelangeli et al., 2008; Caddah, 2013; Reginato & Michelangeli, 2016), a clade 136 with 55 species (3,269 occurrence points, ca. 100% of the AF species included in this lineage) of 137 clearwing butterflies of tribe Ithomiini in the subfamily Danainae (Nymphalidae), a clade of 19 138 species (227 occurrence points; ca. 76% of the AF species included in this lineage) of treefrogs 139 from the genus Boana from the family Hylidae, a clade of 19 species (2,065 occurrence points; ~ 140 67% of the AF species included in this lineage) of the horned frogs of the *Proceratophrys* genus 141 in the Odontophrynidae family (Brown et al., in press; Vasconcelos et al., 2014), and a clade of 142 22 species of birds (8,501 occurrence points; ca. 100% of the AF species included in this lineage) 143 from tanagers belonging to the subfamily Thraupinae (Burns et al., 2014). In addition, we 144 gathered three plant datasets (including species presence data and maximum likelihood 145 phylogenies produced with alternative markers): the *Fridericia* and allies group of the tribe 146 Bignonieae, in the plant family Bignoniaceae (hereon referred to as "bignones," Kaehler et al., 147 2019), and two clades of melastomes, the *Bertolonia* genus (Bacci et al., 2020), and the 148 Cambessedesieae tribe (Bochorny et al., 2019). The bignones dataset includes 65 species (ca. 149 72% of the AF species included in this lineage), 5,115 presence points, and a phylogeny built 150 from one plastid marker (*ndh*F) and one nuclear marker (*PepC*). The *Bertolonia* dataset contains 151 31 species (ca. 88% of the AF species included in this lineage), 744 points, and a phylogeny built 152 from nine nuclear, ribosomal and plastid markers (*nr*ITS and *nr*ET, *atp*F-*atp*H, *ndh*F, *psb*K-153 *psbL*, *rbcL*, *rpl*16, and *trnS-trnG*, *ADH*, and *PCRF1*). The Cambessedesieae dataset contains 54 154 species (ca. 81% of the AF species included in this lineage), 1,167 presence points, and a 155 phylogeny built from six ribosomal, plastid, and nuclear markers (nrITS, nrETS, *atpF-atpH*, 156 psbK-psbL, trnS-trnG, and waxy).

157

158 Mapping species richness, phylogenetic diversity, and phylogenetic endemism

159

To assess which environmental variables (reflecting both past and present conditions) and landscape descriptors best explain diversity patterns in the target clades, we first superimposed distribution maps for every species in each clade. To avoid circularity in testing for environmental predictors of biodiversity, we did not use correlative species distribution models for this step. Instead, we created alpha hulls (Burgman & Fox, 2003) using all occurrence data available for each individual species with more than three locality points (we added all others as 166 individual points). All occurrence data were vetted by the co-authors, who bring expertise in the 167 systematics and natural history of each of the groups sampled here. The alpha hulls were built 168 using the R package rangeBuilder (Rabosky et al., 2016) and a dynamic selection of alpha for 169 each species with alpha varying in steps of 1 (Meyer et al., 2017). Once done, those distribution 170 maps were rasterized to match the spatial resolution of the predictor variables (~10 km). We 171 acknowledge that this method might overestimate individual species distributions (or omit 172 unsampled populations), but such problems are more limiting at finer spatial scales (Peterson, 173 2017; Peterson et al., 2018).

174 Maps of species richness, phylogenetic diversity, and phylogenetic endemism were then 175 built for all nine groups, following the methods described in Brown et al. (in press). In brief, 176 input data consisted of community composition matrices based on the superimposed maps of 177 species ranges (alpha hulls and points when <= 3 points). For each group, the species maps were 178 stacked and converted to a composition matrix in R. For phylogenetic diversity and phylogenetic 179 endemism, we compiled phylogenetic trees, including branch lengths. For each of the nine 180 groups, we imported the community composition matrix and the phylogeny to Biodiverse 181 (Laffan et al., 2010). We then used the spatial analysis tab to calculate species richness, Faith's 182 PD index (Faith, 1992), and phylogenetic endemism (Rosauer et al., 2009) for every pixel in the 183 AF (Figure 1). Species richness was estimated by summing all species present in each pixel. 184 Phylogenetic diversity was computed by summing branch lengths leading to all species present 185 in a given pixel. Phylogenetic endemism, which combines endemism (estimated from the range 186 of the species and the fraction contained in a given cell), and phylogenetic diversity, to estimate 187 its level of restriction (Rosauer et al., 2009).

188

189 Concordance between measures: Mapping the residuals of phylogenetic diversity

190 Because measures of PD are highly correlated with SR, particularly Faith's PD (Forest et al.,

191 2007), we regressed those two maps (PD onto SR) and mapped the residuals of the regression for

192 each clade. The mapped residuals highlight areas where the information from these two diversity

193 measures is different. In the residual maps, values higher than 0 represent areas in which PD is

194 higher than expected given SR. Negative values depict areas with less PD than expected given

195 SR. Hereon, we refer to this variable for each clade as the PD residuals.

197 Environmental variables

198 To assess how much of the spatial patterns of SR, PD, and PE are explained by 199 environmental descriptors, we compiled environmental data for the entire extension of the forest. 200 We opted to use 30 variables, each one describing a climatic or landscape feature that has the 201 potential to correlate with local biodiversity metrics (Table 1). Twenty-one of those were directly 202 obtained from public databases. The variables include a 90m Digital Elevation Model (DEM) 203 from the Shuttle Radar Topography Mission (SRTM, Farr et al., 2007), the Topographic wetness 204 index (TWI) from Envirem (Title & Bemmels, 2018), and 19 bioclimatic variables reflecting 205 temperature and precipitation, downscaled to a 30" (~ 1km) resolution, using climatologies at 206 high resolution for the Earth's land surface algorithm (CHELSA; Karger et al., 2017) and the 207 ANUCLIM method (Xu & Hutchinson, 2010).

208 We generated the remaining nine Atlantic Forest specific layers, two describing terrain, one for 209 cloud cover, and six reflecting climatic stability over the last 120,000 years. The map of Atlantic 210 Forest Domain was created in a GIS environment based on the Vegetation Map of Brazil at a 211 1:5,000,000 scale (IBGE, 2004). Through an interactive discussion with experts having wide 212 experience in the field, known Atlantic Forest areas were selected and combined into a multi-213 polygon vector geometry. The map is referenced to the WGS84 datum with geographic 214 coordinates and is available in the Dryad repository associated to this manuscript. The first two 215 variables, Slope and Rugosity (calculated as the standard deviation of Slope, Grohmann et al. 216 2011), were derived from the DEM (SRTM, Farr et al., 2007). Mean cloud coverage was derived 217 from NASA's Moderate Resolution Imaging Spectroradiometer (MODIS 09GA, Vermote et al., 218 2015) based on satellite data collected from 2000 to 2017, using Google Earth Engine (Gorelick 219 et al., 2017). To determine whether historical climates were important in predicting present-day 220 patterns of diversity, we built six layers to reflect climatic stability over the last 120,000 years. 221 For that, we used existing bioclimatic descriptors available every 4,000 years for the past 222 120,000 years and obtained through the Hadley Center model (HadCM3, Singarayer & Valdes, 223 2010; Carnaval et al., 2014). We summarized long-term variation in three temperature attributes 224 [i.e., Annual Mean Temperature (bio 1), Mean Temperature of the Warmest Quarter (bio 10), 225 and Mean Temperature of the Coldest Quarter (bio 11)], and three precipitation attributes [i.e., 226 Annual Precipitation (bio 12), Precipitation of the Wettest Quarter (bio 16), and Precipitation of 227 the Driest Quarter (bio 17)]. For each one of the six variables, we computed the coefficient of

variation over the past 120,000 years. For downstream analyses, all variables were resampled to
a 5' resolution (~10km) using the resample function of the R package raster 3.0-7 (Hijmans,
2019).

231 The complete dataset of environmental descriptors (30 layers; Table 1) reflected variables 232 that may be highly correlated in the Atlantic Forest area. We thus ran a Variance Inflation Factor 233 (VIF) analysis to reduce collinearity, using the R package usdm 1.1-18 (Naimi et al., 2014), and 234 keeping only those variables with VIF<5. After eliminating the highly co-linear variables, we 235 were left with a dataset including 13 environmental descriptors, which were used in all machine 236 learning analyses. Together, these variables represent present-day climate (Mean diurnal range 237 (bio 2), Mean temperature of the wettest quarter (bio 8), precipitation of the wettest month (bio 238 13), precipitation of the warmest quarter (bio 18), and precipitation of the coldest quarter (bio 239 19)), climatic stability over the past 120,000 years (CV bio 1, CV bio 10, CV bio 16 and CV bio 240 17), topography (Altitude, Rugosity and TWI), and cloud cover (Table 1).

241

242 Correlates of biodiversity

243 To determine which of the environmental descriptors are the best predictors of SR, PD, PE, and 244 the residuals of the PD, we used four machine learning algorithms to generate correlative models 245 of each biodiversity metric. We then combined the four resulting models in an ensemble 246 prediction for each metric. The machine learning algorithms were Random Forests (rf from Liaw 247 & Wiener, 2002), Neural Network (nnet from Venables & Ripley, 2002), Support Vector 248 Machines (svmRadial from Karatzoglou et al., 2004), and Generalized Linear Models. While 249 running each algorithm, we randomly split each dataset (each map of a given diversity metric for 250 a given group) into two sets: one containing 70% of the pixels (for model training), and one 251 containing 30% of the pixels (completely withheld for model testing). For the training of each 252 model, we randomly split the training data into 10 subgroups (folds). We used each fold in turns 253 as an internal validation dataset, utilizing the others for training. We repeated this procedure 254 three times (repeat crossvalidation). All models were built with the R package caret 6.0-84 255 (Kuhn, 2016). A final ensemble model, built from a linear combination of the four algorithms 256 based on the RMSE values, was built with the caretEnsemble function in R. Finally, we used the 257 withheld 30% of pixels for model testing. To obtain an estimate of variable importance in the 258 ensemble model, we computed a weighted average of the variable contributions estimated from

the individual models, using the weight of the models in the ensemble. For that, we used the*varImp* function of the caret package for R.

- 261
- 262 **Results**

263 Patterns of SR, PD, and PE are different among groups (Figure 2). Bignones and 264 Cambessedesieae show a concentration of SR and PD in the interior forests from Bahia to Minas 265 Gerais, a result even more striking in the phylogenetic diversity maps (Figures 1, 2). Both the 266 Miconieae and tanagers have higher diversity (both SR and PD) in the Serra do Mar coastal 267 forest (coastal mountains, Figures 1, 2). On the other hand, the butterflies show peaks for both 268 PD and SR in the coastal forests from São Paulo to central Bahia. Bertolonia shows the opposite 269 pattern, with higher SR and PD both south (from São Paulo state to Santa Catarina) and north-270 east (Bahia coastal forest) of the high diversity areas for the butterflies. The bromeliads have 271 their peak diversity for both PD and SR in the north-east of the Bahia coastal forest, with the 272 *Proceratophrys* frogs showing peak diversity in the north of Bahia and in part of the Serra do 273 Mar coastal forest. Frogs of the genus *Boana* have two peaks of higher SR, one in the Serra do 274 Mar coastal forest, and a second one in the Bahia interior forests; with the highest PD mainly in 275 the Serra do Mar coastal forest.

276 In the case of PE, we detected two different general patterns with either small 277 concentrations of high PE or more widespread areas of high PE values (Figure 2). Two groups, 278 the butterflies and the tanager birds, show a pattern of widespread high PE, inland to the north of 279 Rio de Janeiro for the butterflies and in the Serra do Mar coastal forest for the tanager birds. 280 Both the bromeliads and the *Bertolonia* have very small areas of high PE in the north of the 281 Bahia state. The Miconieae, the Proceratophrys frogs, and the Cambessedesieae, all show high 282 PE in small areas of Espírito Santo and the border between Rio de Janeiro and São Paulo states. 283 The Cambessedesieae also shows a small area of high PE in the forests of inland Bahia. Finally, 284 frogs of the Boana genus and bignones show no apparent areas of high PE.

All groups show spatial concentration of residuals (positive or negative, Figure 3). Most groups have areas where PD is higher than expected given species richness, with *Proceratophrys* frogs showing this pattern along the entire mapped distribution (Figure 3). Butterflies have higher PD than expected given the number of species in the coastal region that extends from Alagoas to Paraiba, and in the southern interior region of the forest (red and dark orange areas in

290 Figure 3). We found a spatial concentration of higher PD than expected in the southern Atlantic 291 Forest for another three groups: (i) the tanagers in the Paraná, Santa Catarina, and part of São 292 Paulo states, (ii) both the Boana frogs, and (iii) bignones from the Serra do Mar Coastal Forest 293 and the Santa Catarina and Parana states. The bignones, Miconieae, and Cambessedesieae also 294 showed higher PD than expected in the north, around the Bahia Interior Forests region. Negative 295 residuals, differently from positive residuals, are more spread out in geographical space (Figure 296 3, green and blue respectively). The few exceptions, with a concentration of negative residuals, 297 are observed in tanager birds, butterflies, and the plant tribe Miconiae. For the tanagers, areas 298 holding less phylogenetic diversity than expected are concentrated in the north, mostly north of 299 Minas Gerais state, including the states of Espírito Santo and Bahia. In the butterflies, these areas 300 are found in the state of Bahia but also in small clusters in the southern portion of the forest. In 301 the Miconieae, they are mostly found in the Serra do Mar Coastal Forests (Figure 3, blue areas).

Of all biodiversity metrics, SR PD ($R^2 0.86-0.98$) and the residuals of PD ($R^2 0.86-0.98$, Figure 4) are best predicted by the environmental models. This metric is followed by SR (R^2 0.79-0.98. Model predictions of PE were more heterogeneous and generally lower ($R^2 0-0.96$, Figure 4), with three main exceptions: the bignones, the tanager birds, and the butterflies (R^2 of 0.94,0.95 and 0.96 respectively). Two other groups, the *Proceratophrys* frogs and the Miconieae (R^2 of 0.36 and 0.49 respectively), showed some predictive power.

The ability of the models to predict SR, PD, and PE also varied across clades. For instance, they were consistently high in butterflies ($R^2>0.96$), bignones ($R^2>0.93$), and tanager birds ($R^2>0.95$), but lower in the plants of the *Bertolonia* genus (lowest R^2 for all but PE and second to last for PE; Figure 4), and *Proceratophrys* frogs (which had one of the lowest R^2 values for predictions for SR, PD diversity and residuals; Figure 4).

313 Climatic variables, reflecting both present-day and past conditions, contributed highly to 314 predictions of SR, PD, and phylogenetic residuals. In particular, precipitation-related variables 315 were consistently identified as those of higher importance to predict SR and PD (blue in Figure 316 5). In eight out of the nine clades, variables reflecting current precipitation are those of highest 317 importance for predicting SR; in one clade (bromeliads), stability in past precipitation was 318 ranked first, but closely followed by current precipitation (Figure 5). Current temperatures were 319 the second or third predictors of SR. Conversely, historical stability in temperature contributed 320 less to predictions of diversity, ranking fourth to last for SR (except from bignones, which

ranked second) (Figure 5a). For PD, precipitation was of higher importance in seven out of the
nine clades. In one clade, the *Boana* frogs, past precipitation closely followed current
precipitation. For the bromeliads, current temperature was the most important variable, followed
by both current and past precipitation. For the other groups, current temperature ranked second,
third, or fourth, while temperature stability ranked third or fourth in importance (last for the
bromeliads; Figure 5b).

327 Variables related to precipitation also had higher importance to explain the residuals of 328 PD in eight out of the nine target groups (Figure 5a, b), with a slightly higher contribution of 329 climate stability as a correlate of the residuals relative to the other metrics (Figure 5a, b, d). The 330 importance of current temperatures as predictors of PD residuals was mixed, being ranked first in 331 the case of tanagers and butterflies, second in the case of Bertolonia, bromeliads and 332 *Proceratophrys*, and third to sixth in all other clades (Figure 5d). Although generally low, cloud 333 distribution was relevant to predicting the residuals for tanagers and *Boana* frogs, and slightly 334 relevant for the bignones and the Cambessedesiae (Figure 5).

335 Unlike the other biodiversity metrics, PE does not seem to be better predicted by one 336 specific type of environmental variable. Variables with highest contributions to the model are 337 related to current precipitation (five out of the nine groups), current temperature (one out of the 338 nine goups), and terrain (three out of nine). Of the groups with some predictive power for PE, 339 four out of five have current precipitation as the main predictor (bignones, birds, Miconieae, and 340 *Proceratophrys*); in one, topography was closely followed by current precipitation. Only in this 341 metric there is a more meaningful correlation with topography, which was recovered in at least 342 four groups. Topography is the highest contributor for patterns observed in Cambessedesiae, 343 bromeliads, and butterflies, but ranks second and third for *Bertolonia* and Miconieae, 344 respectively (Figure 5c).

345

346 Discussion

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Models based on environmental variables describing temperature and precipitation represented good predictors of different dimensions of diversity in the Atlantic Forest, based on the nine focal clades (Figure 4). Nevertheless, predictions of PE were poorer than those of SR or PD, a result likely associated to the spatial restriction of this biodiversity measure in relation to 352 the broader environmental predictors used, other studies have indeed suggested endemism might 353 be explained by variation within broader study regions (Rosauer et al., 2009; Crisp et al., 2011). However, predictions were still good in half of the cases, with R² values ranging from 0.36 to 354 355 0.96. In other words, to a certain extent, we can still predict PE based on our set of predictor 356 variables for some groups, although not as reliably. Our sampling for the AF was fairly 357 complete, however, some narrow endemics are missing in the datasets and thus PE can be 358 underestimated in certain cells. We acknowledge our threshold of >3 points for creating alpha 359 shapes might be considered small. However, we created the distribution maps with alternative 360 thresholds of 10 and 30 and have changes in only few groups of restricted distributions 361 (Appendix A). We also ran analyses with an alternative method for map building, through 362 minimum convex polygons with very similar predictive power and environmental predictors 363 selected (Appendix A). Sampling multiple clades within a single geographic space allowed us to 364 identify variables that are consistently important (or not) predictors of diversity in the Atlantic 365 Forest, highlighting the importance of precipitation (both past and present), but limited 366 contribution of topography.

367 Traditionally, temperature has been considered as the most important driver of diversity 368 patterns, given the importance of this variable in the temperate zones (Rohde, 1992; Erwin, 369 2009; Peters et al., 2016). However, our results point to precipitation as a main predictor of 370 species richness and phylogenetic diversity, while contributing to the mismatch between those 371 variables. This result is in line with other recent tropical clade-based studies that highlight the 372 importance of rainfall as a driver of species richness in small mammals (Mason-Romo et al., 373 2017), trees (Krishnadas et al., 2016), bats (Grimshaw & Higgins, 2017), fruit-feeding butterflies 374 (Santos et al., 2020), and anurans (Vasconcelos et al., 2010). Here we show that studying several 375 clades in the same area does improve our ability to find general patterns in the potential 376 environmental drivers of diversity, highlighting the importance of precipitation.

Contrary to our expectations of historical climatic stability and topography as strong
predictors of evolutionary history, PD was not explained by any of those variables. However,
climatic stability was an important predictor of the other two measures that reflect evolutionary
history (i.e., PE and residuals), with topographic variables contributing to the explanation of PE.
This result might be explained by the relative importance of evolutionary history in each
measure. In this case, PD is highly correlated with SR, and thus the predictors may also be driven

383 by the latter. The residuals, however, show areas of mismatch between the two, highlighting 384 areas where evolutionary history is providing different information (Forest et al., 2007; Devictor 385 et al., 2010), and might therefore give us better insights into predictors of evolutionary history. 386 This relationship has been studied at a global scale for birds, where abiotic correlates of these 387 residuals are spatially heterogeneous, suggesting that elevation is an important predictor of PD 388 residuals in the tropics, along with contact among biomes (Voskamp et al., 2017). Here, we 389 found relatively high importance of climatic stability as a predictor of PD residuals in the AF, a 390 tropical realm that may point to further heterogeneity in the relative importance of environmental 391 variables within realms.

392 We found some discrepancies among study groups in the strength of predictions and the 393 importance of variables. More specifically, *Proceratophrys* frogs showed the lowest predictive 394 power in all models, which might be linked to the more restricted distributions of its species. 395 Another example is the increased contribution of temperature stability (and sometimes 396 precipitation stability) over the Quaternary to explain PE in bromeliads, tanagers, and frogs. This 397 result is congruent with a previous study highlighting the importance of climatic stability for 398 predicting PE in African frogs (Barratt et al., 2017). These discrepancies suggest that 399 environmental correlates of diversity may be more similar among study groups with similar 400 natural histories, rather than region dependent. Indeed, life-history traits have been proposed to 401 explain differences in how shared barriers lead to different levels of intraspecific isolation (or 402 gene flow) in co-distributed species, as well as differences in the impact of climatic changes in 403 population history and demography (Pabijan et al., 2012; Paz et al., 2015; Zamudio et al., 2016; 404 Carstens et al., 2018).

405 By mapping diversity metrics of multiple taxonomic groups that co-occur in a single 406 domain, our approach led not only to strong predictions of different dimensions of biodiversity 407 based on past and present abiotic variables, but also highlighted the importance of precipitation 408 in determining diversity patterns. In the face of global climatic changes, a similar framework 409 may be useful as a biodiversity monitoring tool, particularly if tied to periodically retrieved 410 remote sensing data (e.g., Vermote et al., 2015). Akin to near-real time snapshots of habitat 411 change (Diniz et al., 2015), they can provide fairly accurate predictions of expected changes in 412 diversity patterns driven by climatic shifts. Our analyses suggest that these near-time models, at 413 least in the Atlantic Forest, and potentially in the tropics, will profit from the inclusion of

- 414 climatic data describing current and past precipitation, replacing the need to include topography
- 415 as an independent variable. Models may be refined for specific groups (or life-history
- 416 characteristics) and diversity measures including, for example, stability measures to predict
- 417 evolutionary history, or data on landscape configuration as an additional proxy to explain
- 418 diversity distribution (Santos et al., 2020).
- 419
- 420

421 **Table 1.** Thirty environmental variables compiled for this study. Grey shading indicates the

422 thirteen variables used for Machine Learning analyses, after eliminating variable with high

423 collinearity given the original 30-variable dataset. All variables were resampled at a ~10km

424 resolution. The table depicts the source and the category of each variable. Acronyms:

425 SRTM: Shuttle Radar Topography Mission (Farr et al., 2007); TWI: Topographic wetness

426 index; Environmental rasters for ecological modeling (Title & Bemmels, 2018);

- 427 CHELSA: climatologies at high resolution for the Earth's land surface algorithm (Karger et
- 428 al., 2017); MODIS: NASA's Moderate Resolution Imaging Spectroradiometer (Vermote et
- 429 al., 2015); Hadley Center: Hadley Center model (HadCM3, Singarayer & Valdes, 2010;
- 430 Carnaval et al., 2014).
- 431

Layer	Description	Source	Category
Bio 1	Annual Mean Temperature	CHELSA	
Bio 2	Mean Diurnal Range	CHELSA	
Bio 3	Isothermality (BIO2/BIO7) (* 100)	CHELSA	
	Temperature Seasonality (standard deviation		
Bio 4	*100)	CHELSA	
Bio 5	Max Temperature of Warmest Month	CHELSA	Current
Bio 6	Min Temperature of Coldest Month	CHELSA	temperature
Bio 7	Temperature Annual Range (BIO5-BIO6)	CHELSA	
Bio 8	Mean Temperature of Wettest Quarter	CHELSA	
Bio 9	Mean Temperature of Driest Quarter	CHELSA	
Bio 10	Mean Temperature of Warmest Quarter	CHELSA	
Bio 11	Mean Temperature of Coldest Quarter	CHELSA	
Bio 12	Annual Precipitation	CHELSA	
Bio 13	Precipitation of Wettest Month	CHELSA	
Bio 14	Precipitation of Driest Month	CHELSA	
	Precipitation Seasonality (Coefficient of		Current
Bio 15	Variation)	CHELSA	Precipitation
Bio 16	Precipitation of Wettest Quarter	CHELSA	
Bio 17	Precipitation of Driest Quarter	CHELSA	
Bio 18	Precipitation of Warmest Quarter	CHELSA	

Bio 19	Precipitation of Coldest Quarter	CHELSA	
Altitude	Digital elevation model (DEM)	SRTM	
Slope	Slope calculated from the DEM	Based on SRTM	Topographic
Rugosity	Rugosity calculated from the DEM	Based on SRTM	ropographie
TWI	TWI calculated from the DEM	Envirem	
	Average cloud coverage calculated from		Clouds
Clouds	MODIS imagery	Based on MODIS (2000-2017)	Clouds
	CV of Bio 1 (120,000 years, every 4,000		
CV Bio 1	years)	Based on Hadley Center data	
	CV of Bio 10 (120,000 years, every 4,000		
CV Bio 10	years)	Based on Hadley Center data	Temperature
	CV of Bio 11 (120,000 years, every 4,000		Stability
CV Bio 11	years)	Based on Hadley Center data	
	CV of Bio 12 (120,000 years, every 4,000		
CV Bio 12	years)	Based on Hadley Center data	
	CV of Bio 16 (120,000 years, every 4,000		
CV Bio 16	years)	Based on Hadley Center data	Precipitation
	CV of Bio 17 (120,000 years, every 4,000		Stability
CV Bio 17	years)	Based on Hadley Center data	

- 435 FIGURES
- 436 **Figure 1.** The study area, the Brazilian Atlantic Forest, including the Brazilian states
- 437 encompassed by the Atlantic Forest domain. Lighter shades of gray indicate lower
- 438 elevations, darker shades indicate higher elevations.
- 439



441 Figure 2. Maps of species richness (first column), phylogenetic diversity (middle column), and 442 phylogenetic endemism (last column) for the nine study groups (rows), based on individual 443 species minimum convex polygons around presence points, and molecular phylogenies. 444 Warmer colors represent higher diversity, colder colors depict lower diversity; for all maps, 445 values are stretched to maximum-minimum. Taxa represented from top to bottom are: five clades of plants: Bertolonia, Fridericia and allies (bignones), Bromelioideae. 446 447 Cambessedesiae, Miconieae, and four groups of animals: the Ithomiini butterflies, the 448 *Boana* and, *Proceratophrys* frogs, and the tanager birds of the Thraupinae subfamily. In the 449 species richness maps, the number of species for each group varies as follows: Bertolonia 450 (1-5), Fridericia and allies (bignones; 1-33), Bromelioideae (1-8), Cambessedesiae (1-8), 451 Miconieae (1-81), Ithomiini (2-39), Boana (1-7), Proceratophrys (1-5), and Thraupinae (1-452 19). For the phylogenetic diversity maps, the value of PD for each group varies as follows: 453 Bertolonia (0.11-0.21), Fridericia and allies (bignones; 0.04-0.23), Bromelioideae (5.37-454 15.48), Cambessedesiae (0.17-0.52), Miconieae (25.73-424), Ithomiini (0.14-1.52), Boana 455 (0.74-1.86), Proceratophrys (0.41-0.72), and Thraupinae (39.64-345.69). For the 456 phylogenetic endemism maps, the value of PE for each group varies as follows: Bertolonia (4.35*10⁻⁵-0.029), *Fridericia* and allies (bignones; 2.31*10⁻⁶-0.0013), Bromelioideae 457 (0.003-0.54), Cambessedesiae $(1.87*10^{-5}-0.041)$, Miconieae (0.001-3.64), Ithomiini 458 $(2.69*10^{-6}-0.0045)$, Boana $(6.17*10^{-5}-0.1)$, Proceratophrys $(1.9*10^{-5}-0.0074)$, and 459 460 Thraupinae (0.00034-0.046).



- 462 **Figure 3.** Maps of residuals of phylogenetic diversity regressed on species richness (PD
- 463 residuals) for the nine study groups. Values of positive residuals are shown through a red scale;
- 464 values of negative residuals are depicted through a blue scale. Taxa represented from left to right
- 465 and top to bottom are: five groups of plants (Bertolonia, Fridericia and allies (bignones),
- 466 Bromelioideae, Cambessedesiae, and Miconieae), and four groups of animals (the Ithomiini
- 467 butterflies, the *Boana* and *Proceratophrys* frogs, and the tanagers of the Thraupinae subfamily).







- 469 **Figure 4.** R² of ensemble machine learning models for each studied group and measure of
- 470 diversity. From top to bottom, diversity measures are species richness (SR), phylogenetic
- 471 diversity (PD), phylogenetic endemism (PE), and residuals of phylogenetic diversity regressed
- 472 on species richness (residuals). The study groups from left to right are Bertolonia, Fridericia and
- 473 allies (bignones), Bromelioideae, Cambessedesiae, Miconieae, Ithomiini butterflies, Boana,
- 474 *Proceratophrys* frogs, and the tanagers of the Thraupinae subfamily.



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Figure 5. Relative importance of predictors of diversity for all studied groups and measures of
diversity. Each panel corresponds to one measure of diversity a) Species richness (SR), b)
Phylogenetic diversity (PD), c) Phylogenetic endemism (PE), and d) Residuals of the
PD/SR regression. The 13 predictor variables are grouped in six categories, from top to
bottom: clouds, current temperature, current precipitation, precipitation stability,
temperature stability, and topography (more details in Table 1).



488 Data availability statement

- 489 All used raw data are available in the individual referenced publications. A table pointing to each
- 490 individual data source, all maps in raster format, the shapefile of the Atlantic Forest boundaries,
- 491 as well as scripts for analyses, can be found in Dryad [link to be added after acceptance].
- 492

493 **Author contributions**

494 AP, JLB, ACC and TSS conceived the ideas; AP and JLB analyzed the data with help from TSS; 495 all authors contributed data and participated in discussions that led to data analyses and 496 interpretations of results; AP and ACC led the writing, and all authors read and approved the

- 497 final version of the manuscript.
- 498

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728 Biosketch

- Andrea Paz is a PhD candidate at the CarnavalLab at the City College of New York. She is
- 730 interested in the geographical patterns of biodiversity and the processes generating and
- 731 maintaining those patterns. In particular, she seeks to understand how species change their
- 732 distributions in response to environmental changes both in the past and present.