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Ecological specialization, rather than the island effect, explains morphological diversification in an ancient radiation of geckos

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Island colonists are often assumed to experience higher levels of phenotypic diversification than continental taxa. However, empirical evidence has uncovered exceptions to this 'island effect'. Here, we tested this pattern using the geckos of the genus Pristurus from continental Arabia and Africa and the Socotra Archipelago. Using a recently published phylogeny and an extensive morphological dataset, we explore the differences in phenotypic evolution between Socotran and continental taxa. Moreover, we reconstructed ancestral habitat occupancy to examine if ecological specialization is correlated with morphological change, comparing phenotypic disparity and trait evolution between habitats. We found a heterogeneous outcome of island colonization. Namely, only one of the three colonization events resulted in a body size increase. However, in general, Socotran species do not present higher levels or rates of morphological diversification than continental groups. Instead, habitat specialization explains better the body size and shape evolution in Pristurus. Particularly, the colonization of ground habitats appears as the main driver of morphological change, producing the highest disparity and evolutionary rates. Additionally, arboreal species show very similar body size and head proportions. These results reveal a determinant role of ecological mechanisms in morphological evolution and corroborate the complexity of ecomorphological dynamics in continent-island systems.

1. Introduction

The life history and population biology of continental and insular taxa of a specific evolutionary radiation are fundamentally distinct [1,2]. In the continent, communities are often assumed to be complex and composed of many species that share a long history of coevolution [3]. In such a scenario, most of the ecological niches will be filled, and high levels of interspecific competition are expected [4]. These factors, together with higher predation pressures, will tend to limit niche expansion and, consequently, morphological diversification [5]. By contrast, insular groups are usually exposed to higher levels of ecological opportunity, and thus they can occupy the new or relatively unexploited adaptive landscapes that islands provide [6,7]. As a result, island species may experience increased rates of phenotypic diversification and higher levels of morphological disparity compared to continental taxa [8]. Moreover, the divergence in body size found in insular taxa relative to their continental counterparts is a pervasive and widely studied pattern across vertebrates, known as the 'island rule' [9]. However, empirical evidence outlines a more complex scenario in which island colonists

might not necessarily experience great levels of evolutionary divergence [10]. Instead, these potential outcomes of island colonization (i.e. body size and shape divergence, increased phenotypic disparity and increased rates of morphological evolution; from now on referred to as 'island effect') might depend on multiple *extrinsic* factors (mostly modulated by the geography or geology of the island), as well as *intrinsic* factors (i.e. the biological characteristics of the group concerned) [3,11]. Thus, ecological specialization is expected to be more pronounced when island colonization results in an expansion into novel ecological contexts, and such specialization might carry morphological changes depending on species and system-specific factors [4,6].

Whether ecological specialization follows island colonization or not, the study of habitat occupancy is essential to understand the evolution of associated traits and the structuring of ecological communities [12]. In particular, specialization in substrate use can promote morphological diversity through deterministic body size evolution and diversification [13]. Moreover, microhabitat use can be strongly correlated with convergent phenotypic evolution resulting in recurrent ecomorphs beyond the effect of history and clade membership [14].

Arid regions, generally considered relatively depauperate in terms of animal diversity, have been successfully inhabited by some vertebrate groups and harbour especially high levels of reptile diversity. Among reptiles, geckos are particularly prominent due to their outstanding diversity in ecological features, exhibiting a wide variety of morphological and behavioural adaptations [15]. Afro-Arabian geckos have been prominent in recent studies of the role of arid biomes in generating biodiversity [16-21]. However, the outcomes of morphological diversification in these animals have only been properly investigated within the genus Hemidactylus, which is the best-studied Arabian reptile group with wellresolved taxonomy and reliably reconstructed biogeographic history [18,19,22-27]. In particular, two recent studies including continental and insular taxa from the Socotra Archipelago indicated that the genus Hemidactylus shows an island effect at least regarding body size evolution [28,29]. By contrast, the geckos of the genus Pristurus, which have also colonized and diversified within the same archipelago, seem to show lower rates of body size diversification in Socotra than other genera in the archipelago but also compared to their continental relatives [28]. Despite these preliminary results, a more nuanced analysis of the morphological evolution in Pristurus, including undescribed diversity, morphological and ecological data, is still lacking. Interestingly, the genus Pristurus not only has representatives in both the Socotra Archipelago and continental Africa and Arabia, but it also has diversified ecologically to exploit a variety of habitats, including rocky and sandy surfaces, gravel plains and trees [30,31].

Here we use a recently inferred phylogenetic assessment of Afro-Arabian reptiles including undescribed diversity, together with an extensive morphological sampling and detailed ecological information, to explore the morphological evolution in *Pristurus* geckos. Specifically, we test alternative scenarios of body size and shape evolution in this genus, to determine the role of the colonization of the Socotra Archipelago and the ecological specialization in generating the morphological diversity observed. The independent diversification of both Socotran and continental taxa, the ecological and behavioural diversity and the unique phenotypic dataset compiled in this study, make this group of geckos an exceptional system to investigate keystone dynamics in evolutionary biology such as the island effect and ecological adaptation, and their impact on morphological evolution.

2. Material and methods

(a) Phylogenetic and ecological data

We used a recently published phylogenetic tree of Afro-Arabian squamates [32]. This tree contains all the species of *Pristurus* for which there exist genetic data, including some species currently in the process of being described, resulting in a total of 30 species. We extracted the *Pristurus* clade from the squamate tree, both for the consensus and for 1000 trees randomly selected from the posterior distribution generated in the cited study. Using a sample of posterior trees allowed us to take into account the phylogenetic and temporal uncertainty in the subsequent analyses.

Each species was defined as present in the Socotra Archipelago or in the continent (Africa and Arabia). For habitat specialization, each species was categorized based on substrate preferences into one of three groups: ground-dwelling, rock climber or arboreal [30,33]. Additionally, the ground-dwelling species were divided into 'soft-ground' (sandy surfaces) and 'hard-ground' (gravel plains) categories to further characterize the morphological adaptations to each type of ground habitat. Nevertheless, the disparity dynamics and rates of trait evolution were estimated with the original categorizations (continent— Socotra; and the three habitat states) which, due to the limited number of species, were more appropriate for the analyses.

(b) Ancestral reconstructions

We studied the colonization of the Socotra Archipelago and the habitat specialization through time by reconstructing ancestral states across the phylogeny. First, we fitted several models of character evolution across the phylogeny in order to select the best-fit model for both traits. With such a purpose, we used the function fitDiscrete from the R package geiger v. 2.0 [34,35]. We fitted three models: an equal-rates model (ER), a symmetrical model and an all-rates-different model. We selected the best-fit model in each case based on the Akaike information criterion, correcting for small sample size (AICc) [36]. We then used the function make.simmap from the R package phytools [37], which simulates plausible stochastic character histories after fitting a continuous-time reversible Markov model for the evolution of the character states assigned to the tips of the tree. We ran 1000 simulations with the previously selected model of character evolution (ER model for both traits). Additionally, we randomly selected 100 trees from the posterior distribution, and we ran 100 stochastic character histories on each of them for both traits (presence in Socotra and habitat occupancy).

(c) Phenotypic data

For the 30 species included in the phylogenetic tree, a total of 697 specimens were examined and measured, with a minimum of one, a maximum of 56 and a mean of 23 specimens per species (electronic supplementary material, table S1). All vouchers were obtained from the following collections: Institute of Evolutionary Biology (CSIC-UPF), Barcelona, Spain (IBE), Natural History Museum, London, UK (BM), Museo Civico di Storia Naturale, Carmagnola, Turin, Italy (MCCI), Università di Firenze, Museo Zoologico 'La Specola', Firenze, Italy (MZUF), Oman Natural History Museum (ONHM), Laboratoire de Biogéographie et Écologie des Vertébrés de l'École Pratique des Hautes Etudes, Montpellier, France (BEV) and National Museum Prague, Czech Republic (NMP). The following measurements were taken by the same person (MSR) using a digital caliper with accuracy to the nearest 0.1 mm: snout–vent length (SVL; distance from the tip of

the snout to the cloaca), trunk length (distance between forelimb and hindlimb measured posterior to the forelimb and anterior to the hindlimb insertion points), head length (HL; taken axially from tip of the snout to the anterior ear border), head width (HW; taken at anterior ear border), head height (HH; taken laterally at anterior ear border), humerus length (Lhu; from elbow to axilla), ulna length (Lun; from wrist to elbow), femur length (Lfe; from knee to groin) and tibia length (Ltb; from ankle to knee). Tail length was not measured because most of the specimens had regenerated tails or had lost it.

(d) Morphological differentiation

As body size and shape evolution might be affected by the colonization of Socotra and/or the ecological specialization, we investigated whether Socotran colonists are morphologically distinct from their relatives in the continent (one potential outcome under the 'island effect' framework) and likewise whether the differential habitat use is reflected in species morphology. For each species, the mean of each morphological variable was calculated and \log_{10} -transformed in order to improve normality and homoscedasticity prior to subsequent analyses.

To characterize shape differentiation, we performed a phylogenetic regression of each trait on SVL to remove the effect of the body size on the other variables. The residuals of these regressions were used to implement a phylogenetically controlled principal component analysis (pPCA) using the functions *phyl.resid* and *phyl.pca* from the R package phytools with the method set to 'lambda' [37]. We used the principal components representing 75% of the cumulative proportion of variance as shape variables and we visualized the two-dimensional shape morphospace. Additionally, we performed a PCA with the shape data from all the specimens measured, after correcting for body size through regressions on SVL similarly to our processing of the species data. We generated per-species boxplots of size and shape variation with the specimen data.

For body size, we used the function phenogram from phytools [37] to map and visualize variation in SVL across the species tree. For all the visualizations, we categorized the species according to their presence in the Socotra Archipelago (Socotra or continent) and to their habitat use (ground, rock or tree) separately, to have a detailed perspective of the extent of the morphological differentiation undergone in each category. Furthermore, to test the island rule prediction that body size in insular species is different from the continent, we performed a phylogenetic ANOVA with the phylANOVA function from phytools [37]. Additionally, according to the island rule, we would expect to see body size differences before and after a colonization event. To test this, we performed paired *t*-tests on the estimated body size for each pair of parental and descendent nodes in which a change from continent to Socotra occurred. We used the function OUwie.anc from the R package OUwie [38] to estimate the ancestral body size under a multi-rate Brownian motion model, which was the best-fit model for body size evolution (see below, 'Differences in tempos of phenotypic diversification').

(e) Exploring differences in phenotypic disparities

Since one of the main possible outcomes of island colonization and/or ecological specialization is the increase in phenotypic disparity, we tested this assumption following previous research on other geckos [29] and defining disparity as the average squared Euclidean distance between all pairs of species in a group for a given continuous variable [7], in our case body size (SVL) and two variables of shape (pPC1 and pPC2). First, we tested whether disparity is higher in Socotran species than in the continent, which would indicate the existence of an island effect. Then, considering our results of morphological differentiation according to habitat use, in which ground-dwelling species are widespread throughout most of the morphospace of the genus, we tested whether the phenotype of ground-dwelling species is significantly more disparate than that of species in all the other habitats together (i.e. rock and arboreal species). We calculated the observed disparity ratios (Socotra/continent and ground/no-ground) for each morphological variable, using the function disparity from the R package geiger [34]. In the case of higher disparity in Socotra or in ground species (disparity ratio Socotra/continent or ground/no-ground higher than 1), we then compared the observed ratios with a null distribution of disparity ratios obtained from 10 000 simulations of the evolution of a continuous character according to a Brownian motion model across the phylogeny. These simulations were performed by applying the function sim.char after estimating the empirical rate parameter for body size and shape from the bestfit model (Brownian motion) with the function fitContinuous, both from the package geiger [34]. This approach allowed us to test, in the case of an observed higher disparity in Socotran or ground species, whether this is a significant increase considering the rate of evolution of the character, or rather this is not evidence of effectively increased disparity. In order to test the sensitivity of the results to low intraspecific sampling, we performed the analyses described above also with a dataset in which the species with less than five specimens had been removed: Pristurus adrarensis, Pristurus flavipunctatus, P. sp. 4, P. sp. 9 and P. sp. 12 (see electronic supplementary material, table S1).

(f) Differences in tempos of phenotypic diversification

In order to test the effect of Socotran colonization and ecological specialization in the tempo of phenotypic evolution, we fitted different models of character evolution across the phylogeny in which the evolutionary rates of body size and shape might or might not differ between categories (i.e. between Socotra and continent, and between ground, rock and tree habitats). For body size, we used the R package OUwie [38] to fit three alternative models: BM1 (Brownian motion single rate, i.e. assuming one single-rate regime for all lineages in the phylogeny), OU1 (Ornstein-Uhlenbeck single-rate value with a phenotypic optimum and a selective pressure toward it) and BMS (Brownian motion multirate model, with different rate values for each of the regimes specified, i.e. Socotran different from continental lineages, and differences between habitats). Similarly, we studied the rates of phenotypic evolution for body shape, but in this case, we fitted multivariate models including the first two principal components resulting from the phylogenetic PCA (pPC1 and pPC2; 77% of the variance, see Results section). We used the R package mvMorph [39] to fit four alternative models: BM1, OU1 and BMM (analogous to BMS in the OUwie package) and OUM (multi-rate Ornstein-Uhlenbeck model). We fitted these models in the 1000 stochastic character maps generated for the consensus tree, and also in the 100 character maps on each of the 100 posterior trees used to reconstruct ancestral states (see above, 'Ancestral reconstructions'). We then selected the best-fit models based on the AICc distributions and means, and we extracted the distributions of rate values estimated for each regime (Socotra, continent, ground, rock and tree) in the multi-rate models, to unravel the effect of each trait in morphological rates. All the visualizations of the disparity and phenotypic rate analyses were built with the R packages ggplot2 [40], patchwork [41] and cowplot [42].

3. Results

(a) Ancestral reconstructions and morphological variation

The ancestral reconstructions following an ER model, with the probabilities of each state in ancestral nodes (Socotra

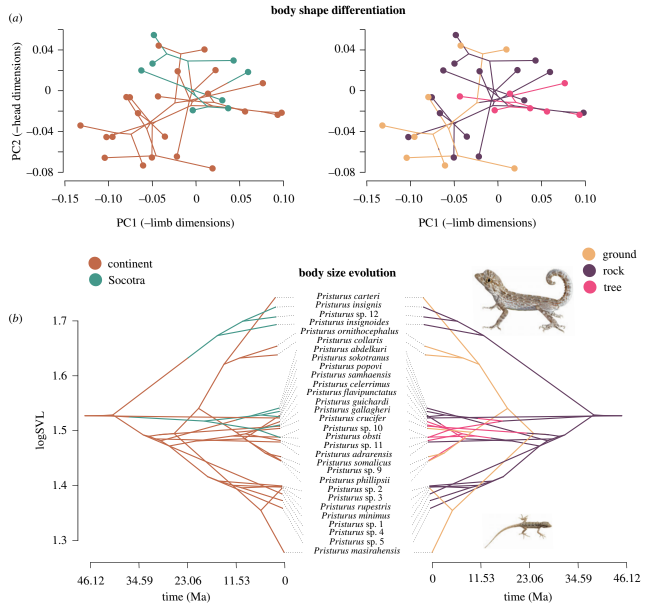


Figure 1. Morphological variability in *Pristurus*, with insight from insularity (left) and habitat use (right). (*a*) Morphospace with phylogenetic relationships between the species, showing body shape differentiation. (*b*) Traitgram showing body size (SVL) through time on the summary phylogenetic tree of *Pristurus*, mapped by the discrete categories of presence in Socotra or the continent (left) and by ecological specialization (right). Photos (proportional to species' SVL): *Pristurus carteri* (top) and *P. masirahensis* (bottom), the largest and smallest species of the genus, respectively. (Online version in colour.)

and continent; ground, rock and tree) can be found in the electronic supplementary material (electronic supplementary material, figure S1). For the Socotra–continent reconstruction, the most probable ancestral state for the root was 'continent' (continent: 0.649; Socotra: 0.351). The ecological reconstruction shows rocky habitats as the ancestral state in *Pristurus* (rock: 0.978; ground: 0.012; tree: 0.010), with several transitions to arboreal habitats and one colonization of the ground in the ancestor of the clade known as 'Spatalura group' [30,33] (electronic supplementary material, figure S1). One of the subclades of this group later colonized more compact, harder substrates, shown in our more detailed analysis separating soft- and hard-ground species (electronic supplementary material, figure S1).

The pPCA analysis of body shape resulted in two first components explaining 77% of the total variance: pPC1 (61% of the variance) mainly representing limb dimensions (variables Lhu, Lun, Lfe, Ltb), and pPC2 (16% of the variance) mostly representing head proportions (variables HL, HW, HH) (figure 1*a*; electronic supplementary material, table S2; see electronic supplementary material, figure S2 and table S3 for results on body size and shape differentiation using the specimen data and PCA). The morphospace occupied by continental species is notably larger than that occupied by Socotran species, and they overlap almost completely (figure 1*a*, left). When visualizing the shape phylomorphospace along with habitat categories, we observe a wide portion occupied by the ground-dwelling species, especially in pPC2 (head dimensions) (figure 1*a*, right). These eight species of the 'Spatalura group' essentially occupy almost as much of the morphospace as all the rest of the species together, with morphologies specialized to arboreal habits localized in a narrow area, especially for head proportions (figure 1*a*, right). 4

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We found a similar pattern for body size. Body size variability of Socotran species is completely contained in the range occupied by continental species (figure 1*b*, left). On the contrary, ground-dwelling species show a size variability higher than all the species from other habitats together, being the largest and the smallest species of *Pristurus* specialized to ground habitats (figure 1*b*, right). As with head proportions, arboreal

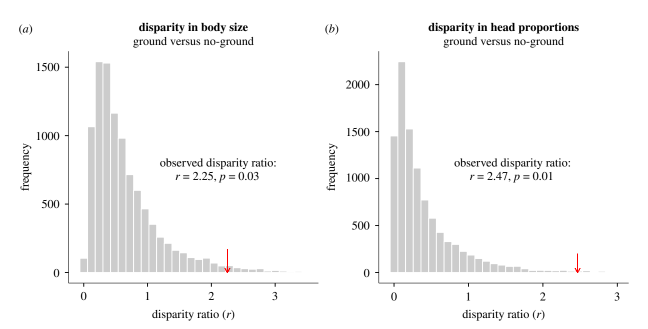


Figure 2. Observed (red arrows) and simulated (grey bars) ratios of phenotypic disparity between ground versus no-ground habitats. (a) Body size disparity ratios. (b) Head proportions (pPC2) disparity ratios. (Online version in colour.)

species have apparently constrained body sizes, being restricted to specific intermediate values within the genus' size range. The phylogenetic ANOVA showed no statistical differences in body size between Socotran and continental species (F = 7.535, p = 0.215). Although one of the colonization events of the Socotra Archipelago was followed by an apparent increase in body size (the clade composed by Pristurus insignis, Pristurus insignoides and P. sp. 12; see electronic supplementary material, figure S3), our paired *t*-test analysis did not find significant differences between parental and descendant nodes in which a change from continent to Socotra occurred (t = -1.201, p = 0.353; electronic supplementary material, figure S3). When separating ground species into hard- and soft-ground habitats, we observed a clear morphological differentiation, especially in body size. Hard-ground species seem to be highly specialized, with some of the largest body sizes of the genus, long limbs and large heads (electronic supplementary material, figure S4).

(b) Phenotypic disparity

We found that the morphological disparity in the Socotra Archipelago is lower than in the continent for the three variables, with disparity ratios Socotra/continent below 1 (SVL: 0.88; pPC1: 0.52; pPC2: 0.53). When comparing disparity between ground and no-ground habitats, we found a higher observed disparity in ground for body size (SVL) and head proportions (pPC2), with disparity ratios ground/no-ground of 2.25 for SVL, 0.86 for pPC1 and 2.47 for pPC2. Furthermore, both for size and head proportions, the increased disparity in ground habitats was significant compared with the null distribution of simulated disparity ratios ($p_{size} = 0.03$; $p_{head} = 0.01$; figure 2). These results were virtually identical to those from the analyses performed after removing the species with less than five specimens (electronic supplementary material, table S4).

(c) Rates of morphological evolution

For body size, a multi-rate BMS was the best fit both for presence in Socotra and for ecological specialization (lowest AICc; figure 3*a*), suggesting differences in the rate of morphological evolution across discrete categories (i.e. Socotra versus continent, and different habitats). For body shape, however, we did not find evidence for differences in evolutionary rates, being the single-rate Brownian motion model (BM1) the best fit both for limb (pPC1) and head (pPC2) dimensions, although the overlap across all models was considerable (figure 3*b*).

We extracted the rates of body size evolution from the Brownian motion multi-rate models, and we found that Socotran species present lower rates than species in the continent (figure 4*a*, top), although the bimodal distribution of the rates in Socotran taxa suggests rate heterogeneity among those clades. For ecological specialization, we found increased rates of body size evolution in the ground-dwelling species relative to the other habitats, with arboreal habitats showing the lowest rates (figure 4*a*, bottom). We also extracted per-category rates of body shape evolution according to the BMM model in mvMorph, even though the multi-rate models were not the best supported, and we found a similar scenario, in which shape evolution (both for limbs and for head proportions) would be notably faster in ground-dwelling species (figure 4b, bottom). Results from the analyses performed with the 1000 stochastic character maps on the consensus tree and with the 100 maps on each of 100 posterior trees lead to the same conclusions, so we show the ones from the posterior trees on the main text. Results from analyses with the consensus tree can be found in the electronic supplementary material (electronic supplementary material, figure S5).

4. Discussion

The present study represents the first comprehensive comparative work on the genus *Pristurus*, including undescribed diversity and extensive morphological (size and shape) and ecological data. We tested the relative roles of the colonization of the Socotra Archipelago and the ecological specialization in the evolution of the phenotypic diversity observed within the genus. We did not find evidence for the existence of an island effect in this radiation of geckos, since Socotran species do not present a notably different morphology, higher disparity, nor increased rates

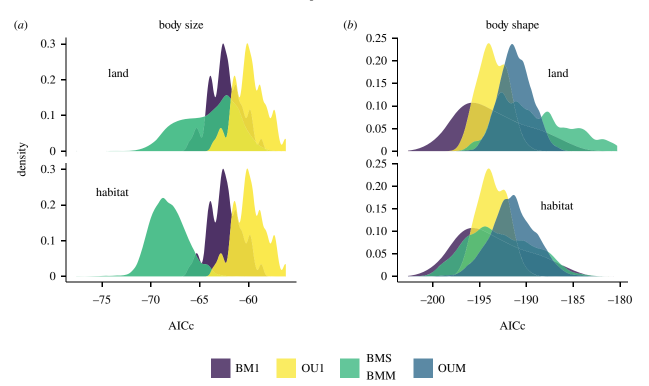


Figure 3. AICc distributions from the model fitting for (*a*) body size and (*b*) shape evolution of the genus *Pristurus*. These results correspond to model fitting on 100 stochastic character maps (insularity in the top panels and habitat use in the bottom) on 100 trees from the phylogenetic posterior distribution. BM1: Brownian motion single rate. OU1: Ornstein–Uhlenbeck single rate. BMS (OUwie)/BMM (mvMorph): Brownian motion multi-rate. OUM: Ornstein–Uhlenbeck multi-rate. For body size, the best supported model is a Brownian motion with rate heterogeneity across categories. For body shape, a single-rate Brownian motion model was the best fit, although there is an extensive overlap across all models. (Online version in colour.)

of morphological evolution, relative to species in the continent. On the contrary, ecological specialization emerges as a determinant factor in generating morphological diversity, with the colonization of ground habitats as the main driver of phenotypic divergence. Our results reveal a complex scenario in which different morphological traits interact with ecological characteristics of the species in different ways, suggesting a differential relevance of body size and shape proportions for the adaptation to specific habitats.

The tendency of island taxa to diverge in morphology compared to their continental relatives is a general pattern in terrestrial vertebrates, especially concerning body size [43,44]. In fact, recent studies on Afro-Arabian geckos colonizing the Socotra Archipelago found support for this island effect, particularly in the genus Hemidactylus [28,29]. Nevertheless, preliminary results on Pristurus geckos failed to find this phenomenon in this genus [28]. Here we corroborated and extended those preliminary results, incorporating the most complete phylogenetic and morphological sampling within Pristurus up to date. We did not find the predicted effects of the colonization of the Socotra Archipelago in phenotypic diversification. Even though one of the Socotran clades (the one including P. insignis, P. insignoides and P. sp. 12) has effectively undergone an increase in body size, we did not find significant differences in body size between the first nodes of each colonization event and their direct ancestral nodes (electronic supplementary material, figure S3). There is also no apparent divergence in body shape, with limb and head proportions of Socotran species being completely encompassed into the morphospace of continental species (figure 1a). Moreover, neither size nor shape disparities observed in the Socotra Archipelago are higher than those of the continental species. Finally, our analyses failed to find another expected outcome of island colonization, as is the increase in rates of phenotypic evolution. Consistent with our results of body size divergence, in which one of the Socotran clades are shown to have undergone a body size increase (electronic supplementary material, figure S3), we found heterogeneity in the rates of body size evolution, with some species evolving faster than others producing a bimodal distribution (figure 4a). Nevertheless, species in the continent show higher rates of body size evolution than all Socotran species (figure 4a). For body shape, our evolutionary model fitting showed no support for differences in shape rates between continental and Socotran species (BM1 was the best-fitting model; figure 3a). When extracting the rate values for shape from the multirate Brownian motion model, we found little or no difference between continental and Socotran taxa (figure 4b). The lack of a general island effect in Pristurus after the colonization of Socotra, opposed to other similar diversifications of geckos, may indicate the existence of different ecological contexts even in the same physical settings, which would imply different ecological opportunities in the same island [11]. Namely, some life-history traits may have limited niche expansion in Pristurus species in the Socotra Archipelago. For instance, the fact that the ground-dwelling clade of Pristurus is also the only one that has adopted partially nocturnal habits [30] may indicate a potentially related development of these two traits. This would explain that diurnal Socotran Pristurus have not colonized ground habitats and, therefore, their morphology has not changed substantially, while some nocturnal Downloaded from https://royalsocietypublishing.org/ on 13 March 2022

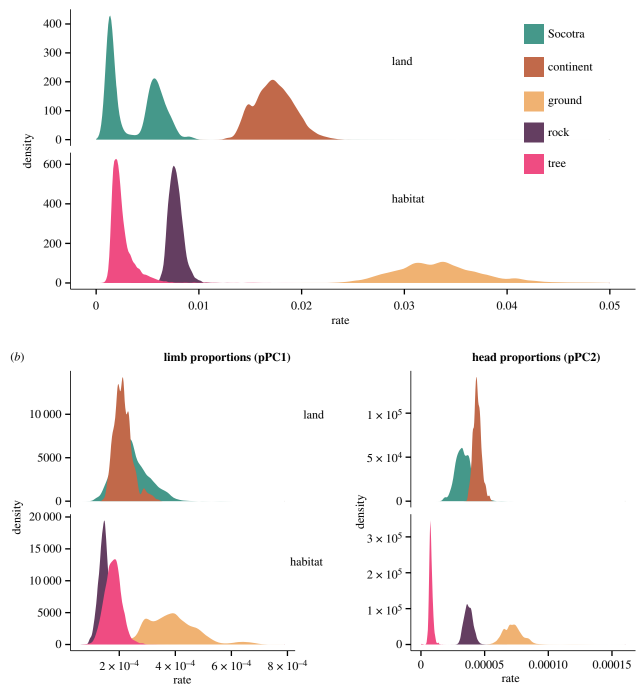


Figure 4. Rates of (a) body size and (b) body shape evolution in the genus Pristurus, extracted from multi-rate Brownian motion models fitted on a total of 10 000 character maps (100 stochastic character maps on 100 posterior trees) of Socotra versus continent and habitat use (ground, rock and tree). (Online version in colour.)

geckos that have colonized the Socotra Archipelago, like *Hemidactylus* or *Haemodracon*, show a marked phenotypic divergence [29,45]. This is consistent with results on global insular vertebrate communities suggesting that the prevalence of the island rule is subjected to system-specific ecological and environmental dynamics [44]. Furthermore, a recent study on the anole radiation in the Greater Antilles did not find evidence for an island effect, pointing instead to ecological opportunity and key innovations as the drivers of the adaptive radiation [46]. In *Pristurus*, one of the clades colonizing the Socotra Archipelago (*P. insignis, P. insignoides* and *P.* sp. 12) seems to have undergone to a certain extent an increase in body size coupled with increased evolutionary rates compared with the other Socotran *Pristurus*. This could be indicative of some kind of ecological release derived from

the island colonization but, in any case, the degree of evolutionary change of this Socotran clade is not comparable to that observed with the colonization of ground habitats.

Following that reasoning, ecological specialization gives us a much more nuanced insight into *Pristurus* phenotypic evolution. The relationship between habitat use and morphological traits is well recognized in lizards [3,47,48]. In fact, preceding observations on *Pristurus* geckos suggested that many morphological changes might be functionally associated with shifts in ecology and behaviour [30]. Our results are consistent with this notion and provide strong evidence that novel ecological opportunities produced high levels of phenotypic disparity associated with increased rates of trait evolution in some forms of *Pristurus*, particularly the species exploiting ground habitats. Even though ground-dwelling species do not show an extremely divergent

body shape relative to species inhabiting other habitats (i.e. rocky and arboreal habitats; figure 1a), they do comprise the largest and the smallest sizes of the genus (figure 1b) and show some extreme values of limb and head proportions (pPC1 and pPC2 respectively; figure 1a), as well as they occupy a very large portion of the genus' entire morphospace (figure 1a). This extreme variability within ground-dwelling species is reflected in our disparity results. While ground species do not present higher disparity in limb dimensions (pPC1; ratio ground/no-ground less than 1), they have more than twice as much disparity as all the rest of the species in body size and head proportions, with these ratios being highly significant compared to the null model generated from simulations of character evolution (figure 2). Furthermore, we found increased rates of body size evolution in ground-dwelling species, followed first by rocky and last by arboreal habitats (figure 4a). Although rate heterogeneity across habitat categories was not the best-fit model for body shape (figure 3b), the rate values extracted from the multi-rate Brownian motion model show a similar pattern, especially for head proportions, with highest rates in grounddwelling species (figure 4b). Taken together, these results point to the existence of a morphological response to the ecological context, especially in body size. This is consistent with the idea that the main driver of morphological divergence, even in an island colonization event, is habitat diversity [49,50]. If habitat heterogeneity in the Socotra Archipelago is lower than in continental Africa and Arabia (e.g. no particularly large gravel plains in Socotra), or if the access to those habitats is limited for Pristurus geckos in Socotra and not in the continent (e.g. due to the interplay of other ecological and historical factors), phenotypic evolution after island colonization would not be as extreme as expected under the island effect framework. This could imply a tight relationship between morphology and structural habitat, a pattern observed in other Arabian geckos such as Ptyodactylus or Asaccus, where niche conservatism is associated with a very conserved morphology [17,51-54]. This would be further supported by the fact that within ground habitats, species show a clear morphological segregation between 'hard' and 'soft' substrates, suggesting a particularly conspicuous specialization to the former (large bodies, long limbs and large heads in the hard-ground species: Pristurus carteri, Pristurus collaris, Pristurus ornithocephalus; electronic supplementary material, figure S4). In fact, these hard-ground species that have developed partially nocturnal habits have evolved into rough ecological analogues of some small diurnal ground-dwelling desert iguanians (e.g. the rock agamas of the genus Pseudotrapelus, the toad-headed agamas of the genus Phrynocephalus and the yellow-spotted agama of the genus Trapelus), with the long limbs favouring the movement in the newly colonized grounds and with large heads able to accommodate large eyes advantageous in those open habitats. Alternatively, the lack of an island effect in Socotran Pristurus might be explained by climatic divergences replacing ecomorphological differentiation, or by low morphological evolvability [28].

Another interesting result is the morphological affinities of arboreal species, especially in body size and head proportions, where they present intermediate values within a very restricted range (figure 1). Consistently, we found notably reduced evolutionary rates in body size and head proportions in these species relative to other habitats with the multi-rate models (figure 4). This might corroborate the idea of adaptive processes leading to a tight relationship between ecological traits and phenotype since this scenario is expected if a specific habitat constrains the morphology toward optimum values of body size and shape [14].

Ultimately, our results provide evidence of the determinant role of habitat specialization in phenotypic evolution. This has important implications for understanding the prevalence of the island effect in the context of differential ecological opportunity and, combined with previous results on other similar systems, shows the complex nature of the relationships between ecological mechanisms and morphology and their reliance on systemspecific dynamics. More detailed ecological and morphological data (e.g. dietary habits, geometric morphometrics of head shape) might help for a deeper understanding of the evolutionary dynamics of this and other groups of arid-adapted lizards.

Data accessibility. Electronic supplementary material, data and R scripts used for this study are available from the Dryad Digital Repository: https://doi.org/10.5061/dryad.xwdbrv1f6 [55]. The data are provided in electronic supplementary material [56].

Authors' contributions. H.T.-C.: conceptualization, data curation, formal analysis, methodology, visualization, writing—original draft, writing—review and editing; M.S.-R.: conceptualization, data curation, methodology, writing—original draft, writing—review and editing; I.M.: formal analysis, methodology, visualization, writing—review and editing; S.C.: conceptualization, funding acquisition, project administration, supervision, writing—review and editing. All authors gave final approval for publication and agreed to be held accountable for the work performed therein.

Competing interests. We declare we have no competing interests.

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References

- Baeckens S, Van Damme R. 2020 The island syndrome. *Curr. Biol.* **30**, R338–R339. (doi:10.1016/ j.cub.2020.03.029)
- Foster JB. 1964 Evolution of mammals on islands. *Nature* 202, 234–235. (doi:10.1038/ 202234a0)
- Losos J. 2009 Lizards in an evolutionary tree: ecology and adaptive radiation of anoles. Berkeley, CA: University of California Press.
- Losos JB, Ricklefs RE. 2009 Adaptation and diversification on islands. *Nature* 457, 830–836. (doi:10.1038/nature07893)
- Yoder JB *et al.* 2010 Ecological opportunity and the origin of adaptive radiations. *J. Evol. Biol.* 23, 1581–1596. (doi:10.1111/j.1420-9101.2010. 02029.x)
- 6. Schluter D. 2000 *The ecology of adaptive radiation*. New York, NY: Oxford University Press.

- Harmon LJ, Melville J, Larson A, Losos JB, Oakley T. 2008 The role of geography and ecological opportunity in the diversification of day geckos (*Phelsuma*). *Syst. Biol.* 57, 562–573. (doi:10.1080/ 10635150802304779)
- Whittaker RJ, Triantis KA, Ladle RJ. 2008 A general dynamic theory of oceanic island biogeography. *J. Biogeogr.* 35, 977–994. (doi:10.1111/j.1365-2699. 2008.01892.x)
- Lomolino MV. 1985 Body size of mammals on islands: the island rule reexamined. Am. Nat. 125, 310–316. (doi:10.1086/284343)
- Rundell RJ, Price TD. 2009 Adaptive radiation, nonadaptive radiation, ecological speciation and nonecological speciation. *Trends Ecol. Evol.* 24, 394–399. (doi:10.1016/j.tree.2009.02.007)
- Losos JB. 2010 Adaptive radiation, ecological opportunity, and evolutionary determinism. *Am. Nat.* **175**, 623–639. (doi:10.1086/652433)
- Mahler DL, Revell LJ, Glor RE, Losos JB. 2010 Ecological opportunity and the rate of morphological evolution in the diversification of Greater Antillean anoles. *Evolution* 64, 2731–2745. (doi:10.1111/j.1558-5646.2010.01026.x)
- Reynolds RG, Collar DC, Pasachnik SA, Niemiller ML, Puente-Rolón AR, Revell LJ. 2016 Ecological specialization and morphological diversification in Greater Antillean boas. *Evolution* **70**, 1882–1895. (doi:10.1111/evo.12987)
- Moen DS, Morlon H, Wiens JJ. 2016 Testing convergence versus history: convergence dominates phenotypic evolution for over 150 million years in frogs. *Syst. Biol.* 65, 146–160. (doi:10.1093/sysbio/ syv073)
- Gamble T, Greenbaum E, Jackman TR, Bauer AM. 2015 Into the light: diurnality has evolved multiple times in geckos. *Biol. J. Linnean Soc.* **115**, 896–910. (doi:10.1111/bij.12536)
- Metallinou M *et al.* 2012 Conquering the Sahara and Arabian deserts: systematics and biogeography of *Stenodactylus* geckos (Reptilia: Gekkonidae). *BMC Evol. Biol.* **12**, 258. (doi:10.1186/1471-2148-12-258)
- Metallinou M, Červenka J, Crochet PA, Kratochvíl L, Wilms T, Geniez P, Shobrake MY, Brito JC, Carranzaa S. 2015 Species on the rocks: systematics and biogeography of the rock-dwelling *Ptyodactylus* geckos (Squamata: Phyllodactylidae) in North Africa and Arabia. *Mol. Phylogenet. Evol.* **85**, 208–220. (doi:10.1016/j.ympev.2015.02.010)
- Šmíd J, Moravec J, Kratochvíl L, Nasher AK, Mazuch T, Gvoždík V, Carranza S. 2015 Multilocus phylogeny and taxonomic revision of the *Hemidactylus robustus* species group (Reptilia, Gekkonidae) with descriptions of three new species from Yemen and Ethiopia. *Syst. Biodivers.* 13, 346–368. (doi:10.1080/ 14772000.2014.996264)
- Šmíd J, Shobrak M, Wilms T, Joger U, Carranza S. 2017 Endemic diversification in the mountains: genetic, morphological, and geographical differentiation of the *Hemidactylus* geckos in southwestern Arabia. *Org. Divers. Evol.* 17, 267–285. (doi:10.1007/s13127-016-0293-3)

- Garcia-Porta J, Simó-Riudalbas M, Robinson M, Carranza S. 2017 Diversification in arid mountains: biogeography and cryptic diversity of *Pristurus rupestris rupestris* in Arabia. *J. Biogeogr.* 44, 1694–1704. (doi:10.1111/jbi.12929)
- Machado L, Salvi D, James Harris D, Brito JC, Crochet PA, Geniez P, Ahmadzadeh F, Carranzac S. 2021 Systematics, biogeography and evolution of the Saharo-Arabian naked-toed geckos genus *Tropiocolotes. Mol. Phylogenet. Evol.* **155**, 106969. (doi:10.1016/j.ympev.2020.106969)
- Carranza S, Arnold EN. 2012 A review of the geckos of the genus *Hemidactylus* (Squamata: Gekkonidae) from Oman based on morphology, mitochondrial and nuclear data, with descriptions of eight new species. *Zootaxa* **95**, 1–95. (doi:10.11646/zootaxa. 3378.1.1)
- Gómez-Díaz E, Sindaco R, Pupin F, Fasola M, Carranza S. 2012 Origin and in situ diversification in *Hemidactylus* geckos of the Socotra Archipelago. *Mol. Ecol.* 21, 4074–4092. (doi:10.1111/j.1365-294X.2012.05672.x)
- Šmíd J, Carranza S, Kratochvíl L, Gvoždík V, Nasher AK, Moravec J. 2013 Out of Arabia: a complex biogeographic history of multiple vicariance and dispersal events in the gecko genus *Hemidactylus* (Reptilia: Gekkonidae). *PLoS ONE* 8, e64018. (doi:10. 1371/journal.pone.0064018)
- Šmíd J, Moravec J, Kratochvíl L, Gvoždík V, Nasher AK, Busais SM, Wilms T, Shobrak MY, Carranza S. 2013 Two newly recognized species of *Hemidactylus* (Squamata, Gekkonidae) from the Arabian Peninsula and Sinai, Egypt. *ZooKeys* 25, 79–107.
- Šmíd J, Uvizl M, Shobrak M, AlGethami RH, Algethami AR, Alanazi AS, Alsubaie SD, Busais S, Carranza S. 2020 Swimming through the sands of the Sahara and Arabian deserts: phylogeny of sandfish skinks (Scincidae, *Scincus*) reveals a recent and rapid diversification. *Mol. Phylogenet. Evol.* 155, 107012. (doi:10.1016/j.ympev.2020.107012)
- Vasconcelos R, Carranza S. 2014 Systematics and biogeography of *Hemidactylus homoeolepis* Blanford, 1881 (Squamata: Gekkonidae), with the description of a new species from Arabia. *Zootaxa* 3835, 501–527. (doi:10.11646/zootaxa. 3835.4.4)
- Garcia-Porta J, Morales HE, Gómez-Díaz E, Sindaco R, Carranza S. 2016 Patterns of diversification in islands: a comparative study across three gecko genera in the Socotra Archipelago. *Mol. Phylogenet. Evol.* **98**, 288–299. (doi:10.1016/j.ympev.2016. 02.007)
- Garcia-Porta J, Šmíd J, Sol D, Fasola M, Carranza S. 2016 Testing the island effect on phenotypic diversification: insights from the *Hemidactylus* geckos of the Socotra Archipelago. *Sci. Rep.* 6, 23729. (doi:10.1038/srep23729)
- Arnold EN. 2009 Relationships, evolution and biogeography of Semaphore geckos, *Pristurus* (Squamata, Sphaerodactylidae) based on morphology. *Zootaxa* 21, 1–21. (doi:10.11646/ zootaxa.2060.1.1)

- Badiane A, Garcia-Porta J, Červenka J, Kratochvíl L, Sindaco R, Robinson MD *et al.* 2014 Phylogenetic relationships of Semaphore geckos (Squamata: Sphaerodactylidae: *Pristurus*) with an assessment of the taxonomy of *Pristurus rupestris. Zootaxa* 3835, 33–58. (doi:10.11646/zootaxa.3835.1.2)
- Tejero-Cicuéndez H, Patton AH, Caetano DS, Šmíd J, Harmon LJ, Carranza S. In press. Reconstructing squamate biogeography in Afro-Arabia reveals the influence of a complex and dynamic geologic past. *Syst. Biol.* syab025. (doi:10.1093/sysbio/syab025)
- Arnold EN. 1993 Historical changes in the ecology and behaviour of semaphore geckos (*Pristurus*, Gekkonidae) and their relatives. *J. Zool.* 229, 353–384. (doi:10.1111/j.1469-7998.1993.tb02642.x)
- Pennell MW, Eastman JM, Slater GJ, Brown JW, Uyeda JC, Fitzjohn RG, Alfaro ME, Harmon LJ. 2014 Geiger v2.0: an expanded suite of methods for fitting macroevolutionary models to phylogenetic trees. *Bioinformatics* **30**, 2216–2218. (doi:10.1093/ bioinformatics/btu181)
- R Core Team. 2019 R: a language and environment for statistical computing. Vienna, Austria: R Foundation for Statistical Computing. See https:// www.r-project.org/.
- Akaike H. 1973 Information theory and an extension of the maximum likelihood principle. In *Second international symposium on information theory* (eds B Petrov, F Csaki), pp. 267–281. Budapest, Hungary: Akademia Kiado.
- Revell LJ. 2012 phytools: an R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.* 3, 217–223. (doi:10. 1111/j.2041-210X.2011.00169.x)
- Beaulieu JM, O'Meara B. 2021 OUwie: analysis of evolutionary rates in an OU framework. R package, version 2.6.
- Clavel J, Escarguel G, Merceron G. 2015 mvMORPH: an R package for fitting multivariate evolutionary models to morphometric data. *Methods Ecol. Evol.* 6, 1311–1319. (doi:10.1111/2041-210X.12420)
- Wickham H. 2011 ggplot2. Wiley Interdiscip. Rev.: Comput. Stat. 3, 180–185. (doi:10.1002/wics.147)
- 41. Pedersen T L. 2020 Patchwork: the composer of plots. R package version 1.1.1.
- Wilke CO. 2020 cowplot: streamlined plot theme and plot annotations for 'ggplot2'. R package version 1.1.
- Lomolino MV. 2005 Body size evolution in insular vertebrates: generality of the island rule.
 J. Biogeogr. 32, 1683–1699. (doi:10.1111/j.1365-2699.2005.01314.x)
- Benítez-López A, Santini L, Gallego-Zamorano J, Milá B, Walkden P, Huijbregts MAJ, Tobias JA. 2021 The island rule explains consistent patterns of body size evolution in terrestrial vertebrates. *Nat. Ecol. Evol.* 5, 768–786. (doi:10.1038/s41559-021-01426-y)
- Tamar K, Simó-Riudalbas M, Garcia-Porta J, Santos X, Llorente G, Vasconcelos R, Carranza S. 2019 An integrative study of island diversification: insights from the endemic *Haemodracon* geckos of the Socotra Archipelago.

Mol. Phylogenet. Evol. **133**, 166–175. (doi:10.1016/ j.ympev.2019.01.009)

- Burress ED, Muñoz MM. In press. Ecological opportunity from innovation, not islands, drove the anole lizard adaptive radiation. *Syst. Biol.* syab031. (doi:10.1093/sysbio/syab031)
- Goodman BA, Miles DB, Schwarzkopf L. 2008 Life on the rocks: habitat use drives morphological and performance evolution in lizards. *Ecology* 89, 3462–3471. (doi:10.1890/07-2093.1)
- Higham TE, Russell AP. 2010 Divergence in locomotor performance, ecology, and morphology between two sympatric sister species of desertdwelling gecko. *Biol. J. Linnean Soc.* **101**, 860–869. (doi:10.1111/j.1095-8312.2010.01539.x)
- Lack D. 1976 Island biology illustrated by the land birds of Jamaica. Berkeley, CA: University of California Press.
- 50. Losos JB, Parent CE. 2009 The speciation-area relationship. In *The theory of island*

biogeography revisited (eds JB Losos, RE Ricklefs), pp. 415–438. Princeton, NJ: Princeton University Press.

- Carranza S, Simó-Riudalbas M, Jayasinghe S, Wilms T, Els J. 2016 Microendemicity in the northern Hajar Mountains of Oman and the United Arab Emirates with the description of two new species of geckos of the genus *Asaccus* (Squamata: Phyllodactylidae). *PeerJ* 4, e2371. (doi:10.7717/ peeri.2371)
- Simó-Riudalbas M, Metallinou M, de Pous P, Els J, Jayasinghe S, Péntek-Zakar E, Wilms T, Al-Saadi S, Carranza S. 2017 Cryptic diversity in *Ptyodactylus* (Reptilia: Gekkonidae) from the northern Hajar Mountains of Oman and the United Arab Emirates uncovered by an integrative taxonomic approach. *PloS ONE* **12**, e0180397. (doi:10.1371/journal.pone. 0180397)
- 53. Simó-Riudalbas M, Tarroso P, Papenfuss T, Al-Sariri T, Carranza S. 2018 Systematics, biogeography and

evolution of *Asaccus gallagheri* (Squamata, Phyllodactylidae) with the description of a new endemic species from Oman. *Syst. Biodivers.* **16**, 323–339. (doi:10.1080/14772000.2017.1403496)

- Tamar K, Mitsi P, Carranza S. 2019 Cryptic diversity revealed in the leaf-toed gecko Asaccus montanus (Squamata, Phyllodactylidae) from the Hajar Mountains of Arabia. J. Zool. Syst. Evol. Res. 57, 369–382. (doi:10.1111/jzs.12258)
- Tejero-Cicuéndez H, Simó-Riudalbas M, Menéndez I, Carranza S. 2021 Ecological specialization, rather than the island effect, explains morphological diversification in an ancient radiation of geckos. Dryad Digital Repository. (doi:10.5061/dryad. xwdbrv1f6)
- 56. Tejero-Cicuéndez H, Simó-Riudalbas M, Menéndez I, Carranza S. 2021 Ecological specialization, rather than the island effect, explains morphological diversification in an ancient radiation of geckos. FigShare.

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