Quantifying the roles of ecology and geography in spatial genetic divergence

Ian J. Wang, Richard E. Glor, and Jonathan B. Losos

1Department of Organismic and Evolutionary Biology, Harvard University, Cambridge, MA, 02138, USA
2Department of Biology, University of Rochester, Rochester, NY, 14627, USA

*Correspondence: E-mail: ianwang@fas.harvard.edu

INTRODUCTION

Understanding the factors that contribute to population genetic divergence is a long-standing goal in ecology and evolution (Wright 1921; Mayr 1963; Coyne & Orr 2004). Patterns of genetic variation often reflect spatial variation in gene flow, which can be influenced by ecological landscapes in two important ways. Spatially separated populations may experience isolation-by-distance (IBD; Wright 1943), in which landscape barriers and geographical distances cause restricted gene flow, and isolation-by-environment (IBE; Wang & Summers 2010), in which gene flow among populations inhabiting different ecological environments is limited either by selection against dispersers moving between them or by individual preference to remain in a particular environment due to local adaptation (Dobzhansky 1937). IBD predicts a correlation between genetic divergence and geographical factors such as landscape barriers and geographical distance, and IBE predicts a correlation between genetic divergence and environmental dissimilarity, because greater environmental differences between populations are expected to be associated with stronger divergent selection and reduction in the success of dispersers (Crispo et al. 2006; Lee & Mitchell-Olds 2011). Of course, geographical and environmental isolation are not mutually exclusive, and spatial genetic divergence among populations can result from reduced gene flow associated with both geographical and ecological factors (e.g. Coyne & Orr 2004; Crispo et al. 2006; Thorpe et al. 2008).

Despite the large body of research on patterns of genetic divergence, few studies have quantified the contributions of IBD and IBE to spatial genetic divergence, especially at comparative scales necessary to investigate the strengths and prevalence of these relationships in nature (Thorpe et al. 2008; Schluter 2009; Sobel et al. 2010). Historically, studies of population divergence have focused on IBD (Wright 1943), while recent studies have argued that IBE plays a dominant role in the evolution of genetic divergence (Thorpe et al. 2008; Nosil 2012). The relative contributions of these two factors is still debated, and understanding their effects on the reduction of gene flow among populations can inform our understanding of how landscapes and environments shape patterns of genetic variation in nature (Cushman et al. 2006; Wang & Summers 2010; Lee & Mitchell-Olds 2011).

Because populations that are geographically distant also tend to occupy different environments, disentangling the effects of IBD and IBE is inherently difficult and remains a major challenge. However, the rise of modern spatial statistical methods and the increasing availability of high-resolution geographical and environmental data layers now makes it possible to accurately describe geographical and ecological landscapes and to simultaneously estimate the effects of IBD and IBE on spatial genetic divergence. The genetic consequences of IBD and IBE are often best studied in widespread organisms with spatially structured population divergence (Thorpe et al. 2004; Sobel et al. 2010). Thus, comparative analyses of such species can provide valuable insights into the evolution of genetic divergence among populations.

In this study, we use a novel implementation of SEM to quantify the contributions of IBD and IBE to spatial genetic divergence in 17 widespread species in the diverse adaptive radiations of Anolis lizards on the Greater Antilles. We used this approach to test predictions stemming from three alternative hypotheses: (1) geographical distances and barriers limit gene flow between populations such that IBD contributes significantly to spatial genetic divergence, (2) divergent selection in different environments reduces gene flow between populations such that IBE contributes significantly to spatial genetic divergence, and (3) geographical distances and divergent selection each influence gene flow such that IBD and IBE both contribute significantly to spatial genetic divergence. In all cases, we quantified the relative strengths of IBD and IBE for each species. Previous work suggests that anole species often inhabit heterogeneous environments (Losos & Ricklefs 2009), have substantial levels of genetic variation (e.g. Glor et al. 2003; Kolbe et al. 2004) and adapt to local ecological conditions (e.g. Thorpe et al. 2004; Calisbeek et al. 2006), making them excellent study systems for examining the evolution of spatial genetic divergence (Ogden &
Thorpe 2002; Thorpe et al. 2008). By investigating numerous widespread species across four large islands, we were able to conduct a comparative analysis of the factors affecting spatial genetic variation across ecologically diverse species and landscapes.

MATERIALS AND METHODS

Genetic data

We assembled neutral DNA sequence data sets for 17 widespread species of *Anolis* lizards, each inhabiting one of the four Greater Antillean islands. For 12 species, we acquired sequence data from previously published studies: *Anolis allisoni* (ND2 and tRNAs; Glor et al. 2004), *Anolis cooki* (ND2 and cytB; Jezkova et al. 2009), *Anolis chlorocyamus, Anolis cristatellus, Anolis cybotes* (ND2 and tRNAs; Kolbe et al. 2007), *Anolis distichus dominicensis, Anolis distichus ignigidens* (ND2 and tRNAs; Glor & Laport 2012), *Anolis krugi* (ND2 and tRNAs; Rodriguez-Robles et al. 2010), *Anolis poncensis* (ND2 and cytB; Jezkova et al. 2009), *Anolis porcatus* (ND2 and tRNAs; Glor et al. 2004), *Anolis sagrei* (ND2 and tRNAs; Kolbe et al. 2004) and *Anolis whitemani* (ND2, tRNAs, ND1, COI and control region; Glor et al. 2009). For five other species, *Anolis equestris, Anolis garmani, Anolis grahami, Anolis lineatopus* and *Anolis stratulus*, we collected DNA sequences from the ND2 gene and adjacent tRNA<sub>TRP</sub> and tRNA<sub>ALA</sub> following previously published methods (Glor & Laport 2012). To be included in our study, a species had to have genetic data collected from multiple localities with precise GPS coordinates providing < 100 m accuracy (at least four significant decimal-degree digits) or with locality information detailed enough to acquire coordinates with this accuracy (Fig. 1). In practice, most localities were georeferenced on-site using hand-held GPS units. For this study, we consider the subspecies of *A. distichus* to be distinct species, based on a recent report recommending that they be elevated to species status (Glor & Laport 2012).

For our mtDNA data set, we calculated between-locality genetic divergence using a maximum composite likelihood model of nucleotide evolution that allowed for gamma-distributed rate variation among sites in MEGA (Tamura et al. 2007). We then used these data to construct a matrix of genetic distances between localities in each species. For three species for which nDNA data were available (*A. allisoni, A. chlorocyamus and A. porcatus*), we also constructed nDNA genetic distance matrices and concatenated mtDNA and nDNA distance matrices. We considered point localities to be the relevant sampling units in our study because our method does not rely on using a metapopulation model of reproductively isolated populations and, like many other approaches to examining spatial genetic patterns, can be applied to species with individuals distributed continuously on a landscape (Rousset 2000; Cushman et al. 2006). In practice, *Anolis* lizards are typically highly territorial and have small home ranges (Schoener & Schoener 1982; Johnson et al. 2009), suggesting that these species will have compact genetic neighbourhoods (Wright 1943) that should generally be smaller than the distances between our sampling localities.

Environmental data

We acquired geographic information systems (GIS) data layers for a total of 24 environmental and geographical variables, including 19 bioclimatic variables with 1-km resolution from the WorldClim database (http://www.bioclim.org), four vegetation variables from the base (http://www.bioclim.org), four vegetation variables from the base (http://www.bioclim.org), four vegetation variables from the base (http://www.bioclim.org), four vegetation variables from the base (http://www.bioclim.org), four vegetation variables from the base (http://www.bioclim.org), four vegetation variables from the base (http://www.bioclim.org), four vegetation variables from the base (http://www.bioclim.org), four vegetation variables from the base (http://www.bioclim.org), four vegetation variables from the base (http://www.bioclim.org), four vegetation variables from the base (http://www.bioclim.org), four vegetation variables from the base (http://www.bioclim.org), four vegetation variables from the base (http://www.bioclim.org), four vegetation variables from the base (http://www.bioclim.org), four vegetation variables from the base (http://www.bioclim.org), four vegetation variables from the base (http://www.bioclim.org), four vegetation variables from the base (http://www.bioclim.org), four vegetation variables from the base (http://www.bioclim.org), four vegetation variables from the base (http://www.bioclim.org), four vegetation variables from the MODIS land cover database with 250-m resolution (leaf area index, normalised difference vegetation index, tree density and herbaceous density; http://modis.gsfc.nasa.gov/) and a digital elevation model with 30-m resolution from the USGS EROS database (http://eros.usgs.gov). We extracted values for each variable at every locality for each species using ArcGIS (ESRI) and calculated differences between localities to create a dissimilarity matrix for each variable.

Geographical distances

Resistance-based (also known as cost-weighted) distances have been shown to more closely reflect gene flow between populations than direct distances for a variety of organisms (Cushman et al. 2006; McRae & Beier 2007). These distances account for variation in the ease of movement across a heterogeneous landscape by assigning relative resistances (or costs) to different landscape features (McRae & Beier 2007; Storfer et al. 2010). To assign resistances to each cell in a landscape data layer for each species, we followed a method
for translating niche model suitability scores into resistances (Wang et al. 2008). We first created a niche model for each species using MAXENT (Phillips et al. 2006), which uses occurrence data and environmental layers to predict the environmental suitability of each cell in the study area. As input data, we used the 24 environmental and geographical data layers and an average of 110 occurrence records for each species (Table 1). Occurrence records were collected from our data, previously published studies and the HerpNet/VertNet species database of museum records from 55 institutions (www.vert-net.org). To evaluate how accurately the resulting GIS layer reflects the suitability of the landscape to the species being modelled, we performed a receiver operating characteristic (ROC) analysis (Phillips et al. 2006) and found that it indicated a good fit (AUC > 0.858 in all cases) to the species distribution model for each species (Phillips et al. 2006). MAXENT estimates environmental suitability values ranging from 0–1, and higher scores indicate more suitable habitat. We used the reverse of these values, because lower suitability should have a higher resistance, to assign resistances to each cell in the study area of each species.

We then used the resulting resistance layers to obtain geographical distance matrices using two alternative measures of geographical distance: (1) least-cost path distances calculated using ArcGIS and (2) circuit distances calculated using CircuitScape (McRae & Beier 2007). Least-cost path distance is calculated by searching for the path that minimises the total cumulative cost (or resistance) between two points (Wang et al. 2009), and circuit distance is calculated by summing the costs of all possible paths between two points (McRae & Beier 2007). For least-cost path distances, our estimates were adjusted for the additional distance between points imposed by topographical relief based on the digital elevation model; this adjustment is not currently available in CircuitScape. These analyses resulted in two matrices of the geographical distances between localities for each species.

Although the geographical distance and environmental dissimilarity estimates both use the same set of 24 environmental variables, those variables are actually utilised very differently. In one case (environmental dissimilarity), only the raw values at specific points are considered, whereas in the other (geographical distance), the full data layers are processed to infer another variable, habitat suitability, which is then used in measuring the weighted distances along paths (or sets of paths) between localities. Thus, geographical distance and environmental dissimilarity are not necessarily correlated. For instance, if two populations inhabited localities with identical scores for each environmental variable, they would have an environmental dissimilarity of zero but a geographical distance greater than zero as long as they have any spatial separation. Nevertheless, to make sure that any associations with geographical distance were not driven by the inclusion of environmental variables, we also calculated topographic distances between localities.

### Structural equation modelling

To estimate the relative contributions of geographical distance and environmental dissimilarity to genetic divergence, we utilised a structural equation modelling (SEM) framework (Grace 2006). Originally developed by the geneticist Sewall Wright (1921), SEM is a statistical framework for evaluating complex relationships between multiple variables that uses a series of regression and model-fitting analyses to calculate correspondence among any number of variables whose relationships are hypothesised a priori (Grace 2006; Santos & Cannatella 2011). SEM is, therefore, an ideal framework for testing hypotheses about the contributions of a large number of different variables associated with geographical distance and environmental variation to genetic divergence. Specifically, we estimated the strength of IBD from the effects of geographical distance on genetic divergence while controlling for environmental dissimilarity.

<table>
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and we estimated the strength of IBE from the effects of environmental dissimilarity on genetic divergence while controlling for geographical distance.

One of the advantages of SEM is that it can use latent variables—variables that are not directly measured but instead inferred from a set of observed variables. In our model, we described geographical distance and environmental dissimilarity as latent variables because we wanted to infer the best-fitting values rather than assume that geography or environment could be summarised by some arbitrary method. Essentially, geographical distance and environmental dissimilarity are both complex variables that are difficult to measure, because determining the biologically relevant variables is complicated and we had no a priori expectation for how those variables should be weighted. Thus, modelling these as latent variables was a more sensible and conservative approach. We inferred the geographical distance latent variable from our two measurements of distance (least-cost path distance and circuit distance), and we inferred the environmental dissimilarity latent variable from our set of 24 environmental variables. Prior to analysis, all observed variables were standardised. We also defined regression pathways between geographical distance and genetic divergence, and between environmental dissimilarity and genetic divergence, and a covariance pathway between geographical distance and environmental dissimilarity (Fig. 2). The estimated coefficients of these pathways can be used to quantify the relative effects of IBD and IBE on spatial genetic divergence, while controlling for covariation between geographical distance and environmental dissimilarity (‘Covar.’ in Fig. 2).

We solved the model by simultaneously estimating all parameters using maximum-likelihood estimation (MLE) in the ‘lavaan’ package in R (http://cran.r-project.org/web/packages/lavaan/). By estimating the parameters simultaneously, we were able to optimise the fit of each parameter, including the regression pathways connecting latent variables and the indicator pathways used to manifest the latent variables, while controlling for covariation among variables. So, we estimated the best fit of geographical distance and environmental dissimilarity to genetic divergence while determining the best estimates of geographical distance and environmental dissimilarity from their respective indicator variables. Standard errors for each parameter were determined by calculating Huber–White robust standard errors (White 1980) based on the residuals of the fitted model. We incorporated a matrix permutation procedure for significance testing because of the distance matrix format of our data and assessed the overall goodness of fit for each model using the Akaike information criterion (AIC), which provides a metric for model selection by comparing the likelihoods of a set of models while penalising for their numbers of parameters. Lower AIC scores indicate closer fit to the true model, and models with AIC scores that exceed the lowest model by 10 or more are not supported (Burnham & Anderson 1998).

To test whether geographical distance and environmental dissimilarity contributed significantly to the model, we compared the AIC from the full model to the AICs from a model excluding the geographical variables and from one excluding the environmental variables. We performed the SEM analysis using models that tested three alternative hypotheses: (1) only geographical distance contributes significantly to genetic divergence, (2) only environmental dissimilarity contributes significantly to genetic divergence and (3) geographical distance and environmental dissimilarity each contribute significantly to genetic divergence (Fig. 2). We performed this analysis on mtDNA sequence divergence from all 17 species included in our study. We examined the results for each species individually (Table 1), and, to draw comparisons between islands, we also averaged the results of all species on each island.

Method validation

The development of methods for the study of spatial genetic variation is still very fluid, and universally suitable analytical frameworks are yet to emerge (Storfer et al. 2010). In general, evaluating a new method requires understanding its accuracy, its assumptions and its advantages and disadvantages. To confirm the consistency of our results with other similar methods, we performed two additional multiple regression-based analyses on our mtDNA data set. First, we performed generalised dissimilarity modelling (GDM; Ferrier et al. 2007), using the ‘GDM’ package in R (Ferrier et al. 2007). GDM allows for nonlinear relationships between predictor and response variables, but only currently implements direct distance (instead of resistance-based distances), because the geographical distance matrix is computed from raw GPS coordinates. Second, we performed a variation partitioning analysis (Legendre & Fortin 2010) using the ‘vegan’ package in R (Oksanen et al. 2007). Variation partitioning uses multiple regression to assign proportions of variation in a response variable to a set of explanatory variables, and canonical redundancy analysis to assess the significance of each proportion. Although similar in principal to our SEM method, there are notable differences. For instance, SEM uses latent variables, which allows for some variables to be inferred from others and for the relationships among all variables to be estimated simultaneously. Because GDM and variation partitioning do not use latent variables, for these analyses, we first reduced the number of predictor variables in our data set by performing a principal components analysis (PCA) on the 24 environmental variables using the ‘prcomp’ function in R. In addition, unlike GDM and variation partitioning, SEM estimates error around the maximum-likelihood estimate for each parameter and provides an information-theoretic output for assessing parameter importance. Thus, for our goals, SEM provides several distinct advantages over other similar methods. However, while SEM is powerful for isolating the effects of individual variables, it cannot be used to estimate the effects of variation shared among variables. For

Figure 2  Graphical representation of the structural equation model used to quantify the relationships between geographical distance, environmental dissimilarity and genetic divergence. Observed variables are enclosed in boxes, whereas latent variables are enclosed in circles (geographical distance and environmental dissimilarity). Single-headed arrows show the observed variables used to infer each latent variable. Solid double-headed arrows indicate regression pathways, and dashed double-headed arrows indicate covariance pathways.

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our application, this means that we can use SEM to disentangle the effects of geographical and environmental distances on genetic divergence individually, but that we cannot quantify the effects of shared variation between geographical and environmental distances on genetic divergence, a drawback that also exists with GDM but not variation partitioning. Finally, an assumption of each methods is that inferences made using current data, both environmental and genetic, reflect the processes that produced genetic divergence. Thus, these analyses are most reliably applied to genetic markers that capture recent and ongoing processes and when the contemporary geography and environment of the study area has not changed substantially since patterns of genetic variation were established.

To test whether the form in which we used environmental variables affected the results, we repeated the SEM analysis for each species using environmental PC axes or topographic distances. To visualise the relationships between geographical distance, environmental dissimilarity and genetic divergence, we plotted genetic divergence vs. geographical distance, genetic divergence vs. environmental dissimilarity and geographical distance vs. environmental dissimilarity for each species (Fig. S1). For these plots, we averaged least-cost path and circuit distances to estimate geographical distance and calculated the distances between points for each population plotted against the two major environmental PC axes to estimate environmental dissimilarity.

Finally, to compare results from different molecular markers, we repeated the analysis on mtDNA and nDNA sequences together and on nDNA sequences alone from the three species \((A.\ allisoni, A.\ chloroxyanus\) and \(A.\ porcatus\)) for which extensive nDNA sampling was available. Because nDNA data were not available from all localities for these species, analyses on nDNA alone and concatenated mtDNA and nDNA sequences were performed on a subset of the localities used in mtDNA analyses (Table S3). To allow comparisons between different molecular markers, we also ran the SEM analysis on mtDNA data alone from the subset of localities with nDNA data.

RESULTS

Genetic, geographical and environmental data

We assembled mtDNA sequence data sets for 17 widespread anole species and nDNA sequence data sets for three species. For our mtDNA data sets, we acquired an average sequence length of 1527 bp from 72.5 individuals from 21 localities per species. We found a wide range of variation (from < 0.5% between some localities to > 15% among others) with an average of 5.6% sequence divergence between localities within each species. The 24 environmental GIS layers used in our study indicated that each of the four Greater Antillean islands is environmentally heterogeneous, with each species typically being distributed across a broad range of environmental conditions (Fig. 1). The two cost-weighted distances we calculated (least-cost path and circuit distances) were highly correlated \((r^2 > 0.65\) in all cases) and roughly equally weighted in estimating a geographical distance latent variable (Table S1).

Structural equation modelling

The SEM analysis revealed that IBD explained a significant proportion of genetic divergence for 15 species, and IBE explained a significant proportion of genetic divergence in 13 species (Table 1; Fig. 3). In each case, the full model including both IBD and IBE was a significantly better fit to the data based on model-fit scores (AIC) than a model including only one or the other (Table S2). There was also, as expected, some covariation between geographical distance and environmental dissimilarity, but this was typically low (Mean = 15.2%; Table 1). However, for three species \((A.\ cooki, A.\ distichus\) ignigularis and \(A.\ eques\) triz), covariation between these variables was fairly high (\(> 0.3\)), and these results should be interpreted with caution. On average, IBD explained 36.3% of genetic divergence \((range = 3.0–79.6\%; Table 1; Fig. 3), whereas IBE explained 17.9% \((range = 0.1–47.7\%; Table 1; Fig. 3). We also identified the individual variables that contributed to making up the environmental dissimilarity latent variable in each species (Table 1; Table S1).

Across islands, the average results were fairly consistent, except that IBE actually contributed more than IBD in the species from Hispaniola \((0.173 \pm 0.172\) IBD vs. \(0.226 \pm 0.157\) IBE), and IBE contributed very little in any of the species from Jamaica \((0.510 \pm 0.178\) IBD vs. \(0.004 \pm 0.003\) IBE). For species from Cuba \((0.543 \pm 0.169\) IBD vs. \(0.158 \pm 0.076\) IBE) and Puerto Rico \((0.329 \pm 0.146\) IBD vs. \(0.216 \pm 0.154\) IBE), IBD contributed 3.5 times and 1.5 times as much, respectively, as IBE to spatial genetic divergence.

Method validation

In general, results from GDM and variation partitioning, both of which have been recently described as powerful methods for studying genetic divergence (Ferrier et al. 2007; Thomassen et al. 2010; Legendre & Fortin 2010), were consistent with those inferred by SEM (Table S3). In addition, as expected, when we used simple topographic distances instead of resistance-based distances, the contribution from geography was lower, but the effects of environment were virtually unchanged (Table S3). When we performed a
PCA on the 24 environmental variables and entered the resulting PC axes as environmental dissimilarity variables — instead of simultaneously estimating the factor loadings for each variable when solving the model — the contribution from IBE was slightly lower, whereas the effects of IBD were basically the same (Table S3). Finally, SEM analysis of mtDNA and nDNA data produced concordant results in each case. For *A. porcatus*, analysis on nDNA data independently resulted in estimates that were very similar to those estimated from mtDNA alone for the strengths of IBD (67.3 vs. 77.6%) and IBE (8.8 vs. 5.7%) (Table 2). For *A. allisoni* and *A. chlorocyanaus*, we did not find sufficient variation in nDNA sequences to permit convergence of the MLE during SEM analysis of nDNA alone. However, the addition of nDNA to the mtDNA data set resulted in only minor differences compared with mtDNA alone (Table 2).

### DISCUSSION

**Structural equation modelling**

Disentangling the effects of IBD and IBE has been a major challenge in identifying the processes that drive spatial genetic divergence (Crispo et al. 2006; Thorpe et al. 2008; Wang & Summers 2010). We used a structural equation modelling approach to isolate and quantify the relative strengths of these effects, highlighting the importance of simultaneously considering the roles of both ecological and geographical factors in driving biological diversification. For most species, we found support for the hypothesis that both geography and ecology contribute to spatial genetic divergence, but, on average, IBD explained about twice as much genetic divergence as IBE (mean 36.3 vs. 17.9%; Table 1; Fig. 3). In most cases, these two factors together explained a large proportion of genetic variation (Fig. 3), suggesting that our SEM analysis captured the key underlying processes. Hence, although several recent studies have suggested a dominant role for IBE (e.g. Thorpe et al. 2008; Surget-Groba et al. 2012), our results clearly show that IBD still plays a key role in the evolution of spatial divergence.

There are, however, exceptions to this trend. IBE actually contributed more than IBD for four Hispaniolan taxa (*A. chlorocyanaus, A. distichus dominicanus, A. distichus iguanastris* and *A. whitmanii*) and one Puerto Rican species (*A. cristatellus*). Meanwhile, for three Jamaican species (*A. garmani, A. grahami* and *A. lineatopus*), IBE played a very minor role, explaining an average of 0.4% of genetic divergence. The reasons for these differences are unclear. On one hand, overall environmental heterogeneity (variation averaged across all environmental variables after standardisation) was greatest on Hispaniola, and Hispaniola had the highest average contribution from environmental dissimilarity (22.6%), but on the other, environmental heterogeneity was not significantly different between the remaining three islands. Variation in environmental variables among sampling localities for each species was also not significantly different between Cuba, Jamaica and Puerto Rico, suggesting that the low contributions from IBE were not simply due to low levels of environmental variation.

In general, the results for each species within islands, except for Hispaniola, were fairly consistent, suggesting that closely related species have similar evolutionary responses to a particular landscape. We also found, though, that results were broadly consistent across islands as well, suggesting that how organisms interact with their environments and the underlying features of the landscape both play important roles. On average, and for 11 of 12 species on three of the four islands (Cuba, Jamaica, and Puerto Rico), IBD played a greater role than IBE. This result joins several recent studies in suggesting that the ubiquity of IBE may be overemphasised and that the important drivers of genetic divergence may vary substantially across different organisms (Crispo et al. 2006; Hendry 2009). In fact, in many cases, both factors may contribute to some extent to spatial genetic divergence (Lee & Mitchell-Olde 2011).

For those species for which IBE played a significant role, simultaneously estimating the contribution of each environmental GIS layer in the SEM allowed us to assess the degree to which individual environmental variables contributed to these results (Table 1; Table S1). Spatial variation in adaptations to thermal environment has been shown in some *Anolis* lizards (Hertz & Huey 1981), but morphological and genetic variation within some species seem to be more closely tied with spatial variation in precipitation or vegetation (Thorpe 2002; Calsbeek et al. 2006; Thorpe et al. 2008). For most species in our study, temperature variables played the biggest part in describing the environmental dissimilarity latent variable, but precipitation variables contributed significantly to environmental dissimilarity in eight species. Understanding how individual environmental variables can produce adaptive differences that lead to IBE remains a major challenge (Coyne & Orr 2004; Sobel et al. 2010), and our results demonstrate that doing so can contribute substantially to understanding the evolution of spatial genetic divergence.

<p>| Table 2 | Results of structural equation modelling (SEM) analyses that included nuclear DNA sequence data (nDNA). Data used include the number of localities (Loc.) and the number (Seq.) and length in base pairs (bp) of DNA sequences. For each species, SEM was used to quantify the proportion of genetic divergence explained by isolation-by-distance (IBD) and isolation-by-environment (IBE), presented as maximum-likelihood estimates ± standard errors. Also listed are the primary contributors (Contrib. vars.) to the environmental dissimilarity latent variable (individual factor loading coefficients &gt; 0.7). Results of analyses on mtDNA data alone are included for comparison. |
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<th>Species</th>
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</table>
Method validation

All three approaches we used to validate our novel SEM methodology produced results consistent with those from our method, confirming the robustness of our approach to different assumptions and sources of data. In each case, the slight differences we observed are in line with the differences expected based on previous studies of genetic divergence (McRae & Beier 2007; Wang et al. 2009). Our validation results also showed consistency between the results drawn from mtDNA data sets and those including nDNA data. So, we have some evidence that the inferences drawn from our mtDNA data sets should be good representations of the results from overall genetic variation. In any case, although there are potential drawbacks for using mtDNA-only data sets (or any single gene data set) for some analytical goals, these typically pertain to methods like phylogenetic inference, in which any single gene tree may present an ‘incorrect’ view of population history due to the random nature of lineage sorting. For this study, we used mtDNA sequences as indicators of population genetic structure and genetic divergence, rather than for inferring phylogenetic history, and these data should be robust for these purposes because mtDNA are known to be sensitive indicators of population structure (Zink & Barrowclough 2008; Karl et al. 2012). In addition, simulations have shown that unlinked neutral loci can consistently detect IBE when divergent selection between environments is strong and dispersal is intermediate (Thibert-Plante & Hendry 2010). We do not measure either selection or dispersal directly in this study, but previous work has shown that divergent selection on Anolis lizards can be strong (Calsbeek & Irschick 2007), and FST estimates from some species suggest moderate levels of dispersal (Ng & Glor 2011; Surget-Groba et al. 2012). Thus, because they typically provide more variable sites than nDNA sequences, mtDNA sequence data continue to be valuable for estimating population structure and relative levels of genetic divergence among populations (Zink & Barrowclough 2008), and for the conditions typical in Anolis lizards, they should also be reliable for detecting IBE. In any case, the SEM method we describe could easily be applied to many additional forms of genetic data, including microsatellite, SNP or genomic datasets, and future studies using these data should illuminate the factors influencing spatial genetic divergence even further.

While for this study, we examined patterns of gene flow through neutral genetic divergence, our approach could also easily be extended to examining variation in genes potentially under selection. Studies examining many genetic loci could look for outliers that have a significantly different signal from the background pattern of loci from across the genome. Those linked to genes under selection might be expected to exhibit greater degrees of ecological isolation because of divergent selection in different environments. Likewise, populations in similar environments should have less divergence in genes under selection than in neutral genes. Close examinations of the environmental variables contributing to these effects could prove especially valuable for linking the genes under selection to specific environmental forces of selection, and should provide insights into the geographical and ecological factors that contribute to adaptive genetic divergence.

CONCLUSIONS

Our results provide strong evidence that both IBD and IBE contributed to the evolution of spatial genetic divergence in the adaptive radiations of Anolis lizards on the Greater Antilles. IBD, in which physical distance and barriers reduce gene flow between populations, played the strongest role. IBE, in which adaptive differences or natural selection against dispersers limit gene flow between divergent habitats, played a secondary role. Experimental evidence has shown that anoles avoid microhabitats in which they have reduced performance (Irschick & Losos 1999; Calsbeek & Irschick 2007), and our results suggest that the same phenomenon may also occur among environmentally distinct macrohabitats. Overall, these results show that understanding the evolution of spatial genetic variation requires examining both ways in which ecological landscapes influence patterns of gene flow among populations.

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AUTHOR CONTRIBUTIONS

IJW, REG and JBL conceived and designed the study. REG contributed new genetic and species distribution data. IJW contributed the new spatial statistical methods and analysed the data. The authors wrote the manuscript together.

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