



Similar but different: Revealing the relative roles of species-traits versus biome properties structuring genetic variation in South American marsh rats

Joyce R. Prado¹ | Alexandre R. Percequillo^{1,2} | Andréa T. Thomaz³ |
L. Lacey Knowles³

¹Departamento de Ciências Biológicas, Escola Superior de Agricultura 'Luiz de Queiroz', Universidade de São Paulo, São Paulo, Brazil

²Department of Life Sciences, The Natural History Museum, London, UK

³Department of Ecology and Evolutionary Biology, The University of Michigan, Ann Arbor, Michigan

Correspondence

Joyce R. Prado, Departamento de Ciências Biológicas, Escola Superior de Agricultura 'Luiz de Queiroz', Universidade de São Paulo, São Paulo, Brazil.
Email: joycepra@gmail.com

Present addresses

Joyce R. Prado, Departamento de Ciências Biológicas, Universidade Federal do Espírito Santo, Vitória, ES, Brazil.
Andréa T. Thomaz, Biodiversity Research Centre and Department of Zoology, University of British Columbia, Vancouver, BC, Canada.

Funding information

Fundação de Amparo à Pesquisa do Estado de São Paulo, Grant/Award Number: 2009/16009-1, 2012/24099-3, 2014/22444-0, 2016/20055-2

Editor: Camila Ribas

Abstract

Aim: Wetland habitats, and the ecological restrictions imposed by them, structure patterns of genetic variation in constituent taxa. As such, genetic variation may reflect properties of the specific biomes species inhabit, or shared life history traits among species may result in similar genetic structure. We evaluated these hypotheses jointly by quantifying the similarity of genetic structure in three South American marsh rat species (*Holochilus*), and test how genetic variation in each species relates to biome-specific environmental space and historical stability.

Location: South America.

Taxon: Rodentia.

Methods: Using complementary analyses (Mantel tests, dbRDA, Procrustes, covariance structure of allele frequencies and environmental niche models [ENMs]) with 8,000–32,000 SNPs per species, we quantified the association between genomic variation and geographic and/or environmental differences.

Results: Significant association between genetic variation and geography was identified for all species. Similarity in the strength of the association suggests connectivity patterns dictated by shared species-traits predominate at the biome scale. However, substantial amounts of genetic variation are not explained by geography. Focusing on this portion of the variance, we demonstrate a significant quantitative association between genetic variation and the environmental space of a biome, and a qualitative association with varying regional stability. Specifically, historically stable areas estimated from ecological niche models are correlated with local levels of geographic structuring, suggesting that local biome-specific histories affect population isolation/connectivity.

Main conclusions: These tests show that although species exhibit similar patterns of genetic variation that are consistent with shared natural histories, irrespective of inhabiting different wetland biomes, local biome-specific properties (i.e. varying environmental conditions and historical stability) contribute to departures from equilibrium patterns of genetic variation expected by isolation by geographic distance. The reflection of these biome-specific properties in the genetic structure of the marsh rats provides a window into the differences among South American wetlands with evolutionary consequences for their respective constituent assemblages.



KEYWORDS

climatic changes, environment, next-generation sequencing, phylogeography, rodent, wetland

1 | INTRODUCTION

In the tradition of using concordance across taxa to infer the common effects of abiotic factors in structuring genetic variation, comparative phylogeographic studies usually focus on a specific geographic region with expanded taxonomic sampling. The role of abiotic factors in structuring genetic variation can also be tested by controlling for the potentially confounding influence of biotic factors, for example, by studying a subset of species with similar biological traits, such as similar ecological or life history traits, within a given geographic region (Papadopoulou & Knowles, 2015). However, genetic variation may be shaped by species-specific traits in concert with the abiotic factors (Capurcho et al., 2013; Choueri et al., 2017; Massatti & Knowles, 2014), or regional differences in abiotic factors may mediate the effects of such factors on genetic variation (Papadopoulou & Knowles, 2016). For relatively understudied regions and taxa, accommodating these complexities becomes an important part of the study design because tests focused on a single component structuring genetic variation (i.e. abiotic as opposed to biotic factors) may be giving an especially incomplete picture given the lack of background information for determining appropriate hypotheses.

Here we test for the effects of both abiotic and biotic factors in structuring genetic variation of three species of marsh rats (genus *Holochilus*: Cricetidae) that are restricted to wetland habitats associated with different biomes throughout South America (Figure 1). Marsh rats are semiaquatic, large bodied rodents with a herbivorous diet (Gonçalves, Teta, & Bonvicino, 2015; Hershkovitz, 1955) that inhabit open and seasonally flooded grasslands. They share a number of characters associated with specialization for aquatic life and an herbivorous diet. These include not only morphological traits (fusiform body shape, webbed and elongate hindfeet, loss of molars lophes; Hershkovitz, 1962; Stein, 1988; Dominguez-Bello & Robinson, 1991) but also behavioural traits linked to foraging and how the mammals seek shelter, as well as where they mate (Eisenberg, 1981). As a function of these shared traits, they may also exhibit similar patterns of genetic variation—that is, biotic factors related to being adapted to wetlands may play a role in structuring the genetic variation across the species (Avice, 2000). However, the wetlands they inhabit differ. For example, two of the focal species (*Holochilus sciureus* and *Holochilus vulpinus*) inhabit patchy wetlands across forested biomes, whereas the other taxa (*Holochilus chacarius*) lives in an open contiguous wetland (Figure 1). The forested wetlands include wetlands distributed along rivers throughout the Amazon biome, and wetland patches throughout the Atlantic Forest, where *H. sciureus* and *H. vulpinus* are distributed respectively. However, *H. vulpinus* can also be found in the wetlands of a non-forested biome, the Pampas

(Figure 1). Unlike the forest-associated taxa, *H. chacarius* is distributed throughout the contiguous wetlands of Chaco/Pantanal biomes (Figure 1). These South American wetlands oscillate between flooded and unflooded phases, although the length, depth, frequency and timing of the phases vary across biomes (Junk, 2013).

In addition, the history of the wetlands differs. For example, the Pantanal is a biome whose formation is linked to precipitation changes since the Last Glacial Maximum (LGM; McGlue, Silva, Assine, Stevaux, & Pupim, 2015). For the forest-associated wetlands within the Amazon, while some riverine forest persisted during more arid periods of the Pleistocene (Erwin & Adis, 1982), *terra firme* vegetation expanded, replacing the flooded forests, and interrupting the distribution and availability of flooded environments (Choueri et al., 2017; Thom et al., 2018). Likewise, expansions and contractions of forested areas in association with climatic change and sea level shifts no doubt impacted the wetlands of the Atlantic Forest, although we lack information specific to the wetland habitats. With a history of climatic (or sea level)-driven change, the stability of wetland habitats varied not only across biomes but also within biomes (Bush, Silman, & Listopad, 2007; Quattrocchio, Borromei, Deschamps, Grill, & Zavala, 2008; Whitney et al., 2011), during the Pleistocene. As such, the genetic structure of the marsh rats may reflect biome-specific properties of the different wetland habitats—that is, abiotic factors related to the wetlands themselves that are extrinsic to the shared characteristics of the marsh rat taxa.

We approach the question of the effects of both abiotic and biotic factors in structuring genetic variation of three species of South American marsh rats by quantitatively assessing not only the similarity of genetic structure across species but also the correspondence between patterns of genomic variation and biome-specific aspects of the wetland habitats. Specifically, using 8,000–32,000 SNPs per species, we quantify (a) the similarity of genetic structure in three South American marsh rat species (*Holochilus*), and (b) test how genetic variation in each species relates to (i) environmental and (ii) historical stability differences across the biomes (i.e. biome-specific impacts on genetic structure of the marsh rat species). Such tests involve an evaluation of whether the data conform to levels of gene flow expected under an isolation by distance (IBD) model, where a similar correspondence between geographic and genetic data would suggest a similar migration history among the taxa reflecting the shared traits among the marsh rats (e.g. Peterson & Denno, 1998). In addition, the degree to which the biome-specific environmental properties and historical stability (as characterized by ecological niche models for the past and present) contribute to genetic structure properties is evaluated by multiple procedures for testing for a correspondence between the biome properties and deviations from IBD, including both quantitative and qualitative measures of association (He,



FIGURE 1 Examples of the open biomes of the central, southern and northern region of South America inhabited by *Holochilus* (see Fig. 2 for distributional map), such as (a) Poconé and (b) Miranda, in Pantanal biome, both habitats occupied by *H. chacarius*; (c) Caçapava do Sul in Pampa biome, and (d) São José do Norte in Atlantic Forest biome, within the range of *H. vulpinus*; (e) lower Xingú River, and (f) Japurá River both in Amazon biome, inhabited by *H. sciureus*; (g) *H. chacarius* [Colour figure can be viewed at wileyonlinelibrary.com]

Edwards, & Knowles, 2013; Knowles, Massatti, He, Olson, & Lanier, 2016).

Given the recent origin of the marsh rat taxa (Machado, Leite, Christoff, & Giuliano, 2013), these biome-specific properties, not more ancient Miocene and Pliocene climatic changes (Hoorn et al., 2010), are a logical starting point for testing the potential effects of biome-specific properties on genetic variation in constituent taxa. Moreover, with few information on the physiogamy of regions and potential impact of Pleistocene climatic changes on wetland habitats specifically (see Aleixo, 2006; Ledru, Montade, Blanchard, &

Hély, 2016; Leite et al., 2016; McGlue et al., 2015; Thom et al., 2018), differences in the genetic structure of constituent taxa across biomes, and any correspondence with regional properties can be mutually informative (e.g. Ferreira et al., 2018; Thom et al., 2018). As one of the first (and to our knowledge) only studies that brings the resolution of genomic data to these vast, but relatively understudied biomes, our findings provide insights about not only the history of the taxa but also the features of the different wetlands with evolutionary consequences for their respective assemblages.

2 | MATERIALS AND METHODS

2.1 | DNA extraction, amplification and sequencing

Genomic data were generated for 26 individuals and 9 populations of *H. chacarius*, 24 individuals and 6 populations of *H. sciureus*, and 18 individuals and 6 populations of *H. vulpinus*, sampled across their entire range (see Figure 2, Table S1.1, Appendix S1 in Supporting Information). Tissues were collected or requested from museums (see Appendix S2 in Supporting Information). DNA was extracted from liver, muscle or skin using the Qiagen DNeasy Blood and Tissue Kit.

Four reduced representation libraries were sequenced using ddRADseq method (Peterson, Weber, Kay, Fisher, & Hoekstra, 2012; Appendix S1). Briefly, DNA was double digested with the restriction enzymes EcoR1 and MseI and 150 base pair, single-end reads sequenced on four lanes of Illumina HiSeq2000, which produced 270 million raw reads at the Centre for Applied Genomics, Canada.

2.2 | Processing of illumina data and generating summary statistics

Raw sequence reads were processed separately for each species in STACKS 1.35 (Catchen, Hohenlohe, Bassham, Amores, & Cresko, 2013). The reads were demultiplexed and filtered using *process radtags*. Only reads with Phred score >10, unambiguous barcodes, and individuals with more than 500,000 reads were retained (additional details are given in Appendix S1). A de novo assembly of filtered reads with a minimum coverage depth of six were used to identify putative loci with the USTACKS. A catalogue of consensus loci was constructed for each species with CSTACKS with a distance between individuals for a given locus ≤ 2 . Alleles were called for loci using SSTACKS.

Characterizations of genetic variation and population summary statistics were generated with the POPULATIONS program (Catchen et al., 2013). At this step, all loci present in at least two populations were identified and exported in Variant Call Format (vcf). This first dataset (hereafter referred as the full dataset) was used to calculate population genetic diversity statistics, such as nucleotide diversity (π), major allele frequency, observed heterozygosity (H_O) and Wright's inbreeding coefficient (F_{IS}) at each locus. Population-level summaries of genetic diversity were also characterized (average π , H_O and F_{IS}), and a one-way ANOVA was used to test for significant differences in genetic diversity between species (conducted in R, 'Car' package; R Core Team, 2017). A second dataset with one randomly chosen SNP and a maximum of 20% missing data was generated with the toolset PLINK 1.07 (Purcell et al., 2007) and used for all other analyses (hereafter referred to as the putative unlinked-SNP dataset).

The full dataset consists of 359,728 SNPs and 26 individuals of *H. chacarius*, 357,050 SNPs and 24 individuals of *H. sciureus* and 160,879 SNPs and 18 individuals of *H. vulpinus*. The unlinked-SNPs dataset consists of 32,210 SNPs in 25 individuals of *H. chacarius*,

17,513 SNPs in 20 individuals of *H. sciureus* and 8,035 SNPs in 15 individuals of *H. vulpinus* (Table S1.2 in Appendix S1).

2.3 | Structuring of genetic variation

A series of complementary approaches were used to test for similar structuring of genetic variation across taxa and for a correspondence between properties of the wetland habitats and genetic variation. These tests were selected because they differ in their respective assumptions, and hence potential to capture different aspects of population genetic structure.

2.3.1 | Procrustes analyses

Tests of the similarity among taxa in the structuring of genetic variation, including the fit to, and also the departures from, expectations of IBD were conducted using Procrustes analyses (Wang, Zöllner, & Rosenberg, 2012). Specifically, an association between genetic variation and geography while retaining latitudinal and longitudinal information for sampled individuals was quantified using the *procrustes* and *protest* functions in VEGAN package (Oksanen et al., 2017). The strength of the association was compared across taxa by the similarity statistic t_0 , which ranges from 0 to 1, and provides a basis for investigating the history of migration.

The statistical technique involves a transformation of a genetic map onto a geographic map to maximize their similarity (i.e. minimize the sum of squared Euclidean distances between the two maps), where a principal component analysis (PCA) of the genetic data was used to generate individual-level coordinates of the first two components (PC1 and PC2). The position of individuals in the genetic map can be visualized, highlighting whether individuals depart from expected levels of gene flow (i.e. they are genetically too similar, or less similar to other individuals sampled across the landscape than expected given their geographic position). To explore how aspects of the migration history (i.e. departures from expected patterns of gene flow under IBD) reflect the wetlands, the correspondence between the genetic map and the (a) environmental space and (b) stability of a biome was examined. To test for an association between biome-specific environmental space and genetic variation, the residuals of the Procrustes analysis between genes and geography for each species were paired with a matrix of extracted PCA data of environmental variables (described below) for the sequenced individuals (e.g. Knowles et al., 2016). For examining how the stability/instability of habitat within biomes might impact the history of migration, we compared the general correspondence between the position of individuals in genetic space and areas of projected stability; stability is inferred by comparing ecological niche models for different periods of the past with the present, and is described below. In particular, deviations from expected patterns of gene flow (i.e. departures from IBD) were examined in relation to the mapped areas of historical habitat instability to assess whether there was any qualitative correspondence of genetic variation with this biome-specific property (see also Knowles et al., 2016).

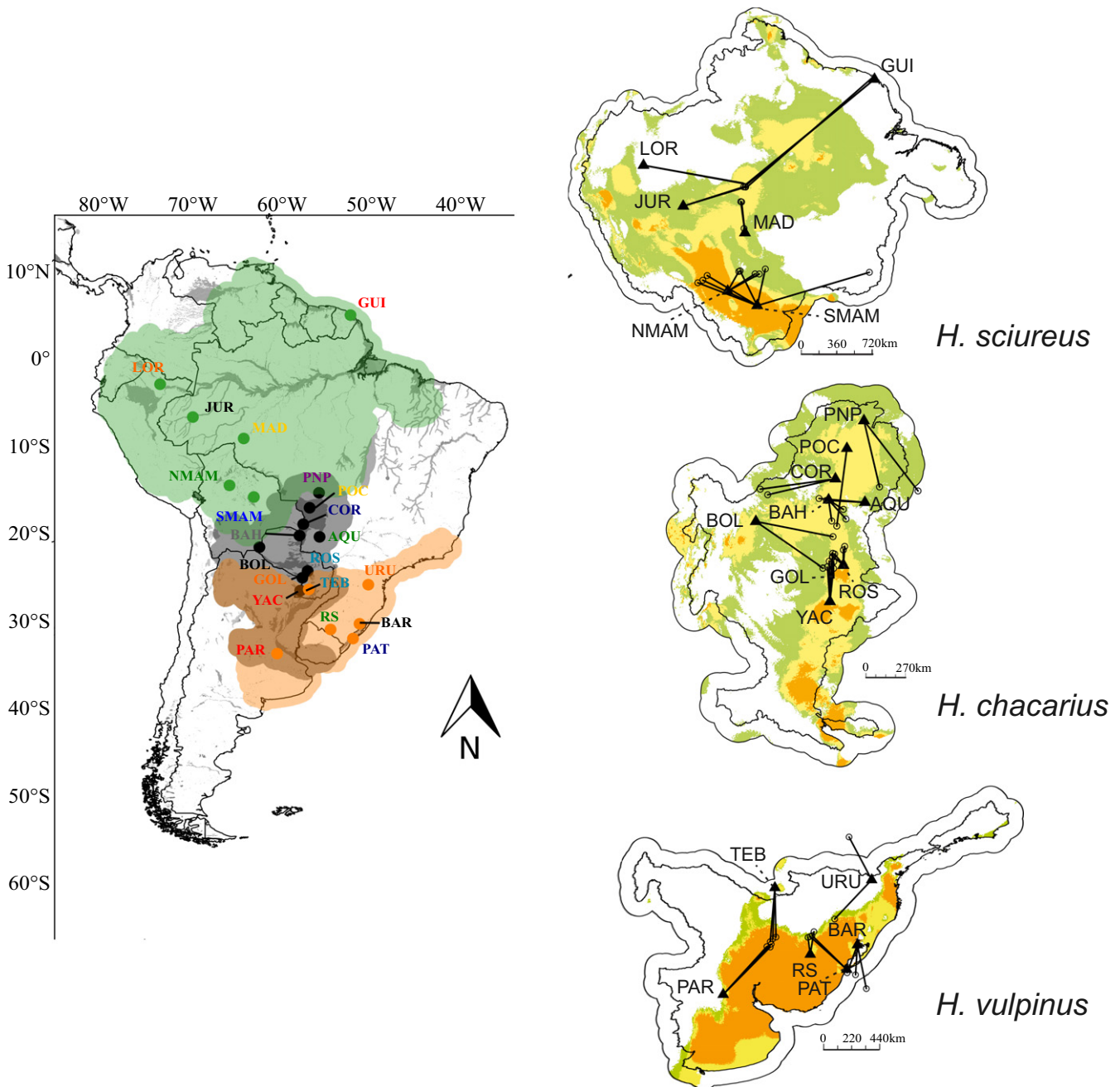


FIGURE 2 Map of the distributions with sampled populations marked for each of the three marsh rats species, with each species colour coded (*Holochilus chacarius*, *Holochilus sciureus* and *Holochilus vulpinus* are shown in black, green and orange respectively); sampling locations span each species' range, which are largely non-overlapping with each taxa occupying different biomes. On the right plots of Procrustes-transformed PCA's of genomic variation of each individual mapped in the genomic PC-space (marked by circles; note the genetic position of some individuals are largely overlapping in some cases) relative to the geographic location of sampled populations (marked by triangles) showing the deviation in the genomic PC-space from the expected pattern of genetic variation based on geography (i.e. the length of the line connecting individuals to their geographic location represents the magnitude of the deviation), as well as how the association between genes and geography differed among species, with $t_0 = 0.69$, 0.64 and 0.52 for *H. vulpinus*, *H. chacarius* and *H. sciureus* respectively. Additionally, the plots for each taxon are shown on the projected stability of each region, where stability is defined as areas that remained suitable overtime (see Figure S3.1 in Appendix 3 for distributional maps from ENMs for each geologic period). Stable areas since LGM, 21 ka, are marked in orange, yellow marks areas that have been stable since Holocene, 6 ka, relative to the unstable areas marked in green (i.e. projected suitable areas for the present, but not the past) [Colour figure can be viewed at wileyonlinelibrary.com]

The significance of the similarity statistic t_0 was evaluated for each species based on 10,000 permutations, where geographic locations were randomly permuted across the different sample localities.

In each species, the sensitivity of the similarity statistic t_0 to particular populations was also evaluated by repeating the analyses excluding one population at a time, with replacement (which is represented

by the similarity statistic t , following Wang et al., 2012). We also computed a similarity score (t ; following Wang et al., 2012) between PCA coordinates for the complete dataset and those for datasets in which one population was excluded to assess whether any populations had a disproportionate effect on the relationship between genes and geography (Knowles et al., 2016; Wang et al., 2012).

2.3.2 | Covariance structure of allele frequencies

Spatial patterns of genomic variation were also examined based on an allele frequency covariance matrix using the program *SPACEMIX* (Bradburd, Ralph, & Coop, 2016); comparison with the results from the PCA provides a measure of whether the PCAs, and hence Procrustes analyses, may be biased by sampling effects (see Novembre & Stephens, 2008). Following the developer's recommendations, 10 "fast" independent chains were run for 5×10^6 MCMC iterations, without conditioning populations on their locations and with no admixture. This was followed by a "long" run of 10^8 iterations, with parameters sampled every 10^2 iterations, in which population locations were initiated at the origin (i.e. inferred from the "fast" runs), and all other parameters (α_0 , α_1 , α_2 , η and w) were drawn randomly from their priors at the start of each chain.

2.3.3 | Mantel tests and dbrDA

As a complementary test to the analyses described above, the correlation between pairwise F_{ST} -values and Euclidean geographic distances among populations, as well as associations between genetic distance and environmental resistance, was examined using a Mantel, partial Mantel tests and dbrDA (Legendre & Anderson, 1999) from the R package *VEGAN* (Oksanen et al., 2017). For the Mantel tests, a sequential population dropout procedure, in which the test was repeated excluding one population at a time, was also conducted to confirm that the results were robust. For the partial Mantel test, an environmental resistance matrix among populations was generated based on raster maps obtained with the environmental niche models (ENMs) using *CIRCUITSCAPE* 4.0 (Shah & McRae, 2008). The *capscale* R function was used in the dbrDA analysis to test for the relationship between pairwise genetic distances and corresponding climatic variables (represented by the PC1 of the 19 climate layers used in the ENMs), and removing the effect of geographic distance separating populations (He et al., 2013). In addition, a second partial Mantel test and dbrDA analyses were performed with the environmental variables extracted from the PCA data for the locations of sequenced populations (He et al., 2013) to examine whether the environment might make a significant contribution to patterns of genetic variation, after controlling for the effects of geography.

2.4 | Characterizing historical stability and environmental space across distributional areas

To characterize historical stability and compare the environmental space of the distributional areas of each *Holochilus* taxa, ENMs

were generated from the 19 bioclimatic variables for the present (Bioclim), Holocene (6 ka; MPI-ESM-P; Bioclim) and the LGM (21 ka; MPI-ESM-P; Bioclim; Hijmans, Cameron, Parra, Jones, & Jarvis, 2005) using *MAXENT* 3.3.3k (Phillips, Anderson, & Schapire, 2006). Georeferenced occurrence data representative of the ranges of each species was used; vetted data were obtained by direct examination of specimens (Appendix S2) or from taxon-specific bibliographic sources (D'Elía, Hanson, Mauldin, Teta, & Pardiñas, 2015; Hershkovitz, 1955; Pardiñas & Teta, 2011; Pardiñas, Teta, Voglino, & Fernández, 2013).

A PCA was used to identify a subset of environmental variables with <0.7 of correlation based on analyses from the *prcomp* function in R; among correlated variables, the variable with the highest score was retained. Occurrence data were rarefied using *SDMToolBox* at a resolution of 10 km to reduce spatial autocorrelation. To avoid overfitting, and considering the semiaquatic habit of *Holochilus*, the geographic extent of the environmental layers used as a mask corresponded to a 100-km buffer surrounding the respective river basins where the species are distributed. To achieve a balance between goodness-of-fit and model complexity, we used the *ENMevaluate* function from 'ENMeval' package (Muscarella et al., 2014), and we tested models over combinations of regularization parameters from 0.5 to 3 in intervals of 0.5 and combinations of feature parameters (Auto, Linear, Quadratic, Hinge, Linear+Quadratic and Linear+Quadratic+Hinge, according to *MAXENT* recommendations). Regularization and feature parameters were chosen using Akaike information criterion (AIC; Warren & Seifert, 2011) and area under the receiver-operator curve (AUC; Swets, 1988). Each model parameter class was replicated 10 times for cross-validation. For each model, we extracted binary predictions, where suitable habitat presence was inferred according to the significance threshold for each model from *MAXENT*.

Climatically stable areas were inferred from the intersection of the binary predictions under current and past climate scenarios (see Table S3.1, Figure S3.1, Appendix S3 in Supporting Information for additional details). Variation in current environmental conditions across each biome was quantified with a PCA of the 19 Bioclim variables using the *rasterPCA* function from the *DISMO* package in R (Hijmans, Phillips, Leathwick, & Elith, 2016), with input variables rescaled from 0 to 1 (so that the PCs are not sensitive to differences in the units). For a graphical presentation of this variation across the landscape, the *ggRGB* function from 'RStoolbox' package in R (Leutner & Horning, 2016) was used to construct maps, with the red layer corresponding to PC1, green layer corresponding to PC2 and the blue layer corresponding to PC3. To visualize how the position of sampling localities spans the environmental space, the environmental dispersion of sequenced individuals relative to the total environmental space for each species was mapped using the environmental data (PC1 and PC2 of the PCA performed with the bioclimatic variables) and the occurrence points of sequenced individuals (see Lanier, Massatti, He, Olson, & Knowles, 2015).

3 | RESULTS

3.1 | Population genetic summary statistics

Values of population genetic summary statistics were broadly overlapping among taxa (Table S4.1, Figure S4.1, Appendix S4 in Supporting Information), except for a statistically significant difference in H_{obs} , with higher values in *H. vulpinus* compared with the other species. Genetic diversity, as measured by π , was similar among taxa, varying from 0.1 to 0.2. Inbreeding coefficients, F_{is} , were also similar across taxa with the exception of two populations; specifically, S_MAM in *H. sciureus* and YAC in *H. chacarius* had substantially higher inbreeding coefficients, although neither exhibited reduced genetic diversity.

3.2 | Historical stability and environmental space across distributional areas

The current environmental characteristics of the biomes inhabited by the three species clearly differ (Figure 3). The environmental space of the Atlantic Forest and Pampas area inhabited by *H. vulpinus* differs the most from the other regions. There is some resemblance between the environmental space occupied by *H. vulpinus* and the north-western region of *H. sciureus*. However, these two areas are separated by a large region with different environmental characteristics (see the two bluish areas are separated by a large green area; Figure 3). Comparison of the environmental conditions where populations were sampled to the rest of biome shows that the sites where genetic samples were collected are generally representative of the environmental range encompassed by the species' distributions, spanning the entire dispersion of environmental values of the PCA (Figure 4).

Estimates of habitat stability based on similarity in spatial distribution of suitable habitat from the LGM, the Holocene and the present, show differences in the stability of the habitat across biomes, as well as within each biome. Specifically, the habitat across most of the distributional range of *H. vulpinus* is projected to be stable, whereas only the southwestern Amazon is predicted to be stable for *H. sciureus*, and only small and patchy areas were inferred to be stable for *H. chacarius* (Figure 2).

3.3 | Structuring of genetic variation

Regarding the geographic distribution of genetic variation within each species, several patterns are reinforced across methods. For example, when the position of individuals along PC1 and PC2 from a genetic PCA is projected onto the geographic distribution of sampled individuals (Figure 5), it highlights how similarity in the multidimensional genetic space is related to where an individual occurs geographically, confirming the Procrustes results (Figures 2 and 6). Likewise, this similarity is also evident when comparing the results from the spatial covariance of alleles analyses (SPACEMIX results) with those from a PCA of genetic variation (Figure S4.2 in Appendix S4). This general correspondence suggests that the PCAs (and therefore the Procrustes analyses) are not subject to biases that can be

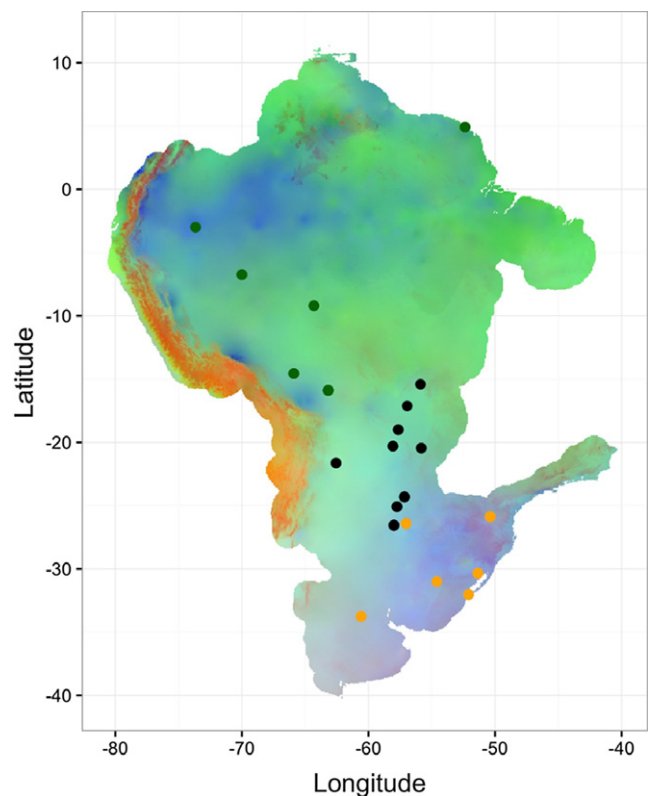


FIGURE 3 Map of the environmental variation across the region where the three species are distributed (orange dots represent *Holochilus vulpinus*, black dots *H. chacarius* and green dots *H. sciureus*), where differences in colour depict geographic regions that differ the most from each other. Specifically, PC1, PC2 and PC3 of bioclimatic variables across the landscape were rescaled between 0 and 1, and the RGB colour composite was calculated and plotted in the map with PC1 set as the red scale, PC2 as the green scale, and PC3 as the blue scale. Coloured dots correspond to the populations presented in Figure 2. Note that the border used to characterize the relative difference in environment corresponded to the same area used to generate the ENMs in each species (see Figure 2) and does not include the Andes [Colour figure can be viewed at wileyonlinelibrary.com]

introduced from the sampling distribution of individuals (Novembre & Stephens, 2008). However, the position of some individuals of *H. chacarius* in multivariate genetic space differs somewhat from the spatial covariance of alleles. Specifically, we note that the GOL population occupies a more distinctive position relative to other populations in the geogenetic map of SPACEMIX compared with its overlap with other populations in the PCA (Figure S4.2 in Appendix S4). However, results from the sequential population dropout procedure in the Procrustes analysis showed that exclusion of the GOL population neither produced large changes in the strength of the association between genes and geography, nor resulted in a significant distortion of the genetic PC map (Figures 5 and 6). We also note that the sampling gap in *H. sciureus*, in which central Amazonian samples were not available (Figure 2), does not appear to bias the position of these individuals in PC space (Bradburd et al., 2016) given the similarity between the two approaches (Figure S4.2 in Appendix S4). Likewise, it is also noteworthy that inclusion of the

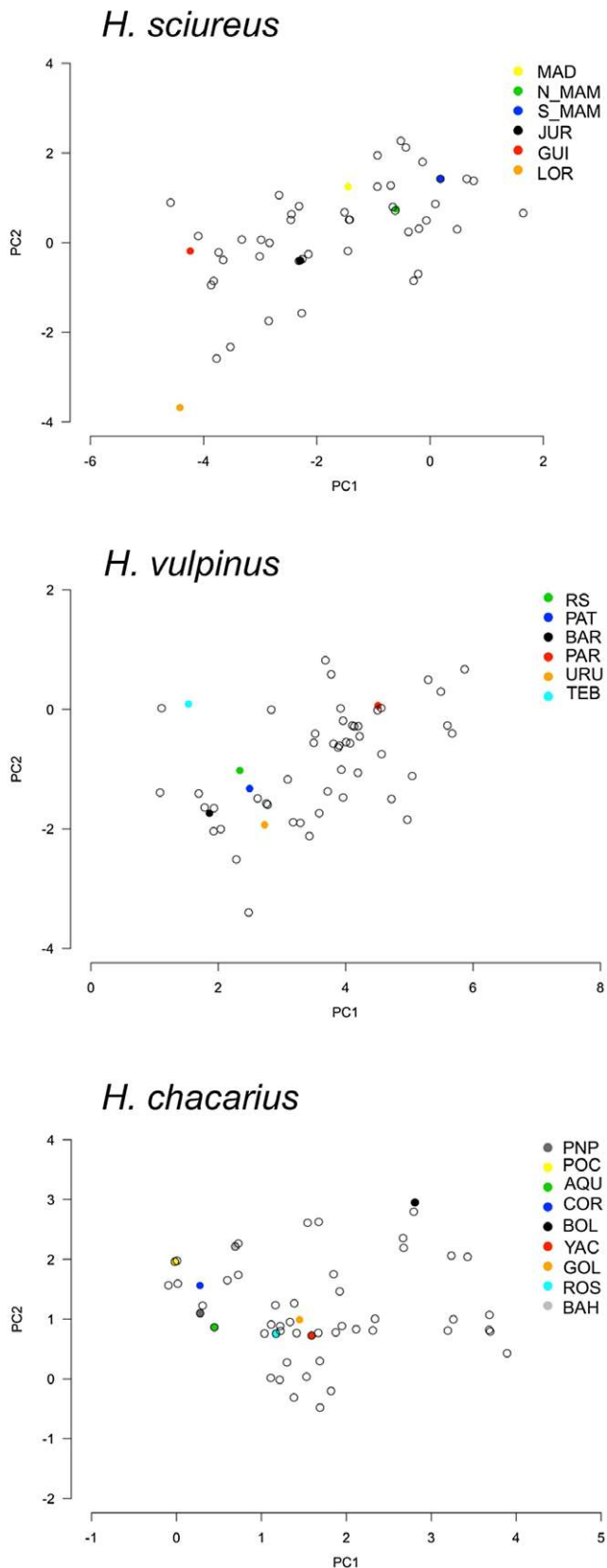


FIGURE 4 Dispersion in environmental space of the sampled populations used in our genetic analyses (marked as coloured dots) relative to the PC values for *Holochilus* sampling locations used in the ENMs. For *H. sciureus*, PC1 is strongly positively correlated with minimum temperature of coldest month (Bio6) and explains most of the variation among populations (i.e. 55.49%), whereas PC2 explains relatively little variation among populations (19.66%). For *H. vulpinus*, PC1 is strongly positively correlated with the annual precipitation (Bio12) and explains most of the variation among populations (48.56%), whereas PC2 explains 24.26% of the variation among populations. For *H. chacarius*, PC1 is strongly positively correlated with the minimum temperature of coldest month (Bio6) and explains most of the variation among populations (54.4%), whereas PC2 explains 21.8% of variation among populations. Names of populations as in Figure 2 [Colour figure can be viewed at wileyonlinelibrary.com]

excluded; within a species, the shift from the t_0 -value (t_0-t') was as high as 0.15 with the exclusion of specific populations (Figure 6, Table S4.2 in Appendix S4). The differences in the strength of the association between genes and geography does not appear to be attributable to the undue influence of any single population on the PCA of genetic variation in any of the species (i.e. t' values were >0.91 in all but four cases; see Table S4.2 in Appendix S4). However, we note that in both *H. vulpinus* and *H. chacarius*, but not *H. sciureus*, the same populations that resulted in higher association between genes and geography when excluded had a disproportionate influence on the PCA of genetic variation (the URU population in *H. vulpinus*; the YAC and BOL populations in *H. chacarius*; in contrast to the GUI population in *H. sciureus*).

Results of Mantel, partial Mantel tests and dbRDA based on geography and environment were not significant for any taxa (Tables S4.4 and S4.5 in Appendix S4), including the Mantel tests with sequential removal of individual populations. The only exception was a significant association between environment and genetic variation based on dbRDA test only in *H. chacarius*; however, the association was not significant after controlling for geography.

The lack of significance based on Mantel tests and dbRDAs contrasts with a significant association between geographic and genetic variation in each species detected by the Procrustes analysis, which unlike the former tests retains the geographic information about latitudinal and longitudinal position of populations relative to patterns of genetic differentiation. The strength of association between genes and geography (t_0) was quite similar across all species (when the outlier population GUI in *H. sciureus* was excluded), ranging from 0.64 to 0.67, out of a maximum of 1, meaning there is also a substantial amount of variation not explained by the geographic position of populations in each species.

3.4 | Genetic deviations/displacements in geographic and environmental space

In *H. vulpinus*, when we consider the deviations of individuals in genetic space from expectations based on their geographic location (i.e. the length of the lines connecting a sampled population to a sequenced individual; Figure 2), the deviations primarily vary along a

geographically disjunct GUI population in *H. sciureus* also did not strongly affect the influence the PCA of genetic variation genetic.

The strength of the association between genes and geography varied substantially when individual populations were sequentially

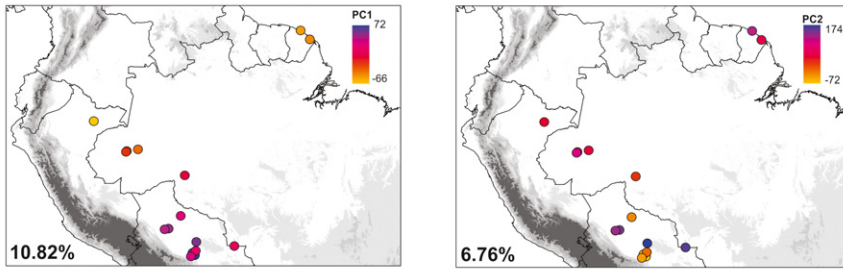
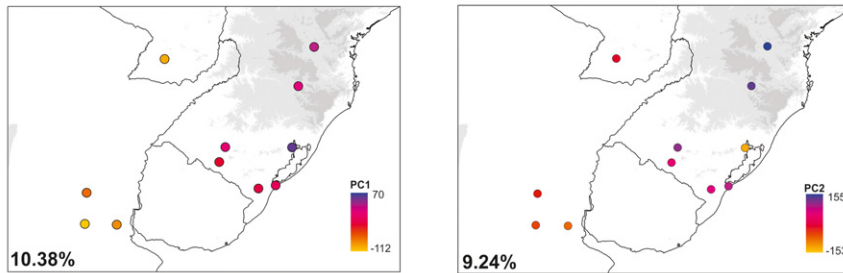
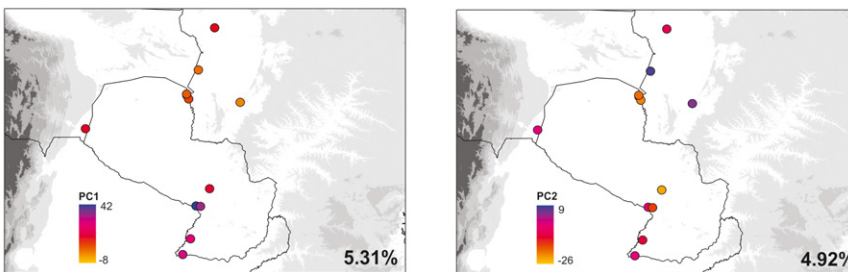
(a) *H. sciureus*(b) *H. vulpinus*(c) *H. chacarius*

FIGURE 5 Distribution of individuals of *Holochilus* along PC1 and PC2 of genomic variation colour coded, and the percentage of variance explained by each PC are shown on a map (i.e. different colours correspond to individuals with the greatest genomic difference along PCs) for each species. Elevation differences are shown in grey scale on each map as well [Colour figure can be viewed at wileyonlinelibrary.com]

latitudinal axis. Moreover, the individuals tend to occupy the central area of the species distribution and the position of these individuals in the PC genetic space show a strong correspondence with areas of stability (Figure 2), with the exception of one population in the southeast (BAR; Figure 2) whose individuals deviate in a southerly direction.

In contrast, the directions of departures in genetic space relative to where individuals of *H. sciureus* were sampled geographically tend to follow a longitudinal axis (Figure 2). Nevertheless, this species also shows some correspondence between the position of individuals in genetic space and areas of projected stability like *H. vulpinus*. Likewise, the deviations of individuals sampled in what is projected to be stable areas historically (N_MAM and S_MAM) tend to show relatively small departures from geographic expectations.

Lastly, in *H. chacarius* the displacement of many individuals to the central part of the species range was observed even though the sampled populations correspond to areas of projected stability (Figure 2). Only some of the northern populations (PNP and COR, but not POC) are somewhat distinct from this general cluster.

Tests of a correlation between the residuals from the Procrustes analysis of genes and geography with the environmental distance among populations suggests that environmental differences contribute to some of the genetic differences observed among

individuals in each species (Table S4.3 in Appendix S4). As with the tests of IBD that did not retain the relative position of sampled populations in geographic space, neither the partial Mantel nor dbrDA detected significant contribution of environmental differences to genetic distances (Tables S4.4 and S4.5 in Appendix S4).

4 | DISCUSSION

Similarity in the structuring of population genetic variation (Figure 2) and population genetic diversities (see Table S4.1, Figure S4.1 in Appendix S4) highlight the contribution of common natural history traits to the similar migration histories of the taxa. However, biome-specific effects on genetic structure explain a substantial amount of genetic variation not explained solely by gene flow associated with shared traits among the taxa. Specifically, quantitative and qualitative analyses (Figures 2 and 3) show that variation of the environment and historical stability within biomes contribute to patterns of genetic structure.

Below we discuss the insights these findings offer about the South American wetlands and their constituent taxa as one of the first, and to our knowledge, only study that brings the resolution of data from

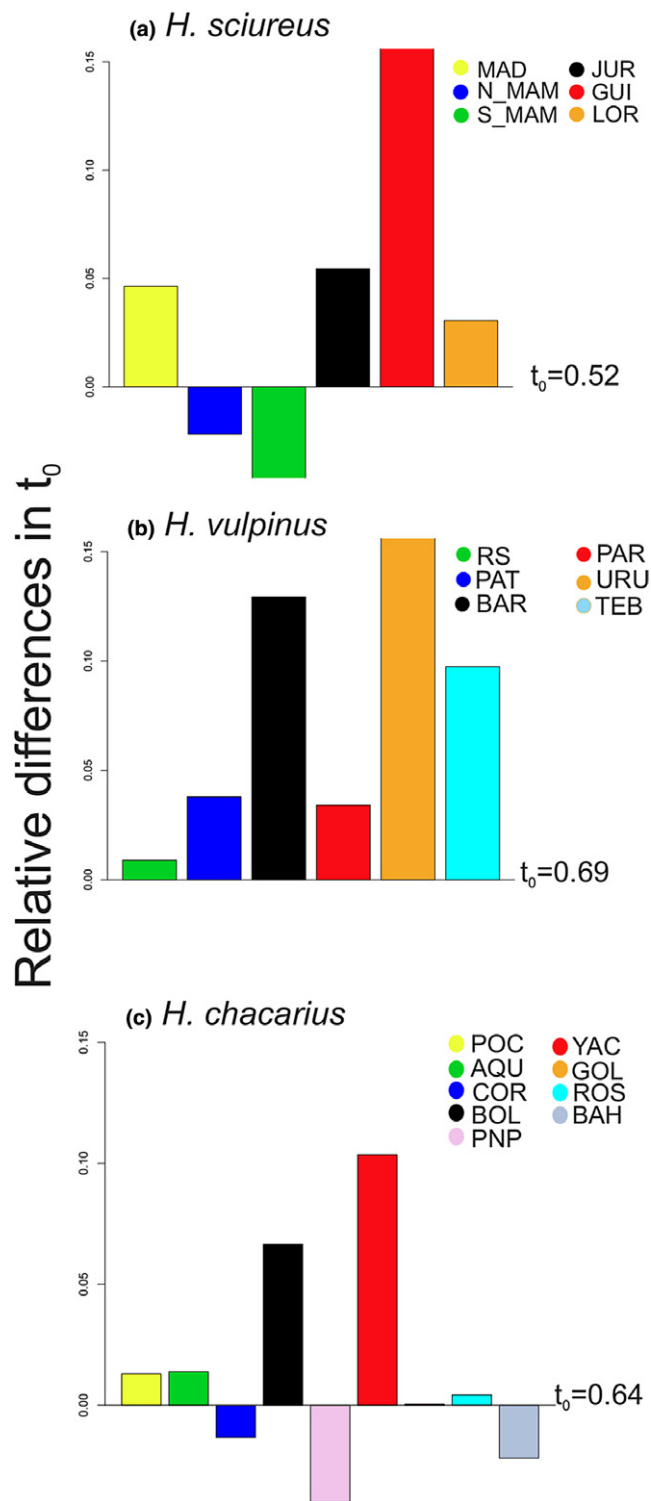


FIGURE 6 Comparison of the changes in the strength of the association between genes and geography with the exclusion of individual populations (i.e. t) relative to when all populations are analysed (i.e. t_0). Values for each species are standardized by t_0 (i.e. 0 on y-axis corresponds to t_0) such that positive values indicate a stronger association between genes and geography when a population is excluded, whereas negative values indicate a weaker association. Bar colours represent sampling populations following the coloured names of populations in Figure 2 [Colour figure can be viewed at wileyonlinelibrary.com]

next-generation sequence technologies to these vast, but relatively understudied, landscapes. In addition, we highlight the niche our work fills in comparative phylogeography: the study of ecologically similar taxa from different regions, as opposed to a tradition of comparing taxa (often with differing ecologies) from a single region, to make inferences about the processes underlying observed patterns of genetic variation (Avice, 2000; Hickerson et al., 2010; Knowles, 2009).

4.1 | Insights from genomic analyses about South American biomes

Despite differences in the distribution of wetland habitats across the biomes inhabited by the taxa (e.g. patches of wetlands vs. large contiguous wetlands), the migration history of the species is similar. Specifically, the strength of the association between genetic variation and geography was similar across the biomes (when the outlier population GUI in *H. sciureus* was excluded). Moreover, even when considering all populations, the Procrustes analyses indicated the genetic isolation between geographically distant populations in the species inhabiting the two biomes of forest-based patches of wetlands was not more similar to each other relative to the species in the contiguous wetlands. Instead, the strength of the association between genes and geography was most similar in *H. chacarius*, which inhabits the large contiguous wetlands, and *H. vulpinus*, which inhabits patches of wetlands in the Atlantic Forest and in the open vegetation of the Pampas. Together, these results emphasize that gene flow decreases with geographic distance similarly across taxa, irrespective of differences across biomes. This similar migration history suggests that shared characters of the marsh rats, that is biotic factors, play a key role in structuring genetic variation when genetic variation is measured at the scale of the species entire distribution. It may be that the effect of the current configuration of habitats (contiguous vs. patchy) may be discernible at different spatial scales, such as at a local landscape level. Such scale-dependencies of biotic traits in migration history is something we would like to investigate in the future; however, we cannot address this question with the available sampling.

Notwithstanding the similar migration histories suggested by the similar strength of the genes and geography association across taxa, the association between biome-specific properties with deviations from expectations under IBD point to potential hypotheses about how the biomes may impact genetic structure. In particular, the magnitude of deviations from a pattern of IBD differs across the landscape in each taxon, suggesting localized differences in population connectivity and/or persistence. The most dramatic effect is across Amazonia, where the more northern population of *H. sciureus* tend to be genetically quite similar despite fairly large distances separating the sampled individuals (Figure 2). This geographic area that individuals map based on their genetic makeup is consistent with an estimated area of habitat stability based on ENMs for the past and present (Figure 2), which suggests a potential role of climatic shifts, and coincidentally is estimated to be less stable relative to the southern sampled populations (Arruda, Schaefer, Fonseca, Solar, & Fernandes-Filho, 2017; Bush et al., 2007; Mayle, Burbridge, & Killeen, 2000; Urrego, Silman, & Bush,

2005). In contrast, in southern regions of the distribution of *H. sciureus*, there is a strong correspondence between the genetic and geographic position of individuals (Figure 2), and their general genetic distinctiveness relative to the more northern populations suggests a history of relative regional isolation within the biome (e.g. Knowles et al., 2016). Differences in fit of northern and southern regions to expected patterns of gene flow under an IBD model, in concert with corresponding differences in duration of stability inferred from the ENMs (Figure 2), suggest variation in the historical stability within a biome may be contributing to genetic structure of constituent taxa. Specifically, our genetic analyses suggest a dynamic of recent expansion may characterize the wetlands from the northern part of the Amazonian biome, in contrast to relative population persistence within the southern region. This pattern agrees with some, but not all, aspects of inferences about past regional stability from other studies. For example, the area corresponding to our LOR, JUR and MAD populations (Figure 2) corresponds to an area that is hypothesized to have been associated with a dynamic geological history (Aleixo & Rossetti, 2007; Leite & Rogers, 2013), and stable past hydroclimate variation (see Cheng et al., 2013). Although the sampling in the east is limited, the genetic clustering of the GUI population with other western Amazonian sampled populations suggests however that the wetlands in the eastern area (or at least the north-east; Figure 2) is also relatively unstable, unlike the results from some landscape genetic and geologic studies (Aleixo & Rossetti, 2007; Leite & Rogers, 2013).

Although less dramatic, patterns of genetic structure in *H. chacarius* show a somewhat similar pattern with populations from the northern part of their range exhibiting the greatest deviations under IBD (individuals from the north tend to be genetically more similar to individuals from the southern than expected), with most of the estimated regions of stability located in the south (Figure 2). This complex history suggests the expansion of wetland across this biome may be fairly recent (see Bezerra & Mozeto, 2008), with more sparse vegetation and intermittent torrential flows along the alluvial fans in the past (Assine et al., 2015; Whitney et al., 2011). In contrast, while *H. vulpinus* shows the larger continuous areas of stability among all species (see Arruda et al., 2017 for proposed expansion scenarios), it nonetheless shows a correspondence between genes and geography that is similar to the other two species (i.e. the t_0 of *H. vulpinus* is 0.69, while the t_0 of *H. sciureus* and *H. chacarius* is 0.52 and 0.64 respectively). Such similarity again emphasizes that differences in stability by itself, or continuity of wetlands (as opposed to being patchy), is not sufficient to override patterns of genetic variation within each species that are consistent with migration history where gene flow decreases with distance (Figure 2).

In addition to the potential effect of historical stability, multiple lines of evidence point to other characteristics of the biomes that may affect patterns of genetic variation, albeit at a local level (as opposed to the regional scale of an entire biome, given similar degrees in the strength in the association between genes and geography). Environmental differences across the landscape show a significant effect on the position of individuals in genetic space, after controlling for the effect of geography. However, it is noteworthy that this effect, as with

test of a general correspondence between genes and geography, is only detectable when retaining the relative distance of populations latitudinally and longitudinally (i.e. with Procrustes analyses). The lack of significance when reducing the geographic separation of populations to a one-dimensional axis (Mantel and dbRDA analyses; Table S4.4 and S4.5 in Appendix S4) highlights how connectivity patterns do not mirror expectations based on random diffusion across a landscape (Excoffier, Foll, & Petit, 2009). That is, a longitudinal distance versus a latitudinal distance is not equivalent in terms of the impact on population differences. As such, our work highlights that within each of the biomes, connectivity patterns vary locally in each species.

Patterns of environmental heterogeneity across the landscape (Figure 3) and/or shifts in habitat stability over time (Figure 2) can cause populations separated by similar geographic distances to differ in levels of connectivity (He et al., 2013; McRae, 2006). Although our tests are insightful in that they identify patterns suggestive of environmental factors influencing connectivity, as correlative analyses they do not provide tests of the process itself (Knowles et al., 2016). Determining which of the dynamics might produce genetic variation consistent with what is observed (i.e. applying a model selection framework to distinguish between the geographic distribution of wetland habitats versus shifting climates; see He et al., 2013; Knowles & Massatti, 2017), is beyond the scope of this study without additional sampling for conducting spatially explicit model-based analyses. However, as with the guiding theme of this work, tests to identify concordant processes across species will be especially exciting to understand the extent to which genetic variation in the taxa might arise from common processes, despite differences in the biomes themselves.

4.2 | Similar taxa, but different biomes, in comparative phylogeography

Although the study of multiple taxa across different regions predominates in historical biogeography and macroecology (Arregoitia, Fisher, & Schweizer, 2017; Leite et al., 2014), comparative phylogeographic analyses have focused primarily on co-distributed taxa from a single area (Avice, 1992; Hickerson et al., 2010; Knowles, 2009). There are several potential explanations for this tradition and one is related to sampling efforts for phylogeographic inference: population sampling poses logistical constraints such that a more circumscribed geographic region is simply more feasible. The focus on more circumscribed region also reflects the traditional motivation behind the comparative phylogeography—making inferences about the history of a region. It is through tests of concordance across taxa from a particular region that specific biogeographic barriers (Avice, 1992) or areas of long-term stability (Carnaval, Hickerson, & Haddad, 2009) might be inferred. In fact, such tests of concordance across taxa with differing ecologies have been used to identify how abiotic factors might supersede any ecological differences among taxa (Naka, Bechtoldt, Henriques, & Brumfield, 2012) or reject the role of historical barriers in driving divergence patterns (Smith et al., 2014).

As a complement to the study of co-distributed taxa, there is a precedent for studying taxa from different regions in comparative

phylogeography. For example, analysis of evolutionarily independent regions can be considered as natural replicates and can be used for making generalizations about geologic events that have broadly affected landscape histories (Bermingham & Moritz, 1998), while controlling for discord that may arise from differences in species-specific traits to mitigate poor predictive power when concordance across regions is not supported (see Papadopoulou & Knowles, 2016). Here we adopted a strategy of similar species, but different regions as an inferential framework. Moreover, with a complementary set of tests and the genetic resolution provided by thousands of loci, we address how genetic patterns vary (i.e. we move beyond concordant vs. discordant binary). As such, we were able to identify concordance in genetic variation (similar strength in the association between genes and geography, despite differences in the wetlands from the different biomes) that suggests a highly specialized semiaquatic life may result in similar genetic IBD. However, inspection of local deviations from IBD identified additional biome-specific effects related to environmental differences and historical stability. That is, rather than biotic versus abiotic factors predominating on the genetic structure, our framework identifies how both components can influence genetic variation, as well as their respective geographic scales of influence. Regarding this latter point, the biotic factors predominate at the regional scale (at the level of genetic structure across the entire biome), whereas abiotic factors appear to contribute to local departures from IBD (see Figure 2). These scale-specific effects are especially interesting considering the qualitative correspondence between regions of instability and departures from IBD, which may be suggestive of particular populations that might have been vulnerable to environmental change (Choueri et al., 2017; Harvey, Aleixo, Ribas, & Brumfield, 2017; Rocha et al., 2014).

With very few genetic studies on organisms that inhabits the flood-dominated landscapes in South America, the present work adds to our knowledge, but there is admittedly much that remains unknown. For example, the effect on terrestrial organisms (as opposed to aquatic or semiaquatic species) may not show the same relative role of biotic and abiotic factors or geographic scale at which they might predominate, given that flooding and rivers may act as barriers in terrestrial organisms rather than as routes for connectivity (Lima, Lima-Ribeiro, Tinoco, Terribile, & Collevatti, 2014; Rocha et al., 2014). This suggests the importance of a nuanced view of interpreting concordance (or the lack thereof) across taxa. Specifically, taxa may show opposing patterns of connectivity and yet this discord could arise from deterministic processes (i.e. not the idiosyncrasies of history; see Massatti & Knowles, 2014) because what constitutes routes of connectivity in a given wetland biome depends upon species-specific traits (e.g. for terrestrial vs. aquatic taxa).

ACKNOWLEDGEMENTS

This work was funded by Fundação de Amparo à Pesquisa do Estado de São Paulo (FAPESP; Proc: 2014/22444-0, 2012/24099-3, 2009/16009-1, 2016/20055-2). We are also grateful to the curators and researchers that generously granted us tissues samples. We also

thank Luciana Resende, Anna Papadopoulou and Melisa Olave for all the help with bioinformatics processing and giving helpful feedback on earlier drafts of the manuscript.

ORCID

Joyce R. Prado  <http://orcid.org/0000-0002-2025-5479>

Alexandre R. Percequillo  <http://orcid.org/0000-0002-7892-8912>

Andréa T. Thomaz  <https://orcid.org/0000-0002-9755-2674>

L. Lacey Knowles  <https://orcid.org/0000-0002-6567-4853>

REFERENCES

- Aleixo, A. (2006). Historical diversification of floodplain forest specialist species in the Amazon: A case study with two species of the avian genus *Xiphorhynchus* (Aves: Dendrocolaptidae). *Biological Journal of the Linnean Society*, 89, 383–395. <https://doi.org/10.1111/j.1095-8312.2006.00703.x>
- Aleixo, A., & Rossetti, D. F. (2007). Avian gene trees, landscape evolution, and geology: Towards a modern synthesis of Amazonian historical biogeography? *Journal of Ornithology*, 148, 443–453. <https://doi.org/10.1007/s10336-007-0168-7>
- Arregoitia, L. D., Fisher, D. O., & Schweizer, M. (2017). Morphology captures diet and locomotor types in rodents. *Royal Society Open Science*, 4(1), 160957. <https://doi.org/10.1098/rsos.160957>
- Arruda, D. M., Schaefer, C. E. G. R., Fonseca, R. S., Solar, R. R. C., & Fernandes-Filho, E. I. (2017). Vegetation cover of Brazil in the last 21 ka: New insights into the Amazonian refugia and Pleistocene arc hypotheses. *Global Ecology and Biogeography*, 27, 47–56. <https://doi.org/10.1111/geb.12646>
- Assine, M. L., Macedo, H. A., Stevaux, J. C., Bergier, I., Padovani, C. R., & Silva, A. (2015). Avulsive rivers in the hydrology of the Pantanal wetland. In I. Bergier & M. L. Assine (Eds.), *Dynamics of the Pantanal wetland in South America* (pp. 83–110). Cham, Switzerland: Springer. <https://doi.org/10.1007/978-3-319-18735-8>
- Avise, J. C. (1992). Molecular population structure and the biogeographic history of a regional fauna: A case history with lessons for conservation biology. *Oikos*, 63, 62–76. <https://doi.org/10.2307/3545516>
- Avise, J. C. (2000). *Phylogeography: The history and formation of species*. Cambridge, MA: Harvard University Press.
- Bermingham, E., & Moritz, C. (1998). Comparative phylogeography: Concepts and applications. *Molecular Ecology*, 7, 367–369. <https://doi.org/10.1046/j.1365-294x.1998.00424.x>
- Bezerra, M. A. D. O., & Mozeto, A. A. (2008). Deposição de Carbono Orgânico na Planície de Inundação do Rio Paraguai durante o Holoceno médio. *Oecologia Brasiliensis*, 12, 155–171.
- Bradbud, G. S., Ralph, P. L., & Coop, G. M. (2016). A spatial framework for understanding population structure and admixture. *PLoS Genetics*, 12, 1–38. <https://doi.org/10.1371/journal.pgen.1005703>
- Bush, M. B., Silman, M. R., & Listopad, C. M. C. S. (2007). A regional study of Holocene climate change and human occupation in Peruvian Amazonia. *Journal of Biogeography*, 34, 1342–1356. <https://doi.org/10.1111/j.1365-2699.2007.01704.x>
- Capurucho, J. M. G., Cornelius, C., Borges, S. H., Cohn-Haft, M., Aleixo, A., Metzger, J. P., & Ribas, C. C. (2013). Combining phylogeography and landscape genetics of *Xenopipo atronitens* (Aves: Pipridae), a white sand campina specialist, to understand Pleistocene landscape evolution in Amazonia. *Biological Journal of the Linnean Society*, 110, 60–76. <https://doi.org/10.1111/bj.12102>
- Carnaval, A. C., Hickerson, M. J., & Haddad, C. F. B. (2009). Stability predicts genetic diversity in the Brazilian Atlantic forest hotspot. *Science*, 323, 785–789. <https://doi.org/10.1126/science.1166955>

- Catchen, J., Hohenlohe, P. A., Bassham, S., Amores, A., & Cresko, W. A. (2013). Stacks: An analysis tool set for population genomics. *Molecular Ecology*, 22, 3124–3140. <https://doi.org/10.1111/mec.12354>
- Cheng, H., Sinha, A., Cruz, F. W., Wang, X., Edwards, R. L., D'Horta, F. M., ... Auler, A. S. (2013). Climate change patterns in Amazonia and biodiversity. *Nature Communications*, 4, 1–6. <https://doi.org/10.1038/ncomms2415>
- Choueri, É. L., Gubili, C., Borges, S. H., Thom, G., Sawakuchi, A. O., Soares, E. A. A., & Ribas, C. C. (2017). Phylogeography and population dynamics of Antbirds (Thamnophilidae) from Amazonian fluvial islands. *Journal of Biogeography*, 44, 2284–2294. <https://doi.org/10.1111/jbi.13042>
- D'Elia, G. D., Hanson, J. D., Mauldin, M. R., Teta, P., & Pardiñas, U. F. J. (2015). Molecular systematics of South American marsh rats of the genus *Holochilus* (Muroidea, Cricetidae, Sigmodontinae). *Journal of Mammalogy*, 96(5), 1081–1094. <https://doi.org/10.1093/jmammal/gyv115>
- Dominguez-Bello, M. G., & Robinson, M. D. (1991). A comparison of digestive adaptations in two Neotropical Cricetid rodents (*Holochilus venezuelae* and *Zygodontomys microtinus*). *Physiological Zoology*, 64(6), 1542–1551. <https://doi.org/10.1086/physzool.64.6.30158229>
- Eisenberg, J. F. (1981). *The mammalian radiations: An analysis of trends in evolution, adaptation, and behavior*. Chicago, IL: The University of Chicago Press.
- Erwin, T. L., & Adis, J. (1982). Amazonian inundation forests: Their role as short-term refuges and generator of species richness and taxon pulses. In G. Prance (Ed.), *Biological diversification in the Tropics* (pp. 358–371). New York, NY: Columbia University Press.
- Excoffier, L., Foll, M., & Petit, R. J. (2009). Genetic consequences of range expansions. *Annual Review of Ecology, Evolution, and Systematics*, 40, 481–501. <https://doi.org/10.1146/annurev.ecolsys.39.110707.173414>
- Ferreira, M., Fernandes, A. M., Aleixo, A., Antonelli, A., Olsson, U., Bates, J. M., ... Ribas, C. C. (2018). Evidence for mtDNA capture in the jacamar *Galbula leucogastra/chalcothorax* species-complex and insights on the evolution of white-sand ecosystems in the Amazon basin. *Molecular Phylogenetics and Evolution*, 129, 149–157. <https://doi.org/10.1016/j.ympev.2018.07.007>
- Gonçalves, P. R., Teta, P., & Bonvicino, C. R. (2015). Genus *Holochilus*. In J. L. Patton, U. F. J. Pardiñas, & G. D'Elia (Eds.), *Mammals of South America* (pp. 325–335). Chicago, IL: University of Chicago Press.
- Harvey, M. G., Aleixo, A., Ribas, C. C., & Brumfield, R. T. (2017). Habitat association predicts genetic diversity and population divergence in Amazonian birds. *The American Naturalist*, 190, 631–648. <https://doi.org/10.1086/693856>
- He, Q., Edwards, D. L., & Knowles, L. L. (2013). Integrative testing of how environments from the past to the present shape genetic structure across landscapes. *Evolution*, 67, 3386–3402. <https://doi.org/10.1111/evo.12159>
- Hershkovitz, P. (1955). South American marsh rats, genus *Holochilus*, with a summary of Sigmodont rodents. *Fieldiana Zoology*, 37, 639–673.
- Hershkovitz, P. (1962). Evolution of Neotropical Cricetine Rodents (Muridae) with special reference to the Phyllotine group. *Fieldiana Zoology*, 46, 1–524.
- Hickerson, M. J., Carstens, B. C., Cavender-Bares, J., Crandall, K. A., Graham, C. H., Johnson, J. B., ... Yoder, A. D. (2010). Phylogeography's past, present, and future: 10 years after Avise, 2000. *Molecular Phylogenetics and Evolution*, 54, 291–301. <https://doi.org/10.1016/j.ympev.2009.09.016>
- Hijmans, R. J., Cameron, S. E., Parra, J. L., Jones, P. G., & Jarvis, A. (2005). Very high resolution interpolated climate surfaces for global land areas. *International Journal of Climatology*, 25, 1965–1978. <https://doi.org/10.1002/joc.1276>
- Hijmans, R. J., Phillips, S., Leathwick, J., & Elith, J. (2016). Dismo: Species distribution modeling (Version 1.0–15). R package. Retrieved from <https://cran.r-project.org/web/packages/dismo/index.html>
- Hoorn, C., Wesselingh, F. P., ter Steege, H., Bermudez, M. A., Mora, A., Sevink, J., ... Antonelli, A. (2010). Amazonia through time: Andean uplift, climate change, landscape evolution, and biodiversity. *Science*, 330, 927–931. <https://doi.org/10.1126/science.1194585>
- Junk, W. J. (2013). Current state of knowledge regarding South America wetlands and their future under global climate change. *Aquatic Sciences*, 75, 113–131. <https://doi.org/10.1007/s00027-012-0253-8>
- Knowles, L. L. (2009). Statistical phylogeography. *Annual Review of Ecology, Evolution, and Systematics*, 40, 593–612. <https://doi.org/10.1146/annurev.ecolsys.38.091206.095702>
- Knowles, L. L., & Massatti, R. (2017). Distributional shifts – not geographic isolation – as a probable driver of montane species divergence. *Ecography*, 40, 1475–1485. <https://doi.org/10.1111/ecog.02893>
- Knowles, L. L., Massatti, R., He, Q., Olson, L. E., & Lanier, H. C. (2016). Quantifying the similarity between genes and geography across Alaska's alpine small mammals. *Journal of Biogeography*, 43, 1464–1476. <https://doi.org/10.1111/jbi.12728>
- Lanier, H. C., Massatti, R., He, Q., Olson, L. E., & Knowles, L. L. (2015). Colonization from divergent ancestors: Glaciation signatures on contemporary patterns of genomic variation in Collared Pikas (*Ochotona collaris*). *Molecular Ecology*, 24, 3688–3705. <https://doi.org/10.1111/mec.13270>
- Ledru, M. P., Montade, V., Blanchard, G., & Hély, C. (2016). Long-term spatial changes in the distribution of the Brazilian Atlantic Forest. *Biotropica*, 48, 159–169. <https://doi.org/10.1111/btp.12266>
- Legendre, P., & Anderson, M. J. (1999). Distance-based redundancy analysis: Testing multispecies responses in multifactorial ecological experiments. *Ecological Monographs*, 69, 1–24. [https://doi.org/10.1890/0012-9615\(1999\)069\[0001:DBRATM\]2.0.CO;2](https://doi.org/10.1890/0012-9615(1999)069[0001:DBRATM]2.0.CO;2)
- Leite, Y. L. R., Costa, L. P., Loss, A. C., Rocha, R. G., Batalha-Filho, H., Bastos, A. C., ... Pardini, R. (2016). Neotropical forest expansion during the last glacial period challenges refuge hypothesis. *Proceedings of the National Academy of Sciences*, 113, 201513062. <https://doi.org/10.1073/pnas.1513062113>
- Leite, R. L., Kolokotronis, S.-O., Almeida, F. C., Werneck, F. P., Rogers, D. S., & Weksler, M. (2014). In the wake of invasion: Tracing the historical biogeography of the South American Cricetid Radiation (Rodentia, Sigmodontinae). *PLoS ONE*, 9, e100687. <https://doi.org/10.1371/journal.pone.0100687>
- Leite, R. L., & Rogers, D. S. (2013). Revisiting Amazonian phylogeography: Insights into diversification hypotheses and novel perspectives. *Organism Diversity & Evolution*, 13, 639–664. <https://doi.org/10.1007/s13127-013-0140-8>
- Leutner, B., & Horning, N. (2016). RStoolbox: tools for remote sensing data analysis (Version 0.1.4). R package. Retrieved from <https://cran.r-project.org/web/packages/RStoolbox/index.html>
- Lima, N. E., Lima-Ribeiro, M. S., Tinoco, C. F., Terribile, L. C., & Collevatti, R. G. (2014). Phylogeography and ecological niche modelling, coupled with the fossil pollen record, unravel the demographic history of a Neotropical swamp palm through the Quaternary. *Journal of Biogeography*, 41, 673–686. <https://doi.org/10.1111/jbi.12269>
- Machado, L. F., Leite, Y. L. R., Christoff, A. U., & Giuliano, L. G. (2013). Phylogeny and biogeography of tetralophodont rodents of the tribe Oryzomyini (Cricetidae: Sigmodontinae). *Zoologica Scripta*, 43, 119–130. <https://doi.org/10.1111/zsc.12041>
- Massatti, R., & Knowles, L. L. (2014). Microhabitat differences impact phylogeographic concordance of codistributed species: Genomic evidence in montane sedges (*Carex* L.) from the Rocky Mountains. *Evolution*, 68, 2833–2846. <https://doi.org/10.1111/evo.12491>
- Mayle, F. E., Burbridge, R., & Killeen, T. J. (2000). Millennial-scale dynamics of southern Amazonian rain forests. *Science*, 290, 2291–2294. <https://doi.org/10.1126/science.290.5500.2291>

- McGlue, M. M., Silva, A., Assine, M. L., Stevaux, J. C., & Pupim, F. do N. (2015). Climate change patterns in Amazonia and biodiversity. In I. Bergier & M. L. Assine (Eds.), *Dynamics of the Pantanal wetland in South America* (pp. 55–81). Cham, Switzerland: Springer.
- McRae, B. H. (2006). Isolation by resistance. *Evolution*, *60*, 1551. <https://doi.org/10.1111/j.0014-3820.2006.tb00500.x>
- Muscarella, R., Galante, P. J., Soley-Guardia, M., Boria, R. A., Kass, J. M., Uriarte, M., & Anderson, R. P. (2014). ENMeval: An R package for conducting spatially independent evaluations and estimating optimal model complexity for Maxent ecological niche models. *Methods in Ecology and Evolution*, *5*, 1198–1205. <https://doi.org/10.1111/2041-210X.12261>
- Naka, L. N., Bechtoldt, C. L., Henriques, L. M. P., & Brumfield, R. T. (2012). The role of physical barriers in the location of avian suture zones in the Guiana Shield, Northern Amazonia. *The American Naturalist*, *179*, 115–132. <https://doi.org/10.1086/664627>
- Novembre, J., & Stephens, M. (2008). Interpreting principal component analyses of spatial population genetic variation. *Nature Genetics*, *40*, 646–649. <https://doi.org/10.1038/ng.139>
- Oksanen, J., Blanchet, F. G., Friendly, M., Kinda, R., Legendre, P., & McGlinn, D., ... Wagner, H. (2017). vegan: Community ecology package (version 2.4-4). R package. Retrieved from <https://cran.r-project.org/web/packages/vegan/index.html>.
- Papadopoulou, A., & Knowles, L. L. (2015). Species-specific responses to island connectivity cycles: Refined models for testing phylogeographic concordance across a Mediterranean Pleistocene Aggregate Island Complex. *Molecular Ecology*, *24*, 4252–4268. <https://doi.org/10.1111/mec.13305>
- Papadopoulou, A., & Knowles, L. L. (2016). Toward a paradigm shift in comparative phylogeography driven by trait-based hypotheses. *Proceedings of the National Academy of Sciences of the United States of America*, *113*, 8018–8024. <https://doi.org/10.1073/pnas.1601069113>
- Pardiñas, U. F. J., & Teta, P. (2011). Fossil history of the marsh rats of the genus *Holochilus* and *Lundomys* (Cricetidae, Sigmodontinae) in southern South America. *Estudios Geológicos*, *67*, 111–129. <https://doi.org/10.3989/egcol.40347.136>
- Pardiñas, U. F. J., Teta, P., Voglino, D., & Fernández, F. J. (2013). Enlarging rodent diversity in west-central Argentina: A new species of the genus *Holochilus* (Cricetidae, Sigmodontinae). *Journal of Mammalogy*, *94*, 231–240. <https://doi.org/10.1644/12-MAMM-A-216>
- Peterson, M. A., & Denno, R. F. (1998). The influence of dispersal and diet breadth on patterns of genetic isolation by distance in phytophagous insects. *The American Naturalist*, *152*, 428–446. <https://doi.org/10.1086/286180>
- Peterson, B. K., Weber, J. N., Kay, E. H., Fisher, H. S., & Hoekstra, H. E. (2012). Double digest RADseq: An inexpensive method for de novo SNP discovery and genotyping in model and non-model species. *PLoS ONE*, *7*, e37135. <https://doi.org/10.1371/journal.pone.0037135>
- Phillips, S. J., Anderson, R. P., & Schapire, R. E. (2006). Maximum entropy modeling of species geographic distributions. *Ecological Modelling*, *190*, 231–259. <https://doi.org/10.1016/j.ecolmodel.2005.03.026>
- Purcell, S., Neale, B., Todd-Brown, K., Thomas, L., Ferreira, M. A., Bender, D., & Sham, P. C. (2007). PLINK: A tool set for whole-genome association and population-based linkage analyses. *American Journal of Human Genetics*, *81*, 559–575. <https://doi.org/10.1086/519795>
- Quattrocchio, M. E., Borromei, A. M., Deschamps, C. M., Grill, S. C., & Zavala, C. A. (2008). Landscape evolution and climate changes in the Late Pleistocene-Holocene, southern Pampa (Argentina): Evidence from palynology, mammals and sedimentology. *Quaternary International*, *181*, 123–138. <https://doi.org/10.1016/j.quaint.2007.02.018>
- R Core Team. (2017). *R: A language and environment for statistical computing*. Vienna, Austria: R Foundation for Statistical Computing.
- Rocha, R. G., Ferreira, E., Fonseca, C., Justino, J., Leite, Y. L. R., & Costa, L. P. (2014). Seasonal flooding regime and ecological traits influence genetic structure of two small rodents. *Ecology and Evolution*, *4*, 4598–4608. <https://doi.org/10.1002/ece3.1336>
- Shah, V. B., & McRae, B. H. (2008). Circuitscape: A tool for landscape ecology. *Proceedings of the 7th Python in Science Conference (SciPy 2008)* Pasadena, CA, August 19–24, 2008, 62–66.
- Smith, B. T., McCormack, J. E., Cuervo, A. M., Hickerson, M. J., Aleixo, A., & Cadena, C. D. (2014). The drivers of tropical speciation. *Nature*, *515*, 406–409. <https://doi.org/10.1038/nature13687>
- Stein, B. R. (1988). Morphology and allometry in several genera of semi-aquatic rodents (*Ondatra*, *Nectomys*, and *Oryzomys*). *Journal of Mammalogy*, *69*, 500–511. <https://doi.org/10.2307/1381341>
- Swets, J. A. (1988). Measuring the accuracy of diagnostic systems. *Science*, *240*, 1285–1293. <https://doi.org/10.1126/science.3287615>
- Thom, G., Amaral, F. R. D., Hickerson, M. J., Aleixo, A., Araujo-Silva, L. E., Ribas, C. C., ... Miyaki, C. Y. (2018). Phenotypic and genetic structure support gene flow generating gene tree discordances in an Amazonian floodplain endemic species. *Systematic Biology*, *67*, 700–718. <https://doi.org/10.1093/sysbio/syy004>
- Urrego, D. H., Silman, M. R., & Bush, M. B. (2005). The Last Glacial Maximum: Stability and change in a western Amazonian cloud forest. *Journal of Quaternary Science*, *20*, 693–70. <https://doi.org/10.1002/jqs.976>
- Wang, C. L., Zöllner, S., & Rosenberg, N. A. (2012). A quantitative comparison of the similarity between genes and geography in worldwide human populations. *Plos Genetics*, *8*, e1002886. <https://doi.org/10.1371/journal.pgen.1002886>
- Warren, D. L., & Seifert, S. N. (2011). Ecological niche modeling in Maxent: The importance of model complexity and the performance of model selection criteria. *Ecological Applications*, *21*, 335–342. <https://doi.org/10.1890/10-1171.1>
- Whitney, B. S., Mayle, F. E., Punyasena, S. W., Fitzpatrick, K. A., Burn, M. J., Guillen, R., ... Metcalfe, S. E. (2011). A 45kyr palaeoclimate record from the lowland interior of tropical South America. *Palaeogeography, Palaeoclimatology, Palaeoecology*, *307*, 177–192. <https://doi.org/10.1016/j.palaeo.2011.05.012>

BIOSKETCHES

The authors share an interest in the study of processes that structure the genetic variation among taxa and across geography. L.L.K., J.R.P. and A.R.P. conceived the idea; J.R.P. collected the genomic data; J.R.P. and A.T.T. processed the genomic data; J.R.P., L.L.K. and A.T.T. performed the analyses; all authors worked on the writing.

SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section at the end of the article.

How to cite this article: Prado JR, Percequillo AR, Thomaz AT, Knowles LL. Similar but different: Revealing the relative roles of species-traits versus biome properties structuring genetic variation in South American marsh rats. *J Biogeogr.* 2019;46:770–783. <https://doi.org/10.1111/jbi.13529>