One of the central goals of evolutionary biology is to understand why species present certain biogeographic distribution patterns (Cox et al. 1973; Schall and Pianka 1978; Cracraft 1983). Biogeographic studies highlight the role of paleogeography and paleoclimate in shaping the current distribution of diversity (Raven and Axelrod 1972; Cox 1974; Gamble et al. 2011; Giribet et al. 2012; Beaulieu et al. 2013). However, geological and environmental processes may drive not only specific biogeographic histories but also diversification dynamics (Flagstad et al. 2001; Drovetts 2003; Kozak and Wiersma 2006; Ricklefs 2006; Fjeldså et al. 2012; Toussaint et al. 2014; Badgley et al. 2017; Oliveros et al. 2019). In this context, inferring the underlying causes of biogeographic patterns can be challenging, since the incidence of certain biogeographic processes (e.g., dispersal or vicariance) is not only subjected to historical factors, but also to changes in the number of species through time (Sanmartín et al. 2001; Sanmartín and Ronquist 2004; Jiang et al. 2019).

Some systems on Earth are particularly interesting from a biogeographic perspective due to the nature of their shifting climatic or geologic histories. Such is the case of biotic interchanges, in which tectonic and paleoenvironmental processes have had an impact on the distribution of biodiversity (Marshall et al. 1982; Barry et al. 1991; Vermeij et al. 1991a,b; Bibi 2011). The region of Afro-Arabia and its biota constitute an ideal system to investigate in general biogeographic patterns and their relationship with past environmental conditions and diversification trends. That is, the geologic history of this region is characterized by a complex succession of landmass fragmentation and reconnection events beginning during the Eocene with the split of the Arabian and African plates around 40–30 Ma, which eventually triggered the onset of the Red Sea formation around 25 Ma (Bohannon 1986; Ghebreab 1998; Redfield et al. 2003; Bosworth et al. 2015; Scotese 2014a,b). The recurrent formation of dispersal barriers and land bridges, together with constantly shifting climatic conditions (Flower and Kennett 1994; deMenocal 1995), has potentially left an imprint on the evolutionary history of the biodiversity inhabiting the region (Delany 1989; Winney et al. 2004; Šmíd et al. 2013; Metalinou et al. 2015; Tamar et al. 2016a, 2018; Simó-Ruidalbas et al. 2019). The desert conditions in the Afro-Arabian region contributed to the extensive diversification of arid-adapted taxa, such as reptiles (Brito et al. 2014;
making them particularly suitable for studying large-scale biogeographic patterns in this area. However, a comprehensive biogeographic assessment of Afro-Arabian reptiles has yet to be conducted.

Here, we conduct an integrative study of the biogeography and macroevolutionary dynamics of squamates in the Afro-Arabian region. We use statistical models to develop an expectation for the incidence of biogeographic events through time and compare it with observed biogeographic patterns. By definition, all models are simplifications of reality and it is naive to expect any statistical model to fully capture all evidence of biogeographical or diversification events provided by empirical phylogenies. Consequently, we use simulations to exploit model inadequacy, pinpointing time-periods during which the observed evolutionary history of squamates significantly deviates from the expectation based on simplistic models. Such periods might be the result of external forces (e.g., paleogeographic and paleoclimatic histories or ecological interactions) affecting the particular biogeographic history of these animals beyond the origin and extinction of lineages.

We found that although a large portion of the observed biogeographic events can be explained by the fitted model and number of lineages through time, some time-periods were more likely shaped by processes other than diversification dynamics alone, being characterized by model inadequacy. We also show that we may unravel how biogeographic history relates to diversification trends by comparing biogeographic expectations to empirical observations and how both are associated with other historical or ecological processes.

**Materials and Methods**

**Taxon Sampling and Definition of Groups of Interest**

In order to investigate the history of biogeographic events involving Africa and Arabia, we first identified all genera within the order Squamata that are currently present in both areas, and then sampled all species in those genera following the most updated taxonomy (Uetz et al. 2020), including species currently present in both areas, and then sampled all genera within the order Squamata that are also known to contain species in those genera following the most updated taxonomy. However, we only had two species, precluding any biogeographic reconstruction. We further sampled at least one species representing each family of Squamata and additional species to serve as outgroups and aid in divergence time calibrations, including the sister species to Squamata, the tuatara (*Sphenodon punctatus*), that was used as outgroup. In total, we sampled 555 species: 430 species belonging to the clades of interest and included in the subsequent biogeographic analyses, and 125 additional species which served for inferring a taxonomically balanced phylogeny and to inform calibrations.

**DNA Amplification, Phylogenetic Reconstruction, and Divergence Times Estimation**

Out of the 2612 DNA sequences used for the phylogenetic analyses, 2490 were retrieved from GenBank, and 122 were produced for this study (Supplementary Table S1 available on Dryad at http://dx.doi.org/10.5061/dryad.wpzgmsbk). We included five mitochondrial (12S, 16S; cytochrome b, cya; and NADH dehydrogenases 2 and 4, ND2 and ND4) and six nuclear markers (the acetylcholinergic receptor M4, ACM4; the oocyte maturation factor MOS, C-MOS; the melanocortin 1 receptor, MC1R; neurotrophin-3, NT3; and the recombination activating genes 1 and 2, RAG1 and RAG2). To generate the new sequences, genomic DNA was extracted from ethanol-preserved tissue samples using the SpeedTissue Tissue DNA Extraction kit (Biotools, Madrid, Spain). Primers and PCR conditions used for the amplification of all genetic fragments are summarized in Supplementary Table S2 available on Dryad. We checked the chromatographs and assembled the contigs in Geneious v.7.1.9 (Kearse et al. 2012). Heterozygous positions in the nuclear genes were coded according to the IUPAC ambiguity codes. We used a supermatrix approach, concatenating all the genetic markers for the 555 samples. The sequence alignments were produced with the online software MAFFT (Katoh et al. 2018) and the concatenated data set had a length of 12,049 base pairs. We reconstructed the phylogenetic relationships and estimated the divergence times in a single Bayesian inference analysis using the software BEAST2 (Bouckaert et al. 2019). We used 21 calibration points (fossils, biogeographic events, and secondary calibrations), avoiding fixing nodes of potential interest for the biogeographic discussion. Information on calibration points is available in Supplementary Table S3 available on Dryad. Additionally, we constrained higher-level clades to

**Naja, Pseumolphis, and Telescopus.** We sampled all the available species from each of these groups except for *Hemidactylus*, for which we considered the “arid clade” only (Carranza and Arnold 2006) as a group of interest for the purpose of this study. This clade is defined by the node including *H. laevis* and *H. granosus* in our phylogenetic reconstruction, and comprises 53 described species. We excluded the Afro-Arabian snake genus *Lystrophus* from the biogeographic analysis because we only had two species, precluding any biogeographic reconstruction. We further sampled at least one species representing each family of Squamata and additional species to serve as outgroups and aid in divergence time calibrations, including the sister species to Squamata, the tuatara (*Sphenodon punctatus*), that was used as outgroup. In total, we sampled 555 species: 430 species belonging to the clades of interest and included in the subsequent biogeographic analyses, and 125 additional species which served for inferring a taxonomically balanced phylogeny and to inform calibrations.
match one of the most recent and supported topologies of squamates published (Streicher and Wiens 2017; see Supplementary Fig. S1 available on Dryad). We tried different partition schemes, including those resulting from preliminary analyses with PartitionFinder (Lanfear et al. 2016) and ModelFinder (Kalyaanamoorthy et al. 2017) as implemented in IQ-TREE (Nguyen et al. 2015), as well as separating the mitochondrial and the nuclear genes in two different partitions. But all of them failed to converge in the BEAST2 analyses. Thus, we performed five individual runs of 100 million generations each, sampling every 10,000 generations, with a Yule process tree prior, a single partition, and a relaxed lognormal clock model to account for heterogeneity in rates across the phylogenetic tree. We used the reversible-jump (RB) method of Bouckaert et al. (2013) to sample across multiple substitution models and accommodated the among-site rate heterogeneity using a discretized Gamma distribution with four categories. We used Tracer v.1.7 (Rambaut et al. 2018) to evaluate the stationarity of the chains and check for sufficient effective sample sizes (ESS > 200), as well as convergence between independent runs. We combined the posterior distributions with LogCombiner, discarding 30% of the posterior trees as burn-in. We obtained the maximum clade credibility tree with median heights using TreeAnnotator (both softwares provided within the BEAST2 package).

Biogeographic Area Delimitation and Ancestral Range Estimation

We used the R package BioGeoBEARS (Matzke 2013) to reconstruct ancestral ranges for each of the 21 monophyletic groups of interest separately, excluding outgroups (see Taxon Sampling and Definition of Groups of Interest section). We assigned each species to their corresponding main biogeographic ranges as a discrete character, delimiting five different areas (Fig. 1): Africa (all the African continent west of the Sinai Peninsula, including Madagascar), Arabia (the Arabian Peninsula south of Jordan and Iraq, including the Socotra Archipelago and Kuwait), Eastern Mediterranean (Israel, Lebanon, Syria and Turkey), Asia/Oceania (east of Iraq), and Europe. This delimitation of biogeographic areas was based on the patterns of squamate distribution in the region (Roll et al. 2017; Šmíd et al. 2021b), and was independently assessed using the interactive online tool for delimiting biogeographical regions from species distribution InfoMap Bioregions with the following settings: cluster cost = 1.0, number of trials = 10 (Edler et al. 2017). The maximum number of areas per node was set to two. We performed ancestral reconstructions using the following models in BioGeoBEARS: Dispersal-Extinction-Cladogenesis (DEC; Ree and Smith 2008), DIVALIKE (Ronquist 1997) and BAYAREA (Landis et al. 2013). The $j$ parameter representing founder event speciation (Matzke 2014) was excluded from each model following recommendations of attending to empirical considerations in the application of statistical models (Ree and Sanmartín 2018; Klaus and Matzke 2020). The models we tested represent alternative biogeographic histories resulting from vicariance, following decades of prior research in the region (e.g., Pook et al. 2009; Šmíd et al. 2013; Portik and Papenfuss 2012), which have never considered a scenario of founder event speciation to be plausible. Fitted models were subsequently selected according to the Akaike information criterion, correcting for small sample size (AICc; Akaike 1975). We extracted ancestral states information at the nodes with the BioGeoBEARS functions get_ML_probs and get_ML_states_from_relprobs. For each node, we selected the state with the highest marginal probability.

Biogeographical Assessment and Incidence of Biogeographic Events Through Time

We tracked the counts of the following types of biogeographic events through time: (i) dispersal: parent node present in Africa or Arabia, and one of the descendant nodes present in the other area (Africa or Arabia, respectively); (ii) vicariance: parent node widespread (Africa+Arabia) and each of the descendant nodes in a different area; and (iii) extirpation or range contraction: parent node widespread and one descendant node only present in Africa or Arabia. As pointed out by Klaus et al. (2016), accounting for uncertainty in phylogenetic calibration of divergence times is essential for the interpretation of biogeographical reconstructions. Therefore, we adopted their approach and included a time range for each biogeographic event, rather than a single time point. We then used the overlap of intervals to estimate the incidence of biogeographic events between Africa and Arabia through time.

For vicariance events, we accounted for temporal uncertainty by including the 95% highest posterior density (HPD) interval of each node inferred to have undergone vicariance between Africa and Arabia. For dispersal and range contraction events, we used time ranges defined as the length of the branch during which the dispersal took place according to our ancestral reconstruction. As a more conservative procedure, we carried out a set of analyses where temporal uncertainty in dispersal and range contraction events was defined as the branch length plus the 95% HPD of the nodes at the beginning and the end of the branch. Combining all clades, we obtained the biogeographic incidence through time by summing the overlapping biogeographic events in each million-year interval. Our metric is analogous to the Maximal number of observed Dispersal Events per million year utilized by Klaus et al. (2016) but includes vicariance and extirpation events besides dispersal.

Model Adequacy and Deviations from the DEC Model

We sought to quantify the extent to which the observed biogeographic history deviates from that expected from
the best-fit model and the inferred phylogenetic history alone. That is, we aimed to identify whether (and when) forces beyond those modeled within the DEC framework led to deviations from model expectations. To do so, we simulated the biogeographic history for each of our clades of interest along their respective phylogenies to obtain an expected distribution of biogeographic events through time. We simulated 1,000 per-clade biogeographic histories, running the function simulate_biegeo_history in BioGeoBEARS, using the phylogenetic trees, states at the root, empirical transition matrices, and best-fit biogeographic models from the reconstruction made for each clade of interest. As we did for the observed biogeographic history, we overlapped the temporal uncertainty of biogeographic events in the simulated biogeographic histories and built 1,000 lines of biogeographic incidence through time by summing the number of events in each million-year period. Then we generated a distribution of the expected number of events through time by calculating the 95% confidence interval of the number of simulated biogeographic events per million-year interval. By comparing the number of observed events through time to those expected under the inferred model, we identified specific time periods when the observed number of biogeographic events significantly differed from that expected under the given model and phylogeny alone. We conducted this comparison for all kinds of biogeographic events together and also separately for dispersal in each direction, extirpation from each area (Africa and Arabia), and vicariance. This allowed us to explore and identify time periods during which biogeographic events deviated significantly from model expectations, implying a role of forces external to that modeled by the DEC framework. Figures were produced in the R environment (R Core Team 2019), with the packages ggtree (Yu et al. 2017), treeio (Wang et al. 2020), and the collection of packages within tidyverse (Wickham et al. 2019). The data and scripts produced for the biogeographic analyses can be found in the Supplementary Material available on Dryad.

RESULTS

Phylogenetic Reconstruction and Divergence Times Estimation

The sequence data matrix was 43% complete, with a minimum of 22% (NT3) and a maximum of 68% (12S). The dated phylogenetic tree of all sampled taxa is shown in Figure 1 (see Supplementary Fig. S1 available on Dryad for details on species names and
calibration points). The topology within our clades of interest and crown age estimates do not deviate substantially from previous results (Supplementary Table S4 available on Dryad). Importantly, our approach allows us to include calibration uncertainty in the analyses, a critical step when addressing biogeographic questions (see Biogeographical Assessment and Incidence of Biogeographic Events Through Time in the Methods section). Furthermore, including all species in the same phylogenetic analysis means that all ages are comparable across clades.

Ancestral Range Estimation and Biogeographical Assessment

The biogeographic histories for each clade are summarized in Supplementary Appendix 1, showing the areas reconstructed for each node according to the most supported model (Supplementary Table S5 available on Dryad). We did not find essential differences between the results of biogeographic incidence through time from both approaches we applied (i.e., one where temporal uncertainty of dispersal and extirpation events was defined as the length of the branch, and a more conservative one where it included also the node uncertainty at the beginning and the end of the branch), and hence we present here the results from the former. Figure 2 shows, for each clade separately, estimated time ranges of biogeographic events involving Africa and Arabia and the type of event (dispersal from Africa to Arabia, dispersal from Arabia to Africa, vicariance, extirpation from Africa, or extirpation from Arabia; see Methods section). Across all clades, the ancestral reconstructions yielded 20 dispersal events from Africa to Arabia and 14 from Arabia to Africa, 26 vicariance events, 8 extirpations from Africa, and 6 extirpations from Arabia (Fig. 2). The probability of biogeographic events began to rise between 50 and 40 Ma within some clades (Fig. 2). The probability distribution of biogeographic events through time, considering only the shape of the phylogeny, the number of lineages through time, and the best-fit biogeographic model. This is shown in Figure 3 (see Supplementary Fig. S2 available on Dryad for the results from the conservative approach, where dispersal and range contraction events included the node uncertainty at the beginning and the end of the branch), including all types of events together (Fig. 3a) and each type separately (Fig. 3b–f). Any deviation of the observed number of events from the expected distribution indicates the inadequacy of the DEC model to fully capture the temporal dynamics of biogeographic events.

When summing across all types of biogeographic events, we find that the observed number of events does not differ substantially from the model expectation, with the exception of the period between 23 and 15 Ma, during which we observed slightly more incidence than expected (Fig. 3a). In contrast, looking at each type of biogeographic event separately gives a much more nuanced understanding. Most notably, the incidence of vicariance events was considerably higher than expected beginning around 30 Ma until about 3 Ma (Fig. 3a), a likely explanation for the excess of biogeographic events observed around the same time (Fig. 2).

The number of observed events of dispersal (both from Arabia to Africa and from Africa to Arabia; Fig. 3b and e, respectively) and range contraction (Fig. 3c and f) have not been significantly different than expected. The levels of dispersal in both directions suggest a potential asymmetry of dispersal rates and exchange among the two regions. Indeed, the interchange between Africa and Arabia has not been symmetrical (Fig. 4). Dispersal started earlier from Africa to Arabia (until around 32 Ma all dispersal happened in that direction). Later, through the Oligocene and the Early Miocene, the relative rate of dispersal from Arabia to Africa intensified, although it never matched the levels of dispersal from Africa to Arabia, which have always been predominant. This asymmetry, however, is indistinguishable from that expected given the phylogeny and fitted biogeographic model.

Discussion

Over the last few decades, historical biogeography as a field has experienced major theoretical and methodological advances, resulting in an increasing number of studies that discuss distributional patterns in the context of past environmental processes according to ancestral range estimations. Recent biogeographic assessments have recognized the need for controlling the effect of phylogenetic diversification through time by randomly sampling nodes from the phylogeny after biogeographic reconstruction (Jiang et al. 2019). Our approach goes one step further and allows us to identify particular periods in the past when biogeographic events happened more or less frequently than predicted by model expectations. That is, we exploit model inadequacies to learn more than by taking fitted models at face value alone. This approach enhances our ability to ask questions central to the study of biogeography, such as what are the historical causes of exceptional historical biogeographic patterns observed in the data.
and how have those patterns unfolded into current species geographic distributions?

Our results show a substantial contrast between the number of reptile biogeographic events involving Africa and Arabia observed in the last 40 myr and the expected number of such events simulated under the best-fit models for the same period. The excess in the incidence of vicariance events between Africa and Arabia during the Oligocene and the Miocene (approximately from 30 to 5 Ma; Fig. 3d) is of particular interest. We hypothesize this pattern is driven by the movement of the Arabian and the African plates (Fig. 4). These two plates, once connected in a single landmass, started splitting around 40–30 Ma (Bosworth et al. 2005; Scotese 2014a), initially with the uplifting of mountain chains along the plates’ limits and eventually resulting in the establishment of the Red Sea around 25 Ma (Ghebreab 1998). This fragmentation is considered to be a major driver of the biogeographic history of taxa currently present in the Afro-Arabian area (Pook et al. 2009; Metallinou et al. 2012; Šmíd et al. 2013; Portik and Papenfuss 2015; Tamar et al. 2016a, 2018). Furthermore, the aridification of the region, that began around the middle Miocene (Flower and Kennett 1994), has been suggested to have

![Biogeographic events through time](image-url)
a significant role in promoting vicariant speciation (Metallinou et al. 2015). Our results suggest this interval of tectonic and climatic instability had a crucial role in shaping the observed biogeographic patterns of Afro-Arabian reptiles, with many clades undergoing vicariance as well as dispersing out of their place of origin from the late Eocene through the middle Miocene (Fig. 2). Consequently, the observed increase in vicariance events beyond DEC model expectations from 30 to 10 Ma (Fig. 3d) is likely related to these geologic and environmental processes involving Africa and Arabia, which is also widely supported in the literature (Smid et al. 2013; Tamar et al. 2016a).

The fact that range contraction, and especially dispersal patterns, do not differ from biogeographic expectations (Fig. 3b, c, d, and f) is also worth discussing, given the complex paleogeographic and climatic history of the region. The fragmentation of the Afro-Arabian plate, which culminated at the end of the Oligocene, would lead us to hypothesize a reduction in the observed rate of dispersal from that time onwards. Likewise, the climatic fluctuations that could have imposed environmental barriers between Africa and Arabia since the Miocene (Tchernov 1992; Pickford and Morales 1994) precluding dispersal and promoting range contraction on the regional fauna, would make us expect a reduced incidence of these biogeographic events compared to the simulated history. These results suggest that dispersal and extirpation patterns might have been shaped by the interplay of diversification dynamics and external forces. Namely, the progressive aridification during the Miocene and the Plio-Pleistocene climatic oscillations have been suggested to trigger diversification in Afro-Arabian faunas (Fu 2000; Douady et al. 2003; Guillaumet et al. 2008; Brito et al. 2014; Tamar et al. 2016a). This, together with the vicariant speciation promoted by the Afro-Arabian plate fragmentation, may have driven the observed dispersal and extirpation patterns. Nevertheless, these results may also reveal the importance of plate tectonics in the region following the formation of the Red Sea. While the expansion of the Red Sea was acting as a potential vicariant agent, the landmasses came into contact several times through different time periods. The connection of
FIGURE 4. Interchange asymmetry between Africa and Arabia over time. Green shaded areas represent, in percentage, the relative intensity of observed dispersal from Africa to Arabia (up, in light green) and from Arabia to Africa (bottom, in dark green). The dark green line shows the observed proportion of dispersal from Arabia to Africa. The darker shaded area in the background encompasses the 95% interval of the expected distribution of dispersal proportion from Arabia to Africa over time according to the 1000 biogeographic simulations. Observed asymmetry meets expectations where the green line falls within the 95% of the expected asymmetry (i.e., throughout the entire time period shown). Paleogeographic maps adapted from Bosworth et al. (2005).

Eurasia and Afro-Arabia through the Gomphotherium landbridge around 18 Ma and the later collision of the Arabian plate with Eurasia around 15 Ma (Rögl 1999; Jolivet and Faccenna 2000; Bosworth et al. 2005) have played an important role as a potential dispersal route in the Miocene faunal interchange (Barry et al. 1991; Joger 1991; Nakaya 1994; Kappelman et al. 2003; Harzhauser et al. 2007; Tassy 1999). This dispersal route has been permanent since the middle Miocene, connecting Africa and Arabia through the Sinai Peninsula. Likewise, the global decline in sea level during the late Miocene (Flower and Kennett 1994) resulted in a temporary southwestern landbridge between the Arabian Peninsula and the Horn of Africa that was in place from approximately 10 to 5.3 Ma (Bosworth et al. 2005). It has been suggested that this land connection might have served as a dispersal route from Africa to Arabia and vice versa (Tassy 1999; Portik and Papenfuss 2012; Smid et al. 2013) until the reopening of the Bab el Mandeb Strait during the Pliocene, which has persisted until the present. Furthermore, aridification trends may have contributed to the dispersal of reptile taxa (Tamar et al. 2018). Here, we found evidence that the landbridge that emerged as a consequence of the connection between the African, Arabian, and Eurasian plates served as a permanent dispersal corridor through the entire geologic history after the Red Sea opening, suggesting an environmental homogeneity between the Sahara and the Arabian deserts. This is consistent with the lack of difference between observed and expected patterns, both for dispersal and range contraction events.

Another interesting aspect of our results is that the reptile interchange between Africa and Arabia shows a clear asymmetry (Fig. 4). The predominant direction of the interchange, from Africa to Arabia, suggests a scenario of classic island biogeography source-sink dynamics, where Africa has primarily acted as the “mainland” source of colonizing species into Arabia. After the establishment of the Red Sea at the end of the Oligocene, the relative levels of dispersal in both directions have remained stable, which further highlights the role of the dispersal corridor in shaping biogeographic patterns and the environmental homogeneity on both sides of the Red Sea.

Overall, our study corroborates the importance of the unstable geologic and climatic history of Afro-Arabia as a key factor to explain the evolutionary history of the fauna in this region. Our results suggest distinct roles of tectonic and climatic processes in different periods since the Oligocene. The fragmentation of the Afro-Arabian plate seems to be the main driver of biogeographic patterns through the Oligocene and
Miocene. Additionally, the landmass movement after the fragmentation, together with diversification dynamics driven by climatic fluctuations, appear to be more relevant in explaining the biogeographic patterns from the middle Miocene onwards.

The influence of this region’s history on evolutionary dynamics is not limited to squamates. Analogous biogeographic patterns have been observed in other animal groups. For example, the evolutionary history of Arabian toads also shows the imprint of ancient vicariance and later dispersal between Africa and Arabia (Portik and Papenfuss 2015). The tectonic events that resulted in the connection of Africa, Arabia and Eurasia shaped dispersal patterns in rodents, artiodactyls, and primates (Barry et al. 1991; Nakaya 1994; Fleagle and Gilbert 2006; Bibi 2011), and the southern connection between Africa and Arabia served as a dispersal route for baboons (Winney et al. 2004). Environmental trends resulting in the aridification of the region have also likely influenced the evolutionary patterns in some mammals (Douady et al. 2003) and birds (Guillaumet et al. 2008). By contrast, plant biogeography has been suggested to be strongly influenced by long-distance dispersal (Clayton et al. 2009) and therefore might be relatively independent from regional dynamics. However, even if the region’s paleogeography has not had a major influence in plant evolutionary patterns, climatic processes in Afro-Arabia do seem to explain some disjoint distributions due to vicariance as a vicariant agent (Thiv et al. 2010). Additionally, some biogeographic patterns in plants are tightly related to Miocene dispersal in animal taxa, such as primates, through the Compsophtherium bridge (Zhou et al. 2012).

The use of null models and adequacy testing is common in areas of evolutionary biology and ecology, such as the study of diversification dynamics (Fitzjohn 2012; Beaulieu and O’Meara 2016; Caetano et al. 2018). Our approach, complemented by recent advances in the analysis of community assembly (Hua and Bromham 2012; Beaulieu and O’Meara 2016) and biogeographic assessments that control for unmodeled processes (i.e., phylogenetic diversification (Jiang et al. 2019), represents a growing recognition of the potential inadequacy of models and the need to test for it. Here, we show that model inadequacy can help us to better understand complex biological patterns. By quantifying model adequacy through time, we have identified specific time periods during which unmodeled processes (i.e., tectonic plate movement) may have led to unexpectedly high rates of vicariance. Our approach made it possible to disentangle extrinsic and intrinsic factors that may have determined the evolutionary history of the groups. Ultimately, such a perspective can provide an important insight from the past to help us face the challenges imposed by the intense changes in our planet’s present and future environmental conditions.

**Supplementary Material**

Data available from the Dryad Digital Repository: http://dx.doi.org/10.5061/dryad.wpzgnsbkn.

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