

Testing the role of ecology and life history in structuring genetic variation across a landscape: a trait-based phylogeographic approach

ANDREA PAZ,* ROBERTO IBÁÑEZ,† ‡ KAREN R. LIPS† § and ANDREW J. CRAWFORD*† ‡

*Department of Biological Sciences, Universidad de los Andes, A.A. 4976, Bogotá, Colombia, †Smithsonian Tropical Research Institute, Apartado 0843-03092, Panama City, Republic of Panama, ‡Círculo Herpetológico de Panamá, Apartado 0824-00122, Panama City, Republic of Panama, §Department of Biology, University of Maryland, College Park MD 20742-4415, USA

Abstract

Hypotheses to explain phylogeographic structure traditionally invoke geographic features, but often fail to provide a general explanation for spatial patterns of genetic variation. Organisms' intrinsic characteristics might play more important roles than landscape features in determining phylogeographic structure. We developed a novel comparative approach to explore the role of ecological and life-history variables in determining spatial genetic variation and tested it on frog communities in Panama. We quantified spatial genetic variation within 31 anuran species based on mitochondrial DNA sequences, for which hierarchical approximate Bayesian computation analyses rejected simultaneous divergence over a common landscape. Regressing ecological variables, on genetic divergence allowed us to test the importance of individual variables revealing that body size, current landscape resistance, geographic range, biogeographic origin and reproductive mode were significant predictors of spatial genetic variation. Our results support the idea that phylogeographic structure represents the outcome of an interaction between organisms and their environment, and suggest a conceptual integration we refer to as trait-based phylogeography.

Keywords: dispersal ability, DNA barcoding, ecological niche modelling, hierarchical approximate Bayesian computation, landscape resistance, phylogeography

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Introduction

A key first step in the generation of species diversity is limiting gene flow between populations, whether by natural selection or by simple physical isolation and allopatry (Mayr 1942). Understanding the factors that determine species' distributions therefore becomes a central goal in evolutionary biology (Gaston 2000; Wiens 2004). Historical processes that may have influenced the history of species' distributions may be inferred from shared geographic or temporal patterns of genetic subdivision among co-occurring species (Avice 1992; Bermingham & Martin 1998). When phylogeographic structure is shared among multiple, codis-

tributed species, the most parsimonious explanation is that they have responded to the same geographic or environmental events (Bermingham & Avice 1986; Zink 1996; Avice 2000; Arbogast & Kenagy 2001).

Comparative phylogeographic studies often support the hypothesis that geographic features such as mountains and rivers serve as vicariant barriers driving allopatric divergence and creating possible geminate species (Avice *et al.* 1987; Marko 2002). By reducing the movement of individuals, barriers reduce gene flow between populations and increase population structure (Wright 1951). Various hypotheses regarding types of geographic barriers have been tested. The riverine barrier hypothesis, for example, states that rivers are important in separating populations (Wallace 1852). A number of studies have been conducted that support (Ayres & Clutton-Brock 1992; Bates *et al.* 2004; Fouquet

Correspondence: Andrea Paz, Fax: (571) 3394949 x 1817; E-mail: paz.andreita@gmail.com

et al. 2012; Ribas *et al.* 2012) or reject (Gascon *et al.* 2000; Gehring *et al.* 2012) this hypothesis for different groups of animals. The Pleistocene refugia hypothesis states that during the cooler and drier period of each Pleistocene climatic cycle, suitable habitat for wet-forest species was broken into isolated blocks (Haffer 1969). While the refugia hypothesis has been supported in a few cases (Waltari *et al.* 2007; Byrne 2008), it has been widely rejected (Lessa *et al.* 2003), especially in cases where lineages are older than the Pleistocene (Rull 2008). Finally, while Pleistocene climate change has provided a general explanation for the phylogeographic patterning of many North American and European taxa (Hewitt 2000; Johnson & Cicero 2004), such geographic mechanisms of lineage diversification seem to fail as a general explanation across the tropical realm (Antonelli *et al.* 2010).

Given the lack of generality of geographic explanations of phylogeographic structure, we hypothesize that variation among codistributed species in phylogeographic histories could be predicted from variation in key ecological traits. Dispersal limitation is caused by the interaction between an organism and its environment, whose outcome is determined as much by the extrinsic environmental conditions as by the intrinsic biological properties of the organism (Bernardo *et al.* 2007). The physical height of a potential montane barrier, for example, may not limit species' distributions so much as the variation in moisture or temperature along an environmental gradient (Janzen 1967). Thus, we can think of the mountain range as a physiological rather than a physical barrier, suggesting that ecological rather than geographic variables may be more predictive of dispersal potential and thus of phylogeographic structure (Crawford *et al.* 2007; Burney & Brumfield 2009; Fine *et al.* 2013; Wang *et al.* 2013).

Amphibians are thought to be physiologically intolerant of novel environments due to their permeable skin (Navas & Otani 2007) as well as evolutionarily conservative in the breadth of their environmental niche (Wiens *et al.* 2006). Amphibians show substantial variation in their life-history characteristics (Crump 1974; Wells 2007); however, we might expect them to respond differentially to variation in environmental factors. Studies of Central American frogs have shown that phylogeographic histories of different species show few common spatial or temporal patterns across a shared landscape, leaving classical vicariant hypotheses with little predictive power (Weigt *et al.* 2005; Crawford *et al.* 2007; Wang *et al.* 2008; Robertson *et al.* 2009). These observations suggest that amphibians may be an ideal group to evaluate whether intrinsic variables can explain the observed divergence where geographic variables have thus far failed.

Here, we examine three frog communities in Panama using multiple co-occurring taxa to evaluate whether variation among species in intraspecific genetic divergence across a common landscape may be predicted by ecological variables potentially related to a frog's ability to disperse, survive and reproduce in a heterogeneous landscape. Each of three communities represents a mid-elevation montane site separated by stretches of lower elevation habitat. We evaluate four hypotheses of increasing complexity to explain average pairwise genetic divergence between localities, our surrogate for phylogeographic structure.

First, the simplest hypothesis assumes that physical landscape features alone are responsible for the separation of populations; thus, a single divergence event (i.e. similar levels of genetic divergence) should apply to all species across the shared landscape. Second, we ask whether variation in the altitudinal distribution of species might predict dispersal ability across the lowlands and therefore cause variation in genetic divergence and divergence times among codistributed species. In other words, is it more difficult for highland species to disperse through lowland habitats? In this analysis, however, elevation may be just a proxy for environmental factors (Janzen 1967); thus, we also sought to quantify variation among species in the environmental suitability of a common landscape. Thus, our third hypothesis is that interspecific variation in genetic divergence is predicted by variation in abiotic niche requirements of species, as quantified by species distribution models (Peterson 2001; Soberón & Nakamura 2009). Rather than looking at single species across a diverse landscape, as commonly done in landscape genetics (Storfer *et al.* 2007), here, we study a diversity of species across a single landscape in a comparative phylogeographic approach. Finally, we ask whether life-history traits themselves, such as developmental mode and body size, limit dispersal ability and thereby predict variation among species in pairwise genetic divergence between populations (Wollenberg Valero 2015). This latter approach is rarely attempted, but could become more common as phylogeographic studies are eventually applied to whole communities, for example, through DNA barcoding campaigns (e.g. Murphy *et al.* 2013), thus providing genetic data on a diversity of codistributed species.

Methods

Geographic and taxonomic sampling

Sampling was performed in three Panamanian national parks: *G. D. Omar Torrijos H.* National Park located in

the province of Coclé, *Chagres* National Park located in the provinces of Panamá and Colón, and Cana field station in the *Darién* National Park located in the Darién Province, referred to here as El Copé, Brewster and Cana, respectively (Fig. 1). We selected 31 species of frogs present in at least two of the three localities and with at least two individuals sampled per locality for a total of 481 individuals, 179 from El Copé, 190 from Brewster and 112 from Cana (Table S1, Supporting information). Sampling at each site varied in elevation: El Copé was sampled at 700–750 m, Brewster was sampled at 135–810 m, and Cana was sampled at 462–1320 m. Samples were collected prior to epizootic outbreaks and population declines at each site (Crawford *et al.* 2010).

Molecular data collection

Molecular data for this study consisted of DNA sequences from two mitochondrial gene fragments: cytochrome oxidase subunit 1 (COI) and the 16S ribosomal RNA gene. Molecular data from El Copé were published previously (Crawford *et al.* 2010), and sequence data for the other two localities were obtained with the same primers and protocols. Full-length COI sequences were 658 base pairs (bp) while full-length 16S fragments ranged from 491 to 684 bp with a species mean of 561 bp. Alignment of conspecific COI and 16S sequences was unambiguous and could be accomplished by eye. Gapped sites were excluded from the analyses. Specimen vouchers and GenBank numbers are provided in Table S2 (Supporting information).

Hierarchical approximate Bayesian computation analysis

One particularly effective method to detect correspondence in the temporal dimension between co-occurring taxa is hierarchical approximate Bayesian computation (HABC). HABC includes data from multiple loci and species in a single analysis within a coalescent framework to estimate interspecific congruence in divergence times (Hickerson *et al.* 2006a, 2010; Hickerson & Meyer 2008). Using coalescent simulations, this method accounts for uncertainty in current and ancestral population sizes and the stochastic nature of the mutational and coalescent processes in estimating genetic divergence among populations (Knowles & Maddison 2002; Hickerson *et al.* 2006b; Nielsen & Beaumont 2009). To test for simultaneous divergence, we performed HABC analyses with the program *MTML-msBayes* (Huang *et al.* 2011), following recent recommendations for implementation and analyses using this software (Oaks *et al.* 2012; Hickerson *et al.* 2014). Thirty species were analysed in the comparison between El Copé and Brewster, and fifteen species were compared between Cana and Brewster (Fig. 1). All pairwise comparisons involved conspecific samples except for four lineages treated as conspecific due to their low genetic divergence or lack of taxonomic resolution: the three species of *Atelopus*, one endemic to each site, *Pristimantis museosus* vs. its close relative, *P. aff. museosus*, *Craugastor opimus* vs. *C. megacephalus* and the as yet unnamed *Diasporus* spp. Conspecific samples included a minimum of two samples per population, as required for estimation of genetic divergences (Hickerson *et al.* 2007).

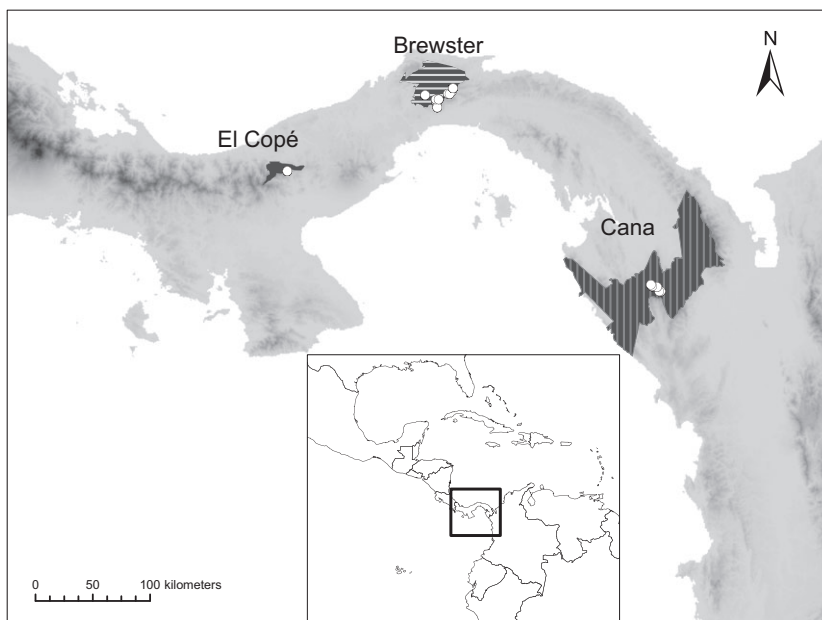


Fig. 1 Map of central and eastern Panama showing the three sites where all frog specimens were obtained: *G. D. Omar Torrijos H. National Park* (in black), *Chagres National Park* (horizontal lines) and *Cana field station in the Darién National Park* (vertical lines), referred to as El Copé, Brewster and Cana, respectively. White points within each site indicate exact localities where tissue samples were collected for genetic analyses.

MTML-msBayes is run as a 3-step process. First, several summary statistics including the net average pairwise differences (π_{net} , Nei & Li 1979), average pairwise differences between populations (π_{B} , Takahata & Nei 1985), average pairwise differences within populations (π_{S} , Takahata & Nei 1985) and Tajima's *D* (D_{T} , Tajima 1989) are calculated from the data (Hickerson *et al.* 2006a). Second, the program performs coalescent simulations of a data set of the same size as the observed data set, drawing randomly parameter values from a prior distribution and calculating summary statistics for each of the simulated data sets. Finally, summary statistics of the simulated data sets are compared to the observed estimates, and those parameter values that produced statistics differing from the observed data by less than a value ϵ are accepted (Hickerson *et al.* 2007; Huang *et al.* 2011). The values obtained from the acceptance/rejection step were subjected to a local linear regression to improve estimation. We used the average pairwise differences between each population pair (π_{B}) as our summary statistic. This statistic is easily computed and biologically relevant even in species with low sample sizes per population (Hickerson *et al.* 2007). π_{B} has been shown to perform well in the face of small sample sizes, and rejecting the null hypothesis of simultaneous divergence becomes more difficult using a single summary statistic. Thus, relying on only π_{B} as our summary statistic makes our test more conservative (Hickerson *et al.* 2007). To evaluate relative posterior support for number of divergence intervals, we used Bayes factors (BF, Hickerson & Meyer 2008). We used the BF results to constrain the number of intervals to their most likely values and reran the HABC analysis to obtain the most likely number of species in each divergence interval and the species membership within each of these temporal intervals.

Ecological niche modelling

We sought a composite measure of species-specific habitat suitability, as data are not available for all life-history variables of potential interest (e.g. resting metabolic rate or moisture uptake efficiency). We used ecological niche models (ENM, Peterson 2001) as a proxy for the abiotic environmental requirements of each species and used these niche models to quantify variation among species in habitat suitability across a common landscape, as measured by 'landscape resistance' (see below). Algorithms for generating models from presence-only data result in a continuous prediction of the species occurrence probability for a given set of environmental conditions (Soberón & Peterson 2005; Elith *et al.* 2006; Elith & Leathwick 2009). Raw data for modelling consisted of a set of georeferenced

localities obtained from the Global Biodiversity Information Facility (<http://www.gbif.org/>), HerpNet (<http://www.herpNet.org/>), Instituto de Ciencias Naturales de la Universidad Nacional de Colombia (<http://www.biovirtual.unal.edu.co/ICN/>), the literature (e.g. Duellman 2001) and our own fieldwork. All previously georeferenced localities were visually screened for obvious coordinate errors, and localities without coordinates were georeferenced by the authors. Number of locality records for each species ranged from 15 for *Sachatamia illex* to 283 for *Engystomops pustulosus*, with only four species having fewer than 25 locality records (Table S1, Supporting information).

We generated one ENM for each of 30 species (*P. museosus* was not used, as we could obtain only four locality records) using the maximum entropy algorithm implemented in *Maxent* 3.3.3 using presence and randomly selected background points (Phillips *et al.* 2006; Phillips & Dudík 2008). All models were generated for an area ranging from southern Honduras to northwestern Colombia. In all cases, data set was randomly divided into two subdata sets, a training group containing 80% of the points and a test group containing 20% of the points for model validation. Although more locality points are preferred, Maxent models perform better than other algorithms with as few as five locality points (Hernandez *et al.* 2006; Pearson *et al.* 2007). Climatic data consisted of the 19 bioclimatic variables available in the WORLDCLIM database with a 30 arc-second resolution (Hijmans *et al.* 2005). These variables are derived from three climate parameters, maximum temperature, minimum temperature and rainfall and are highly correlated (Xu & Hutchinson 2010). We used the receiver operating characteristic or area under the curve (AUC, Phillips *et al.* 2006) and the Akaike information criterion (AIC, Akaike 1973), to compare the performance of models based on all 19 variables vs. a reduced 7-variable model. AUC is expected to favour models with more parameters, while AIC penalizes the use of unnecessary or unidentifiable parameters (Warren & Seifert 2011). The 7-variable model was obtained by first performing a two-tailed Spearman correlation analysis between all pairwise combinations of the 19 variables based on their values at 1000 points selected randomly from across the landscape. Variables with a correlation coefficient ≥ 0.75 were considered redundant, and we removed the variable that least likely limited geographic distributions (e.g. removing mean values rather than maxima or minima; Wiens *et al.* 2006). Both the AUC and AIC criteria suggested that the full 19-variable model performed better (Fig. S1, Supporting information); thus, we performed all further analyses on the 19-variable niche models. All 30 resulting ENMs yielded AUC scores above 0.7 and were considered

good predictors of species distributions (Elith *et al.* 2006).

Palaeoclimatic as well as current climatic conditions may predict the observed genetic divergence among conspecific populations. ENMs based on current distributional records can be projected onto spatial hypotheses of environmental conditions during the last glacial maximum (LGM) or other geological time periods (Martínez-Meyer *et al.* 2004; Peterson *et al.* 2004). We therefore projected or 'back-cast' species distribution models to the LGM, 21 000 years before present under two alternative historical scenarios generated using different general atmospheric circulation models (GCM) available in the WORLDCLIM database (Hijmans *et al.* 2005): the Community Climate System Model (CCSM, Collins *et al.* 2006) and the Model for Interdisciplinary Research on Climate (MIROC, K-1-Model-Developers 2004).

Elevation and geographic range

Using georeferenced localities for each species, we extracted elevation data from the shuttle radar topography mission (SRTM) digital elevation model (DEM) with a 90-m resolution (Jarvis *et al.* 2008). We calculated mean elevation and altitudinal range based on the central 90% of observations (i.e. we discarded the extreme upper and lower 5% of elevation records) to minimize the influence of outliers that may potentially represent sink populations atypical of the species in question (Table S3, Supporting information). Geographic range for all 31 species was computed from the species range shapefiles distributed by the Global Amphibian Assessment program (IUCN 2011).

Landscape resistance

When a species confronts a given landscape, the strength of the potential barrier will depend on the species' intrinsic ability to move through the environment and withstand the prevailing abiotic conditions (Janzen 1967; Navas 2002). If an environment is suitable for a given species, then the passage between two points will be easier and, ignoring biotic factors such as potential competitors, demographic connectivity across the landscape should be higher. If abiotic climatic requirements were relevant to explaining variation among species in phylogeographic structure, then we would expect the degree of genetic isolation of conspecific populations to covary inversely with degree of habitat suitability among species.

To measure the effect of the landscape as a barrier to gene flow, one can use an approach based on circuit theory. An ENM assigns to each pixel a probability of

detection of a given species. If we assume that probability of detection is a measure of habitat suitability, then a circuit theory-based approach estimates the effective resistance (or unsuitability) of the landscape to the passage of an organism between two focal regions (McRae 2006; McRae *et al.* 2008). Although habitat unsuitability is not necessarily equivalent to resistance to movement, this is a reasonable assumption for frogs with low dispersal capabilities and a high dependency on their local environment. This measure of resistance takes into account many possible pathways between two localities and can be interpreted as a measure of the effective environmental distance between foci for a given organism (McRae *et al.* 2008). The analysis was performed using the software, CIRCUITSCAPE 3.5.4 (McRae 2006; McRae *et al.* 2008; Shah & McRae 2008), and comparisons were carried out between two pairs of foci: El Copé vs. Brewster and Brewster vs. Cana (Fig. 1; Table S3, Supporting information).

Life-history variables

The ability of organisms to move across a landscape, survive and successfully reproduce may be determined by a variety of life-history traits. In the case of amphibians, cohorts of individuals may need to move long distances while resisting unfavourably dry conditions, to locate potential breeding sites, and maximize the survival of eggs and juvenile stages. We selected three variables available in the literature that potentially influence the ability of amphibians to disperse, survive and reproduce in a marginal environment: reproductive mode, body size and breeding habitat (Table 1). We predicted that species with larval stages may have lower dispersal ability given their higher dependence on water bodies relative to direct-developing frogs and that larger bodied species will have higher dispersal ability than smaller ones (Austin *et al.* 2004; Van Bocxlaer *et al.* 2010). On the other hand, pond breeders may be expected to move among ponds thus having high migration rates relative to terrestrial breeders that may show higher territorial behaviour and thus lower migration (Zeisset & Beebee 2008), although we are not aware of any previous studies that have made an explicit comparison across a common landscape. Breeding habitat and reproductive mode were treated as a single variable, however, since they were tightly correlated within our data set (i.e. all direct developers are terrestrial breeders). In addition to the above life-history variables, we included the biogeographic origin of each lineage (North or South; Table 1), as recent arrivals on the landscape might be expected to show less genetic structure than older lineages (Cody *et al.* 2010). Geological and paleontological reconstructions suggest that northern

Table 1 Summary of nine independent variables used in general linear modelling to predict the dependent variable, average between-population pairwise genetic distance (π_B) in 31 species of frogs present in at least two of the three sampling sites referred to as El Copé, Brewster and Cana. Prior to statistical analyses (Table 2), six variables were transformed to meet assumptions of normality. As reproductive mode and breeding habitat were tightly correlated in our data set (all terrestrial breeders had direct development), we decided to use only reproductive mode

Variable	Type	Units	Transformation
Resistance present	Continuous	Ohm (Ω)	None
Resistance at LGM* (CCSM) [†]	Continuous	Ohm (Ω)	Logarithmic
Resistance at LGM* (MIROC) [†]	Continuous	Ohm (Ω)	Logarithmic
Body size (maximum snout-vent-length)	Continuous	Millimetres (mm)	Logarithmic
Mean elevation	Continuous	Metres (m)	Logarithmic
Elevation range	Continuous	Metres (m)	None
Geographic range	Continuous	Square Kilometres (km ²)	Logarithmic
Reproductive mode	Categorical	Direct/Indirect	None
Origin	Categorical	North/South	None
π_B	Continuous	Average between-population pairwise genetic distance	Square root

*Last glacial maximum at 21 000 years ago.

[†]CCSM and MIROC are two different global circulation models used to generate predictions of past climate (see Methods).

lineages could have been present on the Isthmus of Panama well before southern lineages (Whitmore & Stewart 1965; Kirby & MacFadden 2005).

Ecological and life-history determinants of genetic diversity

We used a set of nine predictive variables and one response variable, with continuous variables transformed to conform to normality assumptions (Table 1). Based on biological criteria, we defined 28 a priori general linear models (GLM) to explain the observed range of genetic divergences among species (Table 2). Models varied in the number and combination of predictor variables and in the presence of interactions between variables. Linear models were evaluated using the software R 2.14.1 (R Core Team 2013). To evaluate which variables were supported by the data and might be more relevant in determining genetic divergence, we performed an AIC model selection procedure among all models (Table 2) using the base R package.

Results

Hierarchical approximate Bayesian computation

Although sample size (n) varied among species and among sites, n did not significantly predict average pairwise distance between sites (π_B , P -value = 0.48, r^2 = 0.011). For pairwise comparisons, El Copé vs. Brewster and Brewster vs. Cana, the posterior probability for the hypothesis of one simultaneous divergence

time ($\Psi = 1$) was nearly 0 in both cases and was thus rejected (π_B ranged from 0.001438 to 0.152888 and from 0.005482 to 0.111562, respectively, for the two comparisons), implying that the intraspecific divergence times of the various species were spread among multiple temporal intervals (Fig. 2). For the comparison of El Copé vs. Brewster, Bayes factor (BF) analysis showed highest support for three divergence time intervals among 30 taxa (BF = 109, Fig. S2a, Supporting information). For the Cana vs. Brewster comparison among 15 taxa, four divergence intervals received the highest support (BF = 3.4, Fig. S2b, Supporting information). To determine the specific taxa grouped in each divergence time interval, we used the BF results to constrain the number of intervals to their most likely values and reran the analysis (Table S4, Supporting information, Hickerson & Meyer 2008). The timing and specific membership of each interval are given in Figs S3 and S4 and Table S4 (Supporting information).

ENMs and landscape resistance

The effective environmental resistance was estimated for each species using the present ENM and two projections to the LGM (Fig. S5, Supporting information, CCSM and MIROC, see Methods). Resistance for both past models is positively and significantly correlated with each other ($R = 0.62$, P -value = 1.04×10^{-5}). Present resistance is positively and significantly correlated with past resistance both with the MIROC ($R = 0.66$, P -value = 1.13×10^{-6}) and the CCSM models ($R = 0.55$, P -value = 1.50×10^{-4}). For those species present at all

Table 2 28 a priori models hypothesized to explain pairwise between-population genetic distance (π_B) of 31 species of frogs across a common landscape, their biological interpretation, and resulting corrected Akaike information criterion (AICc) scores. Models involved combinations of nine independent variables (Table 1). Following the notation in the statistical package, R, a + indicates a linear combination of variables, while \times between two variables indicates the same as + as well as an interaction between terms. Note, in R, AICc values are often negative, with the smallest AICc value still indicating the preferred model. CCSM and MIROC are two spatial models of environmental variation during the last glacial maximum (LGM). See text for details

Model	Biological significance	AICc	Δ AICc
Body size + Present landscape resistance + Geographic range + Origin \times Reproductive mode	Large body size, low present environmental resistance and generalist life history predict low π_B with contributions from biogeographic origin and its interaction with reproductive mode	-111.58	0.00
Present landscape resistance + Mean elevation + Geographic range + Body size + Reproductive mode + Origin	Full model, minus past resistance variable	-105.15	6.43
Present landscape resistance + Landscape resistance at LGM (MIROC) + Mean elevation + Geographic range + Body size + Reproductive mode + Origin	Full model, assuming past environment based on the MIROC model	-103.61	7.96
Present landscape resistance + Landscape resistance at LGM (CCSM) + Mean elevation + Geographic range + Body size + Reproductive mode + Origin	Full model, assuming past environment based on the CCSM model	-101.94	9.64
Origin \times Reproductive mode	Southern frogs show distinctive life histories although there is a potential bias with six of seven northern species being <i>Craugastor</i>	-96.72	14.86
Reproductive mode + Body size	Intrinsic life-history traits alone determine π_B	-91.96	19.62
Reproductive mode	Higher dependence on water may drive species to move less and stay close to water bodies, leading to higher π_B	-91.04	20.53
Origin \times Geographic range \times Body size	Biogeographic factor interacts with range size and body size	-90.00	21.57
Geographic range	Species with wider geographic ranges are habitat generalists that move more and thus have lower π_B	-89.37	22.21
Body size	Large species disperse more and thus should have lower π_B	-89.33	22.25
Reproductive mode \times Body size \times Present landscape resistance	Life-history variables interact with each other and with landscape resistance	-88.65	22.93
Present landscape resistance + Geographic range + Elevation range	Combined habitat generalist traits plus landscape resistance	-87.89	23.69
Body size + Present landscape resistance + Geographic range	Larger species have less environmental restrictions, and habitat generalists have more dispersal capacity	-87.74	23.84
Body size + Present landscape resistance	Larger species have higher dispersal ability, and environmental resistance also plays a role	-87.70	23.87
Origin	Biogeography as sole driver	-87.38	24.20
Present landscape resistance	Present resistance alone explains π_B . The interaction between the organism and its environment predicts genetic isolation	-86.27	25.30
Geographic range + Elevation range	Species with wider geographic ranges and wider elevation ranges will have wider tolerances and thus be able to disperse more	-86.25	25.33
Body size + Present landscape resistance + Elevation range	Larger species have less environmental restrictions and habitat generalists, as measured by elevation range, have more dispersal capacity	-86.16	25.42
Body size + Present landscape resistance + Geographic range + Origin	Larger species have less environmental restrictions, and habitat generalists have more dispersal capacity, while biogeographic origin also plays a role	-85.14	26.44

Table 2 Continued

Model	Biological significance	AICc	Δ AICc
Mean elevation + Present landscape resistance	Elevation and climatic resistance together predict isolation and divergence	-84.49	27.09
Landscape resistance at LGM (CCSM)	Environmental resistance at the LGM (based on CCSM) is sufficient to explain π_B	-84.11	27.47
Present landscape resistance + Landscape resistance at LGM (CCSM)	Present and LGM (CCSM) environments together predict genetic divergence	-84.04	27.53
Present landscape resistance + Landscape resistance at LGM (MIROC)	Present and LGM (MIROC) environments together predict genetic divergence	-83.90	27.68
Landscape resistance at LGM (MIROC)	Environmental resistance at the LGM (based on MIROC) explains π_B	-83.29	28.29
Body size + Present landscape resistance + Geographic range + Mean elevation \times Origin	Large body size, low present environmental resistance and generalist life history predict low π_B with contributions from biogeographic origin and its interaction with elevation	-83.28	28.30
Mean elevation \times Present landscape resistance	Higher elevation species may experience stronger environmental barriers across lowlands	-82.02	29.55
Mean elevation	Elevation alone may explain genetic divergence, although elevation may be a proxy for environmental factors	-81.89	29.69
Mean elevation \times Elevation range	Narrow elevation range predicts larger π_B at high elevations, but not at low elevations	-77.47	34.10

three sites, present and past landscape resistance was generally higher between Cana and Brewster than between El Copé and Brewster (Fig. 3), suggesting the environment in eastern Panama has been more inhospitable for amphibians than central Panama (Fig. 1).

Ecological and life-history determinants of genetic diversity

Our model selection procedure identified a clearly preferred (Δ AICc = 6.43) best-fit model containing five variables: maximum male body size, geographic range, landscape resistance in the present, reproductive mode and biogeographic origin and an interaction term between biogeographic origin and reproductive mode (Table 2). This model explains a large portion of the variation ($R^2 = 0.64$, Adjusted $R^2 = 0.57$). The second best model lacked the interaction term and included mean elevation. Models with fewer variables but that included past climate (MIROC and CCSM), mean elevation or elevational range tended to perform poorly, suggesting that these parameters may be less informative in predicting genetic divergence across the landscape. Body size and reproductive mode tended to occur across the best-performing models (Table 2).

Discussion

Our results show that, indeed, variation in life-history traits may explain variation in phylogeographic

structure among species. HABC analyses show that among species variation in intraspecific divergence could not be explained by a common barrier or by coalescent stochasticity, as, among species, the observed range of genetic divergences between conspecific population pairs was too great (Fig. 2). If the genetic divergence between populations was strictly stochastic in nature, or caused by independent biogeographic histories of species, we would not expect a relationship between ecological variables and genetic divergence. Instead, ecological traits including body size and reproductive mode explain a large portion ($R^2 = 0.64$, adjusted $R^2 = 0.57$) of the genetic variation among our three frog communities. The significant association of genetic divergence and landscape resistance as estimated from niche modelling also suggests that ENM successfully identifies environmental variation important to the evolution of diverse species of amphibians (Glor & Warren 2011). Thus, by effectively holding geography constant and sampling many taxa from the same geographic localities, we have successfully integrated ecological and life-history variables into a model explaining population divergence (Buckley 2009; Johnson *et al.* 2009). Only one variable showed a phylogenetic signal; thus, phylogenetic correlations did not affect the present analysis, but may need to be addressed in future studies. Subsequent studies should include additional life-history traits, such as metabolic rate or desiccation resistance, as such data are not yet available for most species studied here.

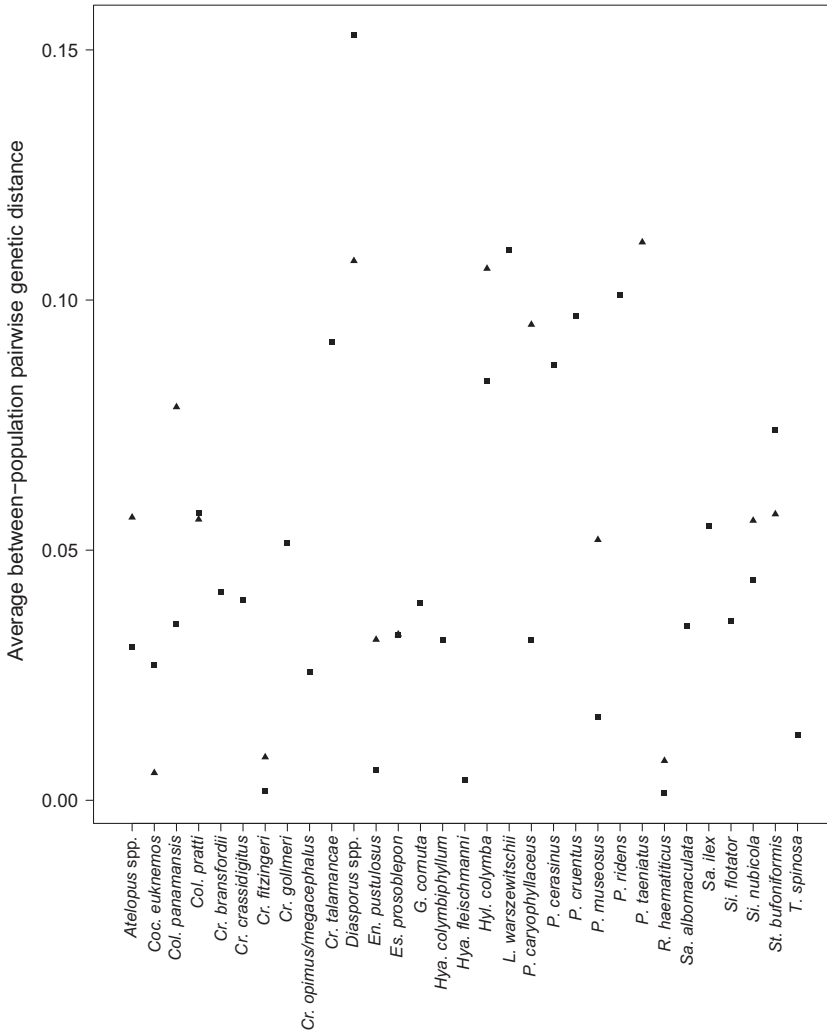


Fig. 2 Average between-population pairwise genetic distance (π_B) computed for each of 31 species of frog based on the concatenated mitochondrial gene sequences, COI and 16S. Data points consist of 30 intraspecific comparisons between El Copé and Brewster (squares) and 15 intraspecific comparisons between Brewster vs. Cana (triangles). Complete generic names and numeric π_B estimates are given in Table S3 (Supporting information).

Our hope is that this study of Panamanian frogs may provide a general approach of wide applicability, as obvious geographic features have failed to constitute a general explanation for observed divergences among allopatric populations, especially in the tropics (Antonelli *et al.* 2010). Vicariant obstacles such as mountains or rivers certainly divide many phylogeographic clades and sister species (Avice *et al.* 1987; Ribas *et al.* 2012), yet there may be a great many pairs of allopatric clades for which there are no apparent barriers separating them. Furthermore, many physical barriers such as mountains do not impede gene flow directly, rather they promote environmental and habitat gradients that may be inhospitable to the dispersing organism (Janzen 1967). The dichotomy between physical and environmental barriers may be a false one, at least in some cases. Recent research has focused not only on the type of barrier but also on the type of mechanism impeding gene flow, that is isolation by distance (IBD) or by physical impassibility vs. isolation by environment

(IBE) or ‘immigrant unviability,’ in which natural selection acts against migrating individuals (Nosil *et al.* 2005; Bradburd *et al.* 2013; Wang 2013). The IBE model highlights why codistributed species may vary widely in their phylogeographic structure, while our comparative approach using general linear modelling allows one to evaluate specific ecological or physiological traits that may contribute to divergence.

Our approach to investigating the effect of ecology on phylogeography, which we refer to as ‘ecophylogeography’, follows Pabijan *et al.* (2012), with some modifications. We employed an HABC analysis to first determine whether the variation in mtDNA divergence cannot be explained simply by coalescent stochasticity in the context of isolation by distance or isolation by physical barrier (Hickerson *et al.* 2007). In addition to life-history variables, we incorporated environmental heterogeneity (IBE) as an explanatory variable through ENM and ‘isolation by resistance’ analyses (McRae 2006). Our data set also covered a larger phylogenetic

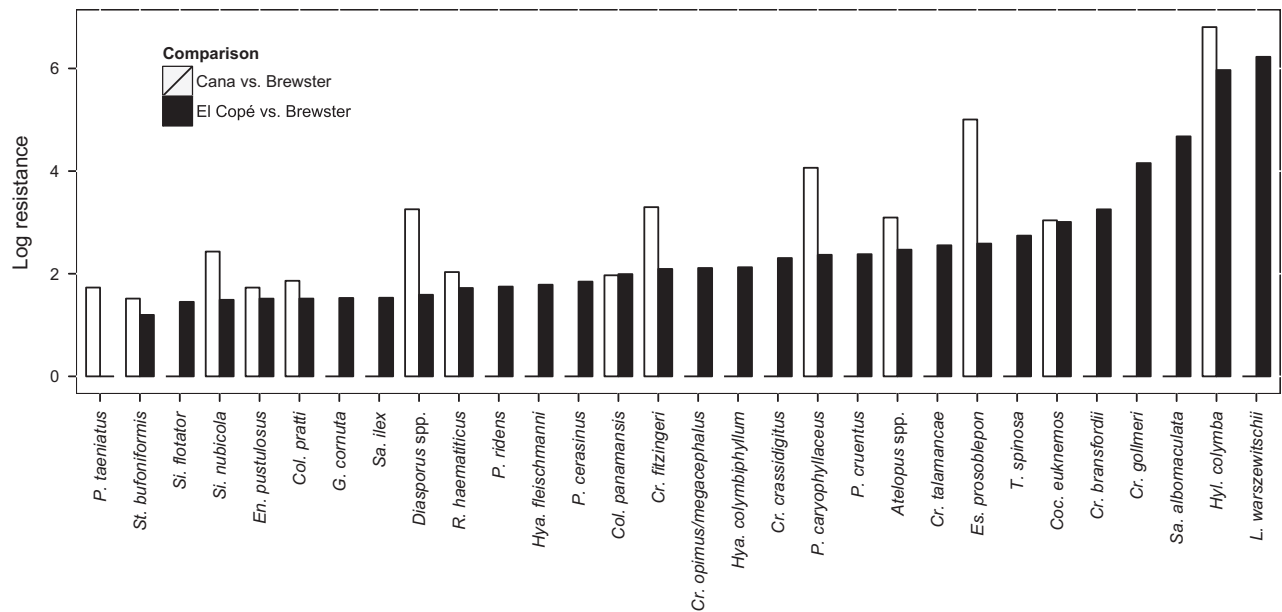


Fig. 3 Landscape resistance in log (ohms) for each of 30 frog species as measured between El Copé and Brewster (black bars), and between Cana and Brewster (white bars), based on present climatic conditions. Resistance was obtained from environmental niche models (see Methods) using the software CIRCUITSCAPE 3.5.4. Resistance could not be estimated for *Pristimantis museosus* as only four locality records were available for this species.

diversity, including species from several taxonomic families, which may explain why we recovered a greater number of significant explanatory variables in addition to body size (Pabijan *et al.* 2012; Wollenberg Valero 2015).

Our approach does not supplant other ecophylogeographic inference methods but does offer some advantages. While landscape genetic studies frequently uncover an association between population genetic structure and environmental heterogeneity (Storfer *et al.* 2007; Richards-Zawacki 2009), our comparative phylogeographic approach allows one to search for common patterns across species and quantify to what extent phylogeographic patterns may be predictable from a knowledge of organismal biology. Previous studies have shown the global importance of dispersal potential in explaining genetic divergence particularly for marine and freshwater organisms as well as some terrestrial invertebrates (Bohonak 1999); however, the importance of continuous variation in specific ecological and life-history traits has not been evaluated. Using a phylogeographic approach, we can infer the action of evolutionary processes acting over longer time scales than those revealed by landscape genetics (Wang 2010; Lawson 2013). By including explicit life-history characteristics in a general linear modelling exercise, our approach evaluated the contribution of specific organismal traits to genetic divergence (Pabijan *et al.* 2012) coupled with a contribution of IBE (Wang 2013).

We estimated phylogeographic structure as genetic divergence at mitochondrial DNA (mtDNA). While variation in mtDNA may not be representative of variation in the nuclear genome (Fay & Wu 1999; Bazin *et al.* 2006; Toews & Brelsford 2012), this standardized marker is widely used in comparative phylogeographic studies (Carnaval *et al.* 2009; Wang *et al.* 2013) due to its particular evolutionary genetic properties. High mutation rates of anuran mtDNA allow one to estimate recent divergence times among species (Crawford 2003). Smaller effective population sizes (N_e) permit a rapid achievement of reciprocal monophyly between populations (Hudson & Coyne 2002), as observed here. The HABC approach takes into account the stochasticity inherent in single-locus coalescence, and moreover, estimation of population divergence time $E(\tau)$ does not improve appreciably until eight nuclear loci are included (Huang *et al.* 2011). While mtDNA data sets are still the norm for comparative ecophylogeographic analyses (Pabijan *et al.* 2012; Wang *et al.* 2013), genomic-scale phylogeographic data sets may soon be routinely available (McCormack *et al.* 2013).

Although ecology was a main predictor of genetic divergence, our model also included an effect of biogeographic origin of species. Species of northern origin showed significantly less genetic divergence across the same landscape than southern lineages, which is the opposite of what we predicted given that southern lineages would be more recent arrivals and would

therefore have had less time to build up spatial genetic structure (Weir *et al.* 2009). Alternatively, many southern lineages likely arrived in Panama perhaps across an early Isthmian connection (Montes *et al.* 2012) around 8 Ma (Pinto-Sánchez *et al.* 2012), which may have been sufficient time to erase the potential genetic imprint of recent colonization.

We found that species ranges predicted genetic divergence, thus revealing that species with wider geographic ranges have less genetic divergence between populations (Duminil *et al.* 2007). Wider geographic ranges are indicative of habitat generalists that, by definition, can tolerate a wider range of environments and should experience lower IBE and increased dispersal. Although this pattern was not detected in Malagasy frogs (Pabijan *et al.* 2012), perhaps because that study covered less phylogenetic diversity, this relationship should be relatively straightforward to test in other species.

The relevance of life-history variables in explaining phylogeographic structure has not been widely tested. In plants, life-history traits such as breeding system, life form and seed mass are known to influence population structure (Hamrick & Godt 1996; Duminil *et al.* 2007). For marine invertebrates, developmental mode is thought to be important in limiting dispersal and promoting population structure (Arndt & Smith 1998; Collin 2001). Habitat preference in birds and more generally their ability to move through the landscape has been shown to predict genetic differentiation and eventually speciation (Burney & Brumfield 2009; Smith *et al.* 2014).

We found two life-history characteristics of anurans that are important in predicting genetic divergence. Larger animals showed less genetic divergence among populations, implying they have higher dispersal capacity than smaller species (Pabijan *et al.* 2012; Wollenberg Valero 2015). Larger frogs may be better dispersers simply because of scale (they can move faster than smaller frogs) or may be better survivors, for example better able to resist desiccation. Reproductive mode was also a significant predictor of genetic divergence, but the effect was in the opposite direction. We predicted that direct developers, being independent of bodies of water for reproduction, would have greater dispersal capabilities, yet it was the toads and tree frogs with larval stages, which showed less genetic divergence and, by implication, greater dispersal potential. Higher phylogeographic structure could be due to low dispersal associated with site faithfulness and territorial behaviour in terrestrial frogs (Stewart & Rand 1991; Ryan *et al.* 2008; Fouquet *et al.* 2012). If true, then behavioural ecology may be another variable that could be taken into account in ecophylogeographic studies, although this may be

difficult using the comparative approach advocated here. Alternatively, the direct developers may simply show greater vulnerability to desiccation and greater niche conservatism, even though they do not need bodies of water to reproduce.

Using our comparative multivariate approach, we were able to study both intrinsic (life history) and extrinsic factors (environmental resistance obtained from ENMs). We recognize, however, that IBE can be interpreted as the interaction between an organism's intrinsic characteristics and the abiotic landscape it confronts. Because we looked at variation in genetic divergence across a common landscape, in effect, we held the landscape constant to reveal variation among species. Thus, we interpret the landscape resistance variable more as a composite property of the organism. The differing responses of organisms to a common landscape has been suggested by a few studies using pairs of species (Goldberg & Waits 2010; Richardson 2012), but rarely is such a high number of multiple codistributed species synthesized in one analysis. To successfully delimit intrinsic vs. extrinsic factors and quantify their relative roles in determining how populations interact with landscape characteristics, biologists may need to incorporate direct studies of organismal physiology, ecomorphology and even behaviour (Janzen 1967; Navas 2002; Bernardo *et al.* 2007). Global DNA barcoding initiatives have begun to provide genetic data on whole communities and regional biotas (Kerr *et al.* 2007; Dick & Kress 2009; Murphy *et al.* 2013). Such massive data sets should provide opportunities to implement the present comparative approach over much larger regions and more diverse species. These data will lend themselves naturally to a phylogeographic approach (Hajibabaei *et al.* 2007). Looking beyond the ENM-based approach (Richards *et al.* 2007), comparative phylogeography should benefit greatly from the direct integration of organismal biology, which may interface with new approaches to studying adaptive genetic variation in a phylogeographic context (Hickerson *et al.* 2010; Günther & Coop 2013).

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A.P. and A.J.C. designed the study. R.I., K.R.L. and A.J.C. collected the data. A.P. and A.J.C. analysed the data and wrote the manuscript with vital input from R.I. and K.R.L.

Data accessibility

DNA sequences: GenBank accession nos. for new sequences of 16S are KR863128–KR863371 and KR911916–KR911918, and for new sequences of COI are: KR862873–KR863113 and KR911913–KR911915. GenBank accession nos. for previously published sequences are provided in Table S2 (Supporting information). Localities and climatic variables used for constructing environmental niche models, DNA sequence alignments per species per locality, R code for generating GLMs and database used in all analyses are available in the Dryad repository (DOI: 10.5061/dryad.t430k).

Supporting information

Additional supporting information may be found in the online version of this article.

Table S1 Number of georeferenced localities used for environmental niche modeling for each species (training, testing and total) and number of samples per species of frog per sampling site in Panama (Cana, El Copé and Brewster) for genetic analyses.

Table S2 Museum voucher number (absent for specimens awaiting accession), field collector number, barcode of life data systems (BoLD).

Table S3 Database of species trait values for 31 frog species.

Table S4 Divergence intervals and times estimated from the HABC analysis.

Fig. S1 Evaluating performance of alternative niche models.

Fig. S2 Number of divergence intervals from the HABC analysis.

Fig. S3 Number of species at each divergence interval for the comparison between El Copé and Brewster obtained from the HABC analysis.

Fig. S4 Number of species at each divergence interval for the comparison between Cana and Brewster obtained from the HABC analysis.

Fig. S5 Ecological niche models constructed for 30 species of frogs.