INTRODUCTION

The Mediterranean islands are populated by rich biotic communities, which comprise a mixture of recently arrived species and ancient insular radiations (Fois et al., 2020). Similar to other regions of the world, the diversity of these island communities has been influenced by geographic isolation and the geophysical characteristics of islands (surface area and topography; Kadmon & Allouche, 2007; MacArthur, 1965). The sea constitutes a powerful barrier to faunal movement, strongly reducing the species that can reach islands (Fattorini, 2002). However, animals can bypass this barrier using intermittent land corridors, or across the sea by swimming or by drifting/floating objects (Spennemann, 2020; Stankiewicz et al., 2006).

The colonization routes followed to reach an island determine the composition of its faunal assemblages. The formation of land corridors allows the almost barrier-free flow of continental fauna, whereas sweepstake dispersal routes are associated with strong filtering and species poor assemblages (Mazza et al., 2013; Simpson, 1940). The great faunal diversity of the communities on Mediterranean islands may be explained by the fact that species reached the islands using several dispersal routes (Poulakakis et al., 2013). However, unlike other regions of the world (e.g., in oceanic islands, such as the Galapagos or Samoa, in which natural colonization from the mainland probably occurred by rafting dispersal; Caccone et al., 1999; Gill, 1993), it is possible that dispersal among the Mediterranean islands through land corridors played a very important role during the colonization process given that this sea is a closed basin.

The importance of dispersal through land corridors is also supported by phylogeographic studies, which suggested that there is no significant phylogenetic signal present in the characteristics of the dispersal paths (i.e., in the distance traveled/bypassed sea depth). Thus, no superior island-colonizer lineages were detected in Mediterranean Squamata. However, our analyses also revealed that small-sized lizards were superior to other groups of squamates at dispersing over long distances on the sea.
a link between island connectivity and the molecular divergence of subpopulations of species/genera across island systems (Kornilios et al., 2019; Thompson, 1999). Colonization following the formation of land bridges occurred during several phases from the Middle Pliocene (17 Mya) to the later glacial eustatic regressions (20 Kya; Parmakelis et al., 2006). However, some islands such as Mallorca, Crete, and Cyprus are surrounded by deep sea regions, and their isolation has possibly remained uninterrupted since the Messinian event (5.33 Mya; Palombo, 2018).

In this study, we evaluated the dispersal patterns of several species of Squamata (Reptilia) throughout the Mediterranean islands. These islands have relatively diverse reptile assemblages, although this species diversity depends greatly on the sizes of the islands and their geographical locations (Chondropoulos, 1986; Mayol, 1997; Figure 1). Most of the squamate island species only occur on a few islands or a single archipelago, thereby indicating a limited ability to disperse across the sea (Hurston et al., 2009). However, a few species have spread among several archipelagos, such as some geckoes, lacertid lizards, and skinks (Di Nicola & Mezzadri, 2018; Stille et al., 2021), and thus, these species may possess adaptive traits that enhance their capacity as island colonizers. If these traits are only shared among evolutionarily related species, their dispersal patterns should exhibit a phylogenetic signal (hypothesis i). We also tested (hypothesis ii) that squamates with small body sizes and located at lower trophic levels would have been more successful at dispersing across the sea (Holt et al., 1999; Krysko & MacKenzie-Krysko, 2016; Lomolino, 2005).

2 | MATERIALS AND METHODS

2.1 | Study region

The study region covered the Mediterranean basin (Figure 2). This basin includes a large number of islands, with broad variability in their isolation and geophysical characteristics (Arnold, 2008; Itescu et al., 2018). In total, 105 species of Squamata occur on these islands and their patterns of occurrence were evaluated based on data obtained from biogeographic atlases and scientific papers (see the references provided in Appendix S1). Squamate species have been separated into endemic and mainland in origin based on recent phylogeographic studies (Kindler et al., 2013; Kornilios et al., 2010, 2019; Kotsakiozi et al., 2018; Senczuk et al., 2019; Spilani et al., 2019; Stöck et al., 2016; Utiger & Schätti, 2004).

2.2 | Dispersal paths

The dispersal patterns were analyzed by decomposing the paths into two components: length of the route (= distance traveled) and depth of the sea floor along the route (= bypassed sea depth). These two components described the progressive difficulty of reaching an island across the sea (Heaney et al., 2005). The distance traveled was estimated with a minimum-cost vector network by connecting the vertices of a graph (Sessions, 1992). This procedure generated the shortest path that connected the island network without assuming any dispersal step (i.e., origin–destination). The distance traveled was also estimated by building a cost surface model, which assumed a direction of dispersal (origin–destination) and that the path followed the gradient of minimum resistance (Carroll et al., 2012).

We defined the origins in a different manner for island endemics and mainland species. For endemic species, we considered the origin as the largest central island (e.g., Ibiza for Podarcis pityusensis or Crete for Podarcis cretensis) or pairs of large central islands (Corsica-Sardinia or Mallorca-Menorca) within an archipelago (because endemic species are not restricted to a single island; Speybroeck et al., 2016). This definition is supported by the dispersal patterns described for some endemic species (e.g., Podarcis lilfordi, Terrasa et al., 2004). In mainland species, we considered the origin as the continental region nearest to the island where these species are distributed, except for those where the origin has been restricted by phylogeographic studies, such as Hemidactylus turcicus, Tarentola mauritanica, and Chalcides ocellatus (Kornilios et al., 2010; Rato et al., 2010, 2011).
In these latter species, only the regions where they are supposed to be native have been considered in the models. The regions of origin were mapped based on Schleich et al. (1996), Geniez (2015), Speybroeck et al. (2016), and IUCN (2021).

The surface resistance was modeled to minimize the traveling cost following the coastline by using three variables: the distance to the coast (greater resistance with distance), terrain elevation (greater resistance with elevation), and depth of the sea floor (greater resistance with sea depth). However, there is still some uncertainty regarding the level of resistance that the sea imposes on the movement of species, and this uncertainty was considered by building three models that assumed various levels of resistance (Beier et al., 2009). Model 1 (M1) assumed that the transmarine route imposed greater resistance than land, although it is still relatively easy to travel across the sea (e.g., in semi-aquatic snakes, or by drifting on wind-driven sea currents; Baker, 2015; Renner, 2004).

**FIGURE 2** Map of the study region, showing the islands included in the study (red dots)

**TABLE 1** Description of the variables and descriptive statistics

| Variable                  | Category       | Description                                              | Mean   | SE  |
|---------------------------|----------------|----------------------------------------------------------|--------|-----|
| Minimum tree              | Distance       | Minimum-cost vector network                              | 986.32 | 135.06 |
| Total travel              | Distance       | Summation of travel lengths                              | 815.59 | 142.83 |
| Average travel            | Distance       | Total travel/number of travels                           | 53.55  | 9.11 |
| Maximum single travel     | Distance       | Longest travel                                           | 279.60 | 40.66 |
| Average depth             | Sea floor depth| Mean sea depth along the travel                          | ~129.60 | 14.22 |
| Maximum depth             | Sea floor depth| Maximum depth along the travel                           | ~640.21 | 61.20 |
| Prop. travel <150 m       | Sea floor depth| Proportion of the travel with sea depth < −150 m         | 0.84   | 0.02 |

Note: The mean values and standard error (SE) were obtained for all species and for the models M1, M2, and M3 (Total travel—Prop. travel <150 m).

**FIGURE 3** PCA scatter plot showing the variation in the characteristics of the paths (blue vectors) among species of insular squamates. Lizards, red; Snakes, blue
FIGURE 4  Squamata species (Mediterranean islands) phylogenetic tree, showing the mapped values of the first two PCA axes (a) PC1 (travel distance); (b) PC2 (travel depth)
**Model 2** (M2) assumed that island colonization occurred mostly during recent eustatic regressions. Species could only travel along the transmarine route until they reached a depth threshold of around −150 m (i.e., minimum sea level during the last glacial cycle; Lambeck & Purcell, 2005). **Model 3** (M3) assumed that the transmarine route was very unlikely, so species maximized their movement across land routes, even by significantly increasing the distances traveled.

These models allowed us to generate a network of paths connecting the islands to each other and to the mainland, and we estimated the following variables from these paths: total distance traveled, average distance traveled, maximum distance traveled in a single trip (i.e., connecting two adjacent points), average sea depth along the trip, maximum sea depth along the trip, and percentage of trip above −150 m sea level (Table 1). The mean value of these variables for each model (M1, M2, M3) and species are shown in Appendix S2.

The cost surface models and species paths were generated using a digital elevation model of the sea floor (Becker et al., 2009) and the package GRASS-GIS (GRASS Development Team, 2020).

### 2.3 | Species data

We constructed a phylogenetic tree using a synthesis-based phylogeny generated from the TimeTree database (Kumar et al., 2017). TimeTree generates an optimal phylogenetic tree compared with other candidate topologies (Hedges et al., 2015). This approach is suitable for testing evolutionary hypotheses, and it usually produces similar results to those generated by purposely constructed phylogenies (Li et al., 2019). The pairwise divergence times between species were used to calculate the 10th percentile of the distances (measured in Mya), evaluating the species phylogenetic isolation relative to the complete pool of insular species (a measure related to the interspecific niche overlap; Münkemüller et al., 2014). This distance will be greater if the species shows a distant phylogenetic relationship to other island species, which could favor the colonization potential of this species. The species were also grouped in the following categories: snakes/lizards, island endemic/mainland, and based on trophic preferences (vertebrates/invertebrates). We also included the average total length (snout tip to the tip of the tail for adult specimens) for each species. These data were obtained from several previously published sources (Baier et al., 2013; Di Nicola & Mezzadri, 2018; Geniez, 2015; Schleich et al., 1996; Speybroeck et al., 2016).

### 2.4 | Data analysis

The analyses tested (i) the strength of the phylogenetic signal in the dispersal patterns, and (ii) the effects of the species groups (i.e., phylogenetic isolation, snakes, endemics, trophic rank, and body size) in the dispersal patterns. Continuous variables with skew and kurtosis values that indicated non-normality were logarithmically transformed prior to the analyses (Sokal & Rohlf, 1995). The species associations with the variables describing the components of the dispersal paths were visualized using principal component analysis (PCA; Pearson, 1901).

| TABLE 2 | Evaluation of the phylogenetic signal (Pagel’s λ and Blomberg’s K) for the dispersion variables, including the uncertainty generated by the three dispersion models (M1–M3) for the variables total travel to prop. travel −150 |
|----------|------------------|------|------|
| Minimum tree | Estimate | λ | K | p-Value |
| | | 0.209 | 0.089 | | 0.0079 | 0.0139 |
| Total travel | Estimate | 0.214 | 0.063 | | 0.153–0.276 | 0.057–0.069 | .4663 | .5044 |
| Average travel | Estimate | 0.008 | 0.081 | | 0.004–0.011 | 0.074–0.087 | .8828 | .4496 |
| Max single travel | Estimate | 0.069 | 0.056 | | 0.053–0.084 | 0.053–0.060 | .4404 | .5182 |
| Average depth | Estimate | 0.003 | 0.162 | | 0.0006–0.0044 | 0.150–0.174 | .9607 | .1938 |
| Max depth | Estimate | 0.012 | 0.066 | | 0.000–0.028 | 0.058–0.074 | .9347 | .5419 |
| Prop. travel −150 | Estimate | 0.137 | 0.059 | | 0.116–0.157 | 0.058–0.061 | .3202 | .3389 |
The intensity of the phylogenetic signal was determined by calculating Pagel's lambda and Blomberg's K statistics (Münkemüller et al., 2012). Pagel's lambda values vary between 0 and 1, where 0 denotes that a trait has evolved independently of the phylogeny and values close to 1 correspond to a Brownian model that indicates a phylogenetic signal during trait evolution (Pagel, 1999). Blomberg's K values vary between 0 and ∞, where values of K < 1 represent less phylogenetic signal than that expected under Brownian motion (Blomberg et al., 2003). When estimating the phylogenetic signal, we considered the uncertainty of the cost surface models and the differences between groups of squamates (if these groups included more than 30 species because the lambda and K parameters are sensitive to small phylogenies; Münkemüller et al., 2012). These analyses were conducted using the sensiPhy package (Paterno et al., 2018) for the R environment (R Development Core Team, 2021).

Associations between the species groups and characteristics of the dispersal paths were tested using phylogenetic linear regression models (PLMs; Revell, 2010). Total length was included in the models as an interacting term with snake/lizard category because most of the lizards in the region have total length values in the lower range for snakes (Speybroeck et al., 2016). PLMs were built by incorporating several phylogenetic structures for the error term: Brownian model (BM), the Ornstein–Uhlenbeck model with fixed root (OU1), the Ornstein–Uhlenbeck model with random root (OU2), Pagel's lambda, Pagel's kappa, Pagel's delta, and the early burst model (EB; Ho & Ané, 2014; Pagel, 1999). In these models, we also accounted for spatial effects (e.g., those associated with the uneven distribution of islands throughout the basin), including a variable generated with the centroids of the geographical coordinates of species. The optimal PLM was selected after comparing them against a null model (without covariance structures) using the delta AIC criterion and AIC weights (Burnham & Anderson, 2002). Models with delta AIC < 2 had great support, and AIC weights close to 1 indicated a higher probability of being the best candidate (Symonds & Moussalli, 2011). These analyses were conducted with the phylolm package (Ho & Ané, 2014) for the R environment.

3 | RESULTS

The first two axes obtained by PCA accounted for a large proportion of the variance (cumulative proportion = 0.916, PC1 = 0.805, PC2 = 0.111; Figure 3). On the first axis, the variables that described the distance traveled (total distance traveled, average distance traveled, and maximum distance traveled in a single trip) accounted for 67.7% of the total variance (Figure 3). On the second axis, the variables that described the bypassed sea depth (average depth and maximum depth along the routes) accounted for 61.9% of the total variance (Figure 3).

The phylogenetic relationships among species on the first two axes obtained by PCA are shown in Figure 4. The estimates of Pagel's lambda and Blomberg's K indicated that no phylogenetic signal was present in any of the path properties (Figure 4; Table 2). In all cases, the phylogenetic signal was either insignificant or significant and close to 0 (i.e., phylogenetically related species shared fewer similarities in terms of their dispersal with each other than to those at greater evolutionary distances; Table 2). We also detected no phylogenetic signal when lizards and snakes were evaluated separately (Table 3).

The model with no phylogenetic covariance structures obtained the best fit for this data set (Table 4). The PLMs showed that the bypassed sea depth had a significant spatial component (average depth, maximum depth, and proportion of trip above −150 m; Table 5), where they indicated a significant negative association with longitude. This result implied that the dispersal paths in the eastern Mediterranean bypassed deeper marine regions (Figure 5). The regression analyses also showed that snakes traveled larger average distances whereas island endemics traveled shorter average distances and that small-sized lizards traveled larger distances than other groups of squamates (Table 5).

4 | DISCUSSION

In this study, we modeled the dispersal paths of squamates throughout the Mediterranean archipelagos. The dispersal models accounted for the uncertainty regarding the resistance imposed by the sea on the movements of species, which could generate a range of routes, particularly when evaluating long traveling distances. However, the

| TABLE 3 Evaluation of the phylogenetic signal (Pagel's λ and Blomberg's K) for the dispersion variables, separating the groups of Squamata snakes and lizards |
|-----------------|-----------------|-----------------|-----------------|-----------------|
|                  | λ               | p-Value         | K               | p-Value         |
| Minimum tree     |                 |                 |                 |                 |
| Snakes           | 0.00007         | 1.00            | 0.083           | .331            |
| Lizards          | 0.00005         | 1.00            | 0.188           | .043            |
| Total travel     |                 |                 |                 |                 |
| Snakes           | 0.00005         | 1.00            | 0.160           | .509            |
| Lizards          | 0.019           | .855            | 0.039           | .652            |
| Average travel   |                 |                 |                 |                 |
| Snakes           | 0.00004         | 1.00            | 0.127           | .732            |
| Lizards          | 0.064           | .360            | 0.035           | .829            |
| Max single travel|                 |                 |                 |                 |
| Snakes           | 0.00006         | 1.00            | 0.187           | .341            |
| Lizards          | 0.053           | .525            | 0.037           | .763            |
| Average depth    |                 |                 |                 |                 |
| Snakes           | 0.00005         | 1.00            | 0.179           | .366            |
| Lizards          | 0.577           | .683            | 0.094           | .017            |
| Max depth        |                 |                 |                 |                 |
| Snakes           | 0.080           | .488            | 0.151           | .585            |
| Lizards          | 0.576           | .138            | 0.061           | .177            |
| Prop. travel ≤150|                 |                 |                 |                 |
| Snakes           | 0.590           | .350            | 0.196           | .286            |
| Lizards          | 0.144           | .171            | 0.052           | .323            |
analyses obtained similar results when the distances traveled were estimated without assuming a dispersal direction and landscape resistance. Our results highlighted the lack of a phylogenetic signal in the dispersal paths when evaluating both the distances traveled and bypassed sea depth. This finding indicated that no squamate lineages in the Mediterranean basin had greater dispersal capacities than others. It was also feasible that more intense interspecific competition between phylogenetic relatives in small islands (Escoriza, 2020) could counteract the effects of traits that might possibly enhance transmarine dispersal.

Snakes and lizards differed in terms of their dispersal patterns. The analyses revealed that snakes show greater average traveled distances. Thus, when snakes traveled similar distances to lizards, they did so with fewer "stops" (i.e., by populating less intermediate islands). The lower number of stops may have been a consequence of the failure of snakes to colonize small islets or to maintain stable populations on these islets for long periods because most of the snakes, even those with small body sizes, occupied higher trophic levels than lizards (Pernetta et al., 2011). However, this result could also have been a human-induced artifact given that the longest paths traveled by Mediterranean snakes are due to translocations (e.g., *Erix jaculus*, *Hierophis viridiflavus*, or *Hemorrhois hippocrepis*; Utiger & Schätti, 2004; Pinya & Carretero, 2011; Insacco et al., 2015), and thus, they do not follow the expected sequence of island chains. Natricine snakes are possible exceptions because they are efficient marine dispersers (Brischoux & Kornilev, 2014; Kyriazi et al., 2013) but confined to large islands due to their requirements for permanent freshwater habitats (Zotos et al., 2021). However, in this group of semi-aquatic snakes, at least two island populations of *Natrix maura* (Mallorca and Sardinia) were introduced (Guicking et al., 2008).

Our analyses also indicated that the island endemics had shorter average traveled distances. Thus, when endemic species traveled similar distances to mainland species, they had a greater number of "stops" (or by populating a larger number of intermediate islands). This is consequence of the process of radial dispersion by following an ordered array of islands (i.e., from the closest to the furthest). This is to be expected considering that very few populations of endemic squamates had anthropogenic origins (Lo Cascio et al., 2006; Pérez-Mellado et al., 2017). The regression models demonstrated the importance of the spatial component in the dispersal patterns. For example, squamates crossed deeper regions of the sea in the eastern Mediterranean region where the density of islands is also greater (Arnold, 2008).

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### Table 4

|                  | BM  | OU1 | OU2 | Lambda | Kappa | Delta | EB | Null |
|------------------|-----|-----|-----|--------|-------|-------|----|------|
| Minimum tree     |     |     |     |        |       |       |    |      |
| ΔAIC             | 91.62 | 0.96 | 0.96 | 2.00   | 31.60 | 73.09 | 93.62 | 0.00 |
| AICw             | 0.00 | 0.24 | 0.24 | 0.14   | 0.00  | 0.00  | 0.00 | 0.38 |
| Total travel     |     |     |     |        |       |       |    |      |
| ΔAIC             | 141.40 | 9.34 | 9.34 | 2.00   | 43.62 | 120.43 | 143.40 | 0.00 |
| AICw             | 0.00 | 0.007 | 0.007 | 0.26   | 0.00  | 0.00  | 0.00 | 0.72 |
| Average travel   |     |     |     |        |       |       |    |      |
| ΔAIC             | 157.92 | 12.67 | 12.67 | 2.00   | 48.38 | 135.25 | 159.92 | 0.00 |
| AICw             | 0.00 | 0.001 | 0.001 | 0.27   | 0.00  | 0.00  | 0.00 | 0.73 |
| Max single travel|     |     |     |        |       |       |    |      |
| ΔAIC             | 136.82 | 9.33 | 9.33 | 2.00   | 44.11 | 115.83 | 138.82 | 0.00 |
| AICw             | 0.00 | 0.007 | 0.007 | 0.26   | 0.00  | 0.00  | 0.00 | 0.72 |
| Average depth    |     |     |     |        |       |       |    |      |
| ΔAIC             | 48.41 | 3.46  | 3.46 | 2.16   | 27.89 | 35.59 | 50.41 | 0.00 |
| AICw             | 0.00 | 0.11  | 0.11 | 0.19   | 0.00  | 0.00  | 0.00 | 0.59 |
| Max depth        |     |     |     |        |       |       |    |      |
| ΔAIC             | 91.69 | 8.10  | 8.10 | 2.00   | 39.73 | 76.74 | 93.69 | 0.00 |
| AICw             | 0.00 | 0.01  | 0.01 | 0.26   | 0.00  | 0.00  | 0.00 | 0.72 |
| Prop. travel<sub>150</sub> |     |     |     |        |       |       |    |      |
| ΔAIC             | 132.48 | 9.35  | 9.35 | 2.00   | 49.40 | 111.28 | 134.48 | 0.00 |
| AICw             | 0.00 | 0.007 | 0.007 | 0.26   | 0.00  | 0.00  | 0.00 | 0.72 |

Note: The best candidate is the one that shows a delta AIC (ΔAIC) <2 and AIC weight (AICw) close to 1.

Abbreviations: BM, Brownian motion model; delta, Pagel’s δ model; EB, early burst model; kappa, Pagel’s κ model; lambda, Pagel’s λ model; OU1, Ornstein–Uhlenbeck model with fixed root; OU2, Ornstein–Uhlenbeck model with random root.
Our analyses indicated that a morphological trait could influence the transmarine dispersal process, at least in lizards. Body size was negatively associated with the distance traveled in lizards, possibly because small lizards can more readily populate tiny intermediate islands (Delaugerre & Corti, 2020; Pafilis et al., 2020) or be transported accidentally by man (Austin, 1999). In addition, small-bodied lizards tend to have larger population sizes, which could favor the probability of successful dispersal by rafting, including the use of a

| Model statistics | Variables | Estimate | p-Value |
|------------------|-----------|----------|---------|
| Minimum tree     |           |          |         |
| $R^2$            | .209      | Latitude | -0.147  | .0003   |
| AIC              | 333.127   | Endemic  | -0.622  | .0524   |
| AICw             | 0.297     | TL:no    | -0.524  | .0353   |
| Total travel     |           |          |         |
| $R^2$            | .067      | TL:no    | -0.580  | .1557   |
| AIC              | 452.252   |          |         |
| AICw             | 0.315     |          |         |
| Average travel   |           |          |         |
| $R^2$            | .150      | Latitude | -0.101  | .0439   |
| AIC              | 346.821   | Snake    | 0.596   | .0282   |
| AICw             | 0.511     | Endemic  | -0.850  | .0129   |
| Max single travel|           |          |         |
| $R^2$            | .062      | TL:no    | -0.554  | .1200   |
| AIC              | 423.253   |          |         |
| AICw             | 0.377     |          |         |
| Average depth    |           |          |         |
| $R^2$            | .144      | Longitude| -0.041  | .0005   |
| AIC              | 327.907   | Snake    | -0.399  | .0933   |
| AICw             | 0.467     |          |         |
| Max depth        |           |          |         |
| $R^2$            | .112      | Longitude| -0.047  | .0035   |
| AIC              | 394.572   | Snake    | -0.719  | .0287   |
| AICw             | 0.572     |          |         |
| Prop. travel     |           |          |         |
| $R^2$            | .236      | Latitude | 0.029   | .0023   |
| AIC              | -13.733   | Longitude| -0.007  | .0044   |
| AICw             | 0.373     | Endemic  | -0.129  | .0296   |

Abbreviations: AIC, Akaike information criterion; AICw, AIC weight; TL :no, interaction between total length and snake category (yes/no).

FIGURE 5 Dispersal paths of Squamata in the Mediterranean islands. (a) Minimum shortest distance; (b) modeled paths in lizards; (c) snakes; (d) island endemics. Blue, M1; yellow, M2; red, M3. When the paths overlap, only M2 is shown.
wider range of raft sizes (Hsu et al., 2021; Novosolov et al., 2016). Examples of highly successful island colonizers with small body sizes are found in several distantly related lizard lineages within the Mediterranean region, including skinks (e.g., Ablepharus), geckoes (Euleptes and Mediodactylus), and lacertids (Ophisops), and thus, a phylogenetic signal was not detected.

5 | CONCLUSIONS

In this study, for the first time, we evaluated the dispersal patterns of Squamata in the Mediterranean islands by estimating several parameters to quantify the difficulty of reaching these islands. The results highlighted the lack of any phylogenetic signal in the dispersal process, thereby indicating that no evolutionary lineages had superior colonization capacities. Our analyses also revealed important differences in the dispersal process for snakes compared with lizards, although these differences could have been human-induced artifacts because some long-distance dispersals of snakes may possibly have been due to accidental introductions. In lizards, small body size possibly enhanced the probability of success over long-distance sea routes.

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CONFLICT OF INTEREST

None declared.

AUTHOR CONTRIBUTION

Daniel Escoriza: Conceptualization (equal); Data curation (equal); Formal analysis (equal); Funding acquisition (equal); Investigation (equal); Methodology (equal); Project administration (equal).

DATA AVAILABILITY STATEMENT

The raw data used in the present study are available within the article or its Supporting Information. Dataset for modeled trajectories is available at Dryad Digital Repository: https://doi.org/10.5061/dryad.zw3r2288c

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REFERENCES

Arnold, C. (2008). Mediterranean Islands. Survival Books.
Austin, C. C. (1999). Lizards took express train to Polynesia. Nature, 397, 113–114. https://doi.org/10.1038/16365
Baier, F., Sparrow, D. J., & Wiedl, H. J. (2013). The amphibians and reptiles of Cyprus. Chimaira.
Baker, J. M. (2015). Marine sightings of grass snakes Natrix natrix. Herpetological Bulletin, 131, 30–31.
Becker, J. J., Sandwell, D. T., Smith, W. H. F., Braud, J., Binder, B., Depner, J., Fabre, D., Factor, J., Ingalls, S., Kim, S.-H., Ladner, R., Marks, K., Nelson, S., Pharaoh, A., Trimmer, R., Von Rosenberg, J., Wallace, G., & Weatherall, P. (2009). Global bathymetry and elevation data at 30 arc seconds resolution: SRTM30_PLUS. Marine Geodesy, 32, 355–371. https://doi.org/10.1080/01490410903297766
Beier, P., Majka, D. R., & Newell, S. L. (2009). Uncertainty analysis of least-cost modeling for designing wildlife linkages. Ecological Applications, 19, 2067–2077. https://doi.org/10.1890/08-1898.1
Blomberg, S. P., Garland, T. Jr, & Ives, A. R. (2003). Testing for phylogenetic signal in comparative data: Behavioral traits are more labile. Evolution, 57, 717–745. https://doi.org/10.1111/j.0014-3820.2003.tb00285.x
Brischoux, F., & Kornilev, Y. V. (2014). Hypernatremia in Dice snakes (Natrix tessellata) from a coastal population: Implications for osmoregulation in marine snake prototypes. PLoS One, 9, e92617. https://doi.org/10.1371/journal.pone.0092617
Burnham, K. P., & Anderson, D. R. (2002). Model selection and multimodel inference: A practical information-theoretic approach. Springer-Verlag.
Caccone, A., Gibbs, J. P., Ketmaier, V., Suatoni, E., & Powell, J. R. (1999). Origin and evolutionary relationships of giant Galápagos tortoises. Proceedings of the National Academy of Sciences, 96, 13223–13228. https://doi.org/10.1073/pnas.96.23.13223
Carroll, C., McRae, B. H., & Brookes, A. (2012). Use of linkage map centrality analysis and population diversity to evaluate connectivity of gray wolf populations in western North America. Conservation Biology, 26, 78–87. https://doi.org/10.1111/j.1523-1739.2011.01753.x
Chondropoulos, B. P. (1986). A checklist of the Greek reptiles. I. The lizards. Amphibia-Reptilia, 7, 217–235. https://doi.org/10.1163/156853886X00028
Delauwere, M. J., & Corti, C. (2020). Tiny but “strong”: The European Leaf-toed gecko, Euleptes europaea, a terrestrial vertebrate able to survive on tiny islets. Israel Journal of Ecology and Evolution, 66, 223–230. https://doi.org/10.1163/22244662-bja10017
Di Nicola, M. R., & Mezzadri, S. (2018). Anfibi e rettili di Sardegna. Libreria della Natura.
Esoriza, D. (2020). Organization of Squamata (Reptilia) assemblages in Mediterranean archipelagos. Ecology and Evolution, 10, 1592–1601. https://doi.org/10.1002/ece3.6013
Fattorini, S. (2002). Biogeography of the tenebrionid beetles (Coleoptera, Tenebrionidae) on the Aegean Islands (Greece). Journal of Biogeography, 29, 49–67. https://doi.org/10.1046/j.1365-2699.2002.06656.x
Fois, M., Podda, L., Médail, F., & Bacchetta, G. (2020). Endemic and alien vascular plant diversity in the small Mediterranean islands of Sardinia: Drivers and implications for their conservation. Biological Conservation, 244, 108519. https://doi.org/10.1016/j.biocon.2020.108519
Geniez, P. (2015). Serpents d’Europe, d’Afrique du Nord et du Moyen-Orient. Delachaux & Niestlé.
Gill, B. J. (1993). The land reptiles of Western Samoa. Journal of the Royal Society of New Zealand, 23, 79–89. https://doi.org/10.1080/03036758.1993.10721219
GRASS Development Team. (2020). Geographic Resources Analysis Support System (GRASS) Software, Version 7.8. Open Source Geospatial Foundation. Retrieved from https://grass.osgeo.org
Guicking, D., Joger, U., & Wink, M. (2008). Molecular phylogeography of the viperine snake Natrix maura (Serpentes: Colubridae): Evidence for strong intraspecific differentiation. Organisms Diversity & Evolution, 8, 130–145. https://doi.org/10.1016/j.ode.2007.05.001
Heaney, L. R., Walsh, J. S. Jr, & Peterson, A. T. (2005). The roles of geological history and colonization abilities in genetic differentiation between mammalian populations in the Philippine archipelago. Journal of Biogeography, 32, 229–247. https://doi.org/10.1111/j.1365-2699.2004.01120.x
Hedges, S. B., Marin, J., Suleski, M., Paymer, M., & Kumar, S. (2015). Tree of life reveals clock-like speciation and diversification. Molecular
Pérez-Mellado, V., Pérez-Cembranos, A., Rodríguez, V., Buades, J. M., Brown, R. P., Böhme, W., Terrasa, B., Castro, J. A., Picornell, A., & Ramon, C. (2017). The legacy of translocations among populations of the Ibizan Wall Lizard, Podarcis pityusensis (Squamata: Lacertidae). Biological Journal of the Linnean Society, 121, 82–94. https://doi.org/10.1093/biolinnean/blw026

Pernetta, A. P., Allen, J. A., Beebee, T. J. C., & Reading, C. J. (2011). Fine-scale population genetic structure and sex-biased dispersal in the smooth snake (Coronella austriaca) in southern England. Heredity, 107, 231–238. https://doi.org/10.1038/hdy.2011.7

Pinya, S., & Carretero, M. A. (2011). The Balearic herpetofauna: A species update and a review on the evidence. Acta Herpetologica, 6, 59–80.

Poulakakis, N., Kapli, P., Kardamaki, A., Skourtanioti, E., Gökmen, B., Ilgaz, Ç., Kumlutaş, Y., Avci, A., & Lymberakis, P. (2013). Comparative phylogeography of six herpetofauna species in Cyprus: Late Miocene to Pleistocene colonization routes. Biological Journal of the Linnean Society, 108, 619–635. https://doi.org/10.1111/j.1095-8312.2012.02039.x

R Development Core Team. (2021). R package vs 4.0.4. Retrieved from https://cran.r-project.org/bin/windows/base/

Rato, C., Carranza, S., & Harris, D. J. (2011). When selection deceives phylogeographic interpretation: The case of the Mediterranean house gecko, Hemidactylus turcicus (Linnaeus, 1758). Molecular Phylogenetics and Evolution, 58, 365–373. https://doi.org/10.1016/j.ympev.2010.12.004

Rato, C., Carranza, S., Perera, A., Carretero, M. A., & Harris, D. J. (2010). Conflicting patterns of nucleotide diversity between mtDNA and nDNA in the Moorish gecko, Tarentola mauritanica. Molecular Phylogenetics and Evolution, 56, 962–971. https://doi.org/10.1016/j.ympev.2010.04.033

Renner, S. (2004). Plant dispersal across the tropical Atlantic by wind and sea currents. International Journal of Plant Sciences, 165, S23–S33. https://doi.org/10.1086/383334

Revell, L. J. (2010). Phylogenetic signal and linear regression on species data. Methods in Ecology and Evolution, 1, 319–329. https://doi.org/10.1111/j.2041-210X.2010.00044.x

Schleich, H. H., Kästle, W., & Kabisch, K. (1996). Amphibians and reptiles of North Africa. Koeltz.

Sencuzk, G., Castiglia, R., & Böhme, W. (2019). Podarcis siculus latastei (Bedriaga, 1879) of the Western Pontine Islands (Italy) raised to the species rank, and a brief taxonomic overview of Podarcis lizards. Acta Herpetologica, 14, 71–80.

Sessions, J. (1992). Solving for habitat connections as a Steiner network problem. Forest Science, 38, 203–207. https://doi.org/10.1093/forestscience/38.1.203

Simpson, G. G. (1940). Mammals and land bridges. Journal of the Washington Academy of Sciences, 30, 137–163.

Sokal, R. R., & Rohlf, F. J. (1995). Biometry (3rd ed.). W. H. Freeman.

Spennemann, D. H. (2020). Cruising the currents: Observations of extra-limital Saltwater Crocodiles (Crocodylus porosus Schneider, 1801) in the Pacific Region. Pacific Science, 74, 211–213. https://doi.org/10.2984/74.3.1

Speybroeck, J., Beukema, W., Bok, B., & Van Der Voort, J. (2016). Field guide to the amphibians and reptiles of Britain and Europe. Bloomsbury.

Sokal, R. R., & Rohlf, F. J. (1995). (3rd ed.). W. H. Freeman.

Stankiewicz, J., Thiari, C., Masters, J. C., & De Witi, M. J. (2006). Did lemurs have sweepstake tickets? An exploration of Simpson’s model for the colonization of Madagascar by mammals. Journal of Biogeography, 33, 221–235. https://doi.org/10.1111/j.1365-2699.2005.01381.x

Stille, M., Gasteratos, I., & Stille, B. (2021). Alien and invasive terrestrial vertebrate species on Corfu, Ionian Islands, Greece. Journal of Vertebrate Biology, 70, 20126–20213. https://doi.org/10.25225/jvb.20126

Stöck, M., Grifoni, G., Armor, N., Scheidt, U., Sicilia, A., & Novarini, N. (2016). On the origin of the recent herpetofauna of Sicily: Comparative phylogeography using homologous mitochondrial and nuclear genes. Zoologischer Anzeiger, 261, 70–81. https://doi.org/10.1016/j.jcz.2015.10.005

Symonds, M. R., & Moussalli, A. (2011). A brief guide to model selection, multimodel inference and model averaging in behavioural ecology using Akaike’s information criterion. Behavioral Ecology and Sociobiology, 65, 13–21. https://doi.org/10.1007/s00265-010-1037-6

Terrasa, B., Picornell, A., Castro, J. A., & Ramon, M. M. (2004). Genetic variation within endemic Podarcis lizards from the Balearic Islands inferred from partial Cytochrome B sequences. Amphibia Reptilia, 25, 407–414. https://doi.org/10.1163/1568538042788960

Thompson, J. D. (1999). Population differentiation in Mediterranean plants: Insights into colonization history and the evolution and conservation of endemic species. Heredity, 82, 229–236. https://doi.org/10.1046/j.1365-2540.1999.00504.x

Utiger, U., & Schätti, B. (2004). Morphology and phylogenetic relationships of the Cyprus racer. Revue Suisse De Zoologie, 111, 225–238. https://doi.org/10.5962/bhl.part.80236

Zotos, S., Stamatiou, M., Naziri, A., Meletiou, S., Demosthenous, S., Perikleous, K., Eroktirou, E., Xenophontos, M., Zavrou, D., Michael, K., & Sergides, L. (2021). New evidence on the distribution of the highly endangered Natrix matrix cypriaca and implications for its conservation. Animals, 11(4), e1077. https://doi.org/10.3390/ani11041077

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