- Evolution along allometric lines of least resistance: Morphological
- differentiation in *Pristurus* geckos
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- 7 14 August, 2023
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Abstract

Species living in distinct habitats often experience unique ecological selective pressures, which can drive phenotypic divergence. However, how ecophenotypic patterns are affected by allometric trends 18 and trait integration levels is less well understood. Here we evaluate the role of allometry in shaping 19 body size and body form diversity in *Pristurus* geckos utilizing differing habitats. We found that patterns of allometry and integration in body form were distinct in species with different habitat 21 preferences, with ground-dwelling *Pristurus* displaying the most divergent allometric trend and high 22 levels of integration. There was also strong concordance between static intraspecific allometry across 23 individuals and evolutionary allometry among species, revealing that differences in body form among individuals were predictive of evolutionary changes across the phylogeny at macroevolutionary 25 scales. This suggested that phenotypic evolution occurred along allometric lines of least resistance, with allometric trajectories imposing a strong influence on the magnitude and direction of size and 27 shape changes across the phylogeny. When viewed in phylomorphospace, the largest rock-dwelling 28 species were most similar to the smallest ground-dwelling species, and vice versa. Thus, in *Pristurus*, 29 phenotypic evolution along the differing habitat-based allometric trajectories resulted in similar body forms at differing body sizes in distinct ecological habitats.

2 1. Introduction

Understanding how phenotypic diversity evolves, and elucidating the forces that generate and maintain this diversity, are major goals in evolutionary biology. Because adaptive evolution is the product of natural selection, changes in ecological selection pressures are expected to affect the evolutionary trajectory of phenotypic traits that facilitate an organism's survival and reproduction in their habitat. The theory of natural selection predicts that differing habitats will exert unique ecological selection pressures on organisms, resulting in associations between ecological and phenotypic traits. Indeed, species inhabiting differing habitats often display functional, behavioral, or phenotypic differences, that have presumably been the result of adaptive diversification in their respective ecological contexts (Collar et al. 2010; Kaliontzopoulou et al. 2015; Price et al. 2015; Martinez et al. 2021; Kolmann et al. 2022).

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One possible evolutionary outcome of ecological specialization is that organisms inhabiting similar environments display common phenotypic characteristics. When such patterns occur repeatedly (Losos 1992; Schluter and McPhail 1992), this convergent evolution is treated as strong evidence of adaptation. Indeed the ecomorphological paradigm (Arnold 1983) is predicated, in part, on such cases, which emphasize the strong association between the phenotypic traits that organisms display (morphological, behavioral, or physiological) and the ecological characteristics of their habitat that mediate organismal performance. In vertebrates, ecomorphological trends have been well studied in numerous taxonomic groups, and include the emblematic 'ecomorphs' of Caribbean Anolis lizards that exploit different microhabitats (Losos 1992, 2009; Mahler et al. 2013), differential beak morphology in species of Darwin's finches (Schluter and Grant 1984; Grant and Grant 2006; Reaney et al. 2020), the recurring phenotypes of African lake cichlids across ecological regimes (Albertson and Kocher 2001; Urban et al. 2022), and the distinct body forms of freshwater fishes in benthic and limnetic habitats (Jastrebski and Robinson 2004; Berner et al. 2008; Stuart et al. 2017), among others.

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9 While the patterns of morphological differences in distinct ecological contexts have been well

documented, less-well understood is how ecomorphological differentiation has been influenced by trait covariation associated with body size differences (i.e., allometry). The study of size-related changes in anatomical traits has a long history in evolutionary biology (Huxley 1932; Jolicoeur 1963; Gould 1966; Klingenberg 1996; Zelditch and Swiderski 2022). One reason for this is that nearly all traits covary strongly with overall body size (Jolicoeur 1963; Gould 1966; Bookstein 2022), and as such, the effects of allometry on patterns of phenotypic diversity are expected to be considerable. Further, allometric patterns manifest widely are widely prominent across differing levels of biological organization: from variation across individuals at differing stages or ages (i.e., ontogenetic allometry), to variation across individuals in a single ontogenetic stage within a population or species (i.e., static allometry), to variation across species of differing sizes, as described by evolutionary allometry (see Cock 1966; Klingenberg and Zimmermann 1992). Indeed, across vertebrates, numerous studies have shown that a sizeable proportion of overall phenotypic variation is related to interspecific differences in body size (Pyron and Burbrink 2009: Piras et al. 2010; Cardini and Polly 2013; e.g., Sherratt et al. 2014; Cardini et al. 2015; Bright et al. 2016; Bardua et al. 2021; Felice et al. 2021; Zelditch and Swiderski 2022). When viewed from this perspective, patterns of ontogenetic both intraspecific and evolutionary allometry are thought to play a decisive role in shaping patterns of phenotypic diversification across the tree of life.

However, allometry can also act as a restraining force on evolution by limiting the breadth of phenotypes that can be realized (Bright et al. 2019). This occurs because trait correlations influence the degree to which phenotypic variation is exposed to selection (Wagner and Altenberg 1996). Thus, the integration among traits can constrain phenotypic change in certain directions, or enhance variation along other phenotypic axes (Schluter 1996; Wagner and Altenberg 1996; Wagner and Zhang 2011; Klingenberg and Marugán-Lobón 2013; Goswami et al. 2014, 2016; Felice et al. 2018; Navalón et al. 2020). Further, because nearly all linear traits covary strongly with overall body size, allometric trends could be considered the quintessential expression of phenotypic integration (Zelditch and Swiderski 2022; Bookstein 2022). Thus, when evaluating ecophenotypic differences among taxa, it is important to consider how allometric trends of trait covariation influence such patterns (e.g., Esquerré et al. 2017; Patterson et al. 2022; Chatterji et al. 2022).

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The Afro-Arabian geckos in the genus *Pristurus* afford the opportunity to elucidate the interdigitating effects of allometry and habitat specialization on clade-level patterns of phenotypic 91 diversity. From an evolutionary perspective, *Pristurus* geckos are an ideal system to investigate the role of different factors in evolutionary history. They are found in both insular and continental 93 settings, which are known to impose differential ecological selection pressures resulting in distinct evolutionary trajectories of species and clades (Losos and Ricklefs 2009). They are also distributed in the contact zone between Africa and Eurasia, a region of high biogeographic interest which has been the epicenter of major faunal interchanges and complex geologic and environmental processes (Kappelman et al. 2003; Tejero-Cicuéndez et al. 2022). The study of evolutionary dynamics in *Pristurus* and other Afro-Arabian taxa is also important biologically, gg as our understanding of the dynamics of biological systems in remote arid regions is generally 100 neglected and understudied (Durant et al. 2012). Furthermore, prior work on this system 101 (Tejero-Cicuéndez et al. 2021a) has revealed that the colonization of ground habitats has been 102 a trigger of morphological change, specifically reflected in an increase in body size and shape 103 disparity. Interestingly, some ground-dwelling species are among the largest of the genus and 104 also show increased relative head sizes and limb proportions, while some other species with 105 this ecological specialization have evolved to be among the smallest of the group. Additionally, 106 among the species exploiting rocky habitats (the most common ecological feature in *Pristurus*), 107 there are also species with both considerably large and small body sizes (Tejero-Cicuéndez et al. 2021a). What remains unexplored, however, is how the evolution of body form is related to dif-109 ferences in body size and whether habitat specialization has an impact in this shape-size relationship. 110

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In this study, we employed a combination of multivariate morphometric and phylogenetic comparative analyses to interrogate macroevolutionary patterns of evolutionary allometry in *Pristurus* geckos of Afro-Arabia. Using phenotypic, phylogenetic, and ecological data, we first characterized allometric trends in body form in the group, to discern the extent to which evolutionary allometric trends across the phylogeny aligned with habitat-based static intraspecific allometry for species occupying distinct ecological regimes. We then examined changes in allometric trends across the phylogeny,

and linked these patterns to overall phenotypic integration, diversification in morphospace, and
habitat utilization among taxa. Our analyses reveal that patterns of evolutionary allometry across
species align with allometric trends among individuals, and that differing habitat-based allometric
trajectories have resulted in similar body forms at differing body sizes in distinct ecological regimes.
Thus, patterns of phenotypic diversification in *Pristurus* are the outcome of an interplay between
ecological specialization and size-form changes evolving along habitat-specific allometric trajectories
in the group.

2. Materials and Methods

126 (a) Data

We used a combination of phenotypic, phylogenetic, and ecological data to characterize and evaluate 127 intra- and interspecific allometric trends in *Pristurus*. The data utilized here were obtained from 128 our prior work on this system (Tejero-Cicuéndez et al. 2021a, 2022), and are briefly described 129 here. First, we used a time-calibrated molecular phylogeny of squamates that included all members 130 of the genus *Pristurus*, including several currently undescribed taxa. The tree was estimated in 131 a Bayesian framework, using five mitochondrial markers, six nuclear markers, and 21 calibration 132 points (Tejero-Cicuéndez et al. 2022). Next, we categorized each species as belonging to one of three 133 ecological habitat groups (ground-dwelling, rock-dwelling, or tree-dwelling). Habitat designations 134 were based on substrate preferences and habitat use as found through extensive field observations 135 described in the primary literature (Arnold 1993; Arnold 2009, and references therein). Finally, 136 we obtained a phenotypic dataset containing body size (snout-vent length: SVL) and eight linear 137 measurements (Figure 1) that described overall body form: trunk length (TL), head length (HL), 138 head width (HW), head height (HH), humerus length (Lhu), ulna length (Lun), femur length (Lfe), and tibia length (Ltb) (Tejero-Cicuéndez et al. 2021a). We restricted our study to those 140 species represented by nine or more individuals; resulting in a dataset of 687 individuals from 25 141 species (invidivuals per species: mean = 27; min = 9, max = 56). All specimens used in this 142 study were adults, and thus patterns of ontogenetic allometry could not be explored. Species in 143 the phenotypic dataset were then matched to the phylogeny, which was subsequently pruned to 144 the final topology. All measurements were log-transformed prior to statistical analyses. Additional 145

details regarding data collection and formal descriptions of each linear measurement may be found in the original sources (Tejero-Cicuéndez et al. 2021a, 2022). The data are available on DRYAD: https://doi.org/10.5061/dryad.xwdbrv1f6 (Tejero-Cicuéndez et al. 2021b).

149 (b) Statistical and Comparative Analyses

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We conducted a series of analyses to interrogate allometric trends, patterns of integration, and 150 macroevolutionary changes in allometry, relative to differentiation in body form. First, we 151 characterized evolutionary allometry in the genus by performing a phylogenetic multivariate 152 regression of body form on body size (i.e., SVL), using the species means as data. We then evaluated 153 patterns of static instraspecific allometry among individuals using a pooled within-species regression 154 (sensu Klingenberg 2016). Here a pooled within-species dataset was generated by obtaining 155 residuals for all individuals relative to their respective species means, which were then pooled across 156 species. This dataset was then used in a multivariate regression to obtain an overall estimate of 157 static intraspecific allometry among individuals. By first removing species-specific differences, this procedure partials out trends of evolutionary allometry from the data, enabling patterns of static 159 intraspecific and evolutionary allometry to be disentangled —(note that because juvenile specimens 160 were not available, it was not possible to disentangle the ontogenetic and static components of 161 allometric trends. Thus we refer to this level as 'intraspecific' allometry to be conservative). From 162 both the species-level and the individual-level analyses, we obtained the set of regression coefficients, 163 which respectively described the trajectories of evolutionary and static intraspecific allometry in 164 morphospace. We then calculated the difference in their direction in morphospace to discern the 165 extent to which patterns of static intraspecific allometry at the individual level were concordant 166 with evolutionary allometric trends across species. 167

Next, we used the pooled within-species dataset to determine whether trends in static intraspecific allometry differed across habitat groups. This was accomplished by performing a multivariate analysis of covariance, with body size (SVL), habitat, and $SVL \times habitat$ as model effects. Significance of model effects was evaluated using 999 iterations of a permutation procedure, where residuals from a reduced model were randomly permuted in each permutation (RRPP), model

statistics were recalculated, and used to generate empirical null sampling distributions to evaluate 174 the observed test statistics (following Freedman and Lane 1983; Collyer and Adams 2007; Collyer et al. 2015). In this analysis, no variation was attributable to the habitat effect, as the pooled-within 176 species data are mean-centered for both the dependent and independent variables. However, any 177 differences in multivariate allometric slopes among habitats will be discernable, and revealed by a 178 significant $SVL \times habitat$ interaction. To evaluate this possibility, we compared the direction of 179 multivariate allometric vectors for each habitat group to one another, and to a vector representing 180 multivariate isometry, by calculating pairwise angular differences in their direction in morphospace, 181 and evaluating these relative to empirical sampling distributions obtained through RRPP (Collyer 182 and Adams 2007; Adams and Collyer 2009; Collyer and Adams 2013). Here, residuals were obtained 183 from a common isometry reduced model, whose common slope component described a pattern of 184 multivariate isometry, and whose intercepts allowed for differences in least-squares means among 185 groups. Patterns of multivariate allometry relative to body size were visualized via regression 186 scores (Drake and Klingenberg 2008) and predicted lines (Adams and Nistri 2010), based on the 187 coefficients and fitted values from the linear model described above. 188

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Additionally, because allometry describes the extent to which traits covary with body size and with 190 each other (i.e., integration), we conducted an analysis of integration. Here we characterized the 191 extent of morphological integration in body form for individuals within each habitat group using the 192 pooled within-species dataset, and by summarizing the dispersion of eigenvalues of their respective 193 trait covariance matrix (Pavlicev et al. 2009). This measure (V_{rel}) was subsequently converted to 194 an effect size (a Z-score), which quantified the strength of morphological integration (Conaway 195 and Adams 2022). We then performed a series of two-sample tests to compare the strength of 196 morphological integration across habitat groups, following the procedures of Conaway and Adams 197 (2022). Additionally and for comparison, we repeated these analyses on the set of size-standardized 198 trait data, found as a set of shape ratios (Mosimann 1970) where each trait was divided by body 199 size (Supplementary Material). 200

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To determine the extent to which static intraspecific and evolutionary allometry were concordant,

we evaluated the directions in morphospace of both the evolutionary (species-level) and static intraspecific (habitat-based) allometric trends. Specifically, we obtained the set of regression coefficients from both the phylogenetic multivariate regression and the multivariate analysis of covariance analyses above, and calculated the angular difference in direction between the evolutionary allometry trajectory and the static intraspecific allometry trend for each habitat group. The observed angles were then statistically evaluated relative to empirical sampling distributions obtained through permutation (RRPP), based on the common isometry model described above.

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Next, to discern how allometric trends resulted in the evolution of distinct body forms, we examined 211 changes in relative body form across the phylogeny. Here we treated the head dimensions and limb 212 dimensions separately, as allometric trends could potentially differ between these body regions 213 due to differential functional or selective constraints (Kaliontzopoulou et al. 2010). Because 214 both the head and limb data were multivariate, we used regression scores (sensu Drake and 215 Klingenberg 2008) of a multivariate regression of head traits versus SVL and limb traits versus 216 SVL to represent the allometric trends in each dataset. We then measured the mean residuals 217 of each species to the inferred allometric trend, which described the extent to which head and 218 limb proportions of species were greater or smaller than expected for their body size. The 219 species residuals were then mapped on the phylogeny of *Pristurus* using a Brownian motion 220 model of evolution, to qualitatively evaluate shifts in head and limb proportionality across the 221 phylogeny for the group. Similarly, within-species patterns of static-patterns of intraspecific allome-222 try were visualized by plotting regression scores versus SVL for both head and limb traits separately. 223

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Finally, to relate within-species allometric trends with patterns of phenotypic diversification in the group, we generated a phylomorphospace (sensu Sidlauskas 2008), based on a principal component analyses (PCA) of the size-standardized species means (i.e., relative body proportions) obtained from a phylogenetic regression(see Tejero-Cicuéndez et al. 2021a). non-phylogenetic regression. Here, phenotypic similarities among species, relative to their phylogenetic relationships and habitat affiliations, were observed. A similar phylomorphospace was constructed for species means with size-standardized species means obtained from a phylogenetic regression, and another one with

species means not corrected for body size, and the. The phenotypic disparity among species 232 means in each habitat was calculated and subsequently compared (Supplementary Material). 233 Additionally, anatomical changes associated with allometric trends across taxa were visually depicted 234 via representative specimens from the largest and smallest ground-dwelling species (scaled to unit 235 size), and specimens from a large and small rock-dwelling species, to aid in describing these allometric trends. All analyses were conducted in R 4.2.1 (R Core Team 2022), using RRPP 237 version 1.3.2 (Collyer and Adams 2018; Collyer and Adams 2022) and geomorph 4.0.5 (Baken 238 et al. 2021a) for statistical analyses and the tidyverse version 1.3.0 (Wickham et al. 2019), 239 phytools version 0.7-77 (Revell 2012), and a modified version of the function ggphylomorpho 240 [https://github.com/wabarr/ggphylomorpho] for data manipulation and visualization, as well as 241 scripts written by the authors (Supplementary Material). 242

3. Results

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Using phylogenetic regression, we found significant evolutionary allometry in body form across species ($N_{sp} = 25$; F = 217.9; Z = 5.53; P < 0.001). Likewise, when static intraspecific allometry in body form was examined across individuals, a similar pattern was observed ($N_{ind} = 687$; F = 1176.9; Z = 8.24; P < 0.001). Further, the vectors of regression coefficients between the two analyses were oriented in a similar direction and were nearly parallel in morphospace ($\theta = 5.64^{\circ}$; Table 1). This revealed that the pattern of multivariate allometry across individuals was largely concordant with macroevolutionary trends of interspecific static allometry among species of *Pristurus* across the phylogeny.

Our analyses also exposed significant differences in the allometry of body form among *Pristurus* utilizing distinct habitats (Table Tables 1 and 2). Further, pairwise comparisons of multivariate allometric vectors revealed that patterns of static-intraspecific allometry in each habitat differed significantly from isometry, indicating the presence of multivariate allometry in each (Table 23).

Additionally, comparisons identified that ground-dwelling *Pristurus* displayed the most distinct allometric trend as compared with *Pristurus* occupying both the rock and tree habitats (Table 23; Figure 2). Here, regression coefficients of each trait *versus* size (Supplementary Material) revealed

that ground-dwelling *Pristurus* exhibited higher coefficients for head traits as compared with 260 rock-dwelling and tree-dwelling taxa ($\beta_{ground} > \beta_{rock}$; $\beta_{ground} > \beta_{tree}$). By contrast, coefficients 261 for limb traits were somewhat smaller for ground-dwelling Pristurus as compared with other 262 taxa ($\beta_{ground} < \beta_{rock}$; $\beta_{ground} < \beta_{tree}$). Thus, these findings implied that within species, larger 263 individuals of ground-dwelling *Pristurus* displayed proportionately larger heads and slightly smaller limbs as compared with large individuals in taxa utilizing other habitat types. Visualizations 265 of the allometric trends (Figure 2) confirmed these statistical findings, and indicated that the 266 allometric trajectory in ground-dwelling *Pristurus* was more extreme as compared with either rock-267 or tree-dwelling *Pristurus*. 268

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Examination of patterns of trait covariation for the pooled within-species data revealed strong 270 levels of morphological integration in the ground and tree ecotypes, with lower levels of integration 271 displayed in the rock habitat. Subsequent two-sample tests revealed that the strength of morphological integration was significantly greater in both ground-dwelling and tree-dwelling 273 $Pristurus \ \ {\rm than \ in \ those \ utilizing \ rock} \ (Z_{ground-rock} \ = \ 6.05; \ P \ << \ 0.001; \ Z_{tree-rock} \ = \ 4.07;$ 274 $P \ll 0.001$). Levels of morphological integration did not differ between ground and tree-dwelling 275 Pristurus ($Z_{tree-rock} = 0.38$; P = 0.702). Finally, when body size was taken into account, levels of 276 integration dropped considerably, though the overall pattern and differences among habitat groups 277 remained the same (Supplementary Material). 278

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Comparisons of evolutionary allometry with static intraspecific allometry in each habitat revealed 280 substantial concordance between allometric trends across these levels. Here, vectors of regression 281 coefficients representing static-intraspecific allometry within habitat groups were oriented in very 282 similar directions with the regression vector representing evolutionary allometry, with small pairwise 283 angles between them $(\theta:5.8^{\circ} \rightarrow 7.2^{\circ}5.8 < \theta < 7.2)$. Subsequent permutation tests indicated no 284 differences in direction between the regression vector representing evolutionary allometry and the 285 static intraspecific allometry vectors for *Pristurus* in both the ground or tree habitats, indicating 286 strong congruence between them (Table 34). By contrast, rock-dwelling *Pristurus* differed most in 287 their static intraspecific allometry trend relative to patterns of evolutionary allometry. Notably, 288

static intraspecific allometry in ground-dwelling *Pristurus* was most similar to trends of evolutionary allometry, displaying the smallest angular difference in direction when compared to evolutionary allometry. Overall, these findings implied that phenotypic evolution across species aligned closely with directions of allometric variation within habitat groups at the individual level, describing a trend where larger individuals – and larger ground-dwelling species – exhibited disproportionately larger heads and limbs, while smaller individuals – and smaller ground-dwelling species – displayed disproportionately smaller heads and limbs.

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Mapping the residuals of phylogenetic regression onto the phylogeny showed that large ground-297 dwelling species displayed greater head proportions than large rock-dwelling species, who exhibited 298 smaller heads relative to body size (Figure 3A). Conversely, the opposite pattern was observed 290 when comparing small species utilizing these habitats: ground-dwelling species showed small 300 relative head proportions while rock-dwelling species displayed generally larger head proportions. In 301 contrast, limb shape showed more variable patterns. Although all large ground-dwelling species 302 consistently displayed large relative limb proportions, large rock-dwelling species were more variable 303 in this trait, with P. insignis exhibiting large and P. insignoides small limb proportions. For 304 small species, shifts in relative limb proportions seemed more independent of habitat utilization, 305 since there were differences in limb residuals both within rock- and ground-dwelling species 306 (Figure 3B). Likeweise, static Likewise, intraspecific allometry trends within species revealed that 307 ground-dwelling species generally displayed steeper allometric patterns in head proportions as compared with rock-dwelling species (Figure 4). Overall, there was general concordance across taxa 309 in terms of trends of multivariate allometry, affirming that the association between evolutionary 310 allometry and habitat-based static intraspecific allometry was robust. 311

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Viewing differentiation in *Pristurus*' relative body proportions in phylomorphospace (Figure 5)
revealed a broad overlap among habitat groups in the first few dimensions, though arboreal
(tree-dwelling) species were somewhat more constrained in morphospace. Rock-dwelling species
occupied a slightly larger region of morphospace as compared with the other groups, though this
pattern was not statistically significant (Supplementary Material). Intriguingly, when viewed

in relation to body size, large *Pristurus* species were not localized to a particular region of morphospace, nor were smaller species. Instead, the largest rock-dwelling species were found in close proximity to the smallest ground-dwelling species, indicating that they were similar in relative body proportions. Likewise, the smallest rock-dwelling species were found close to large ground-dwelling species in morphospace, indicating they displayed similar body proportions as well.

These results did not change when the phylomorphospace was based on size-standardized species means obtained from a phylogenetic regression (Supplementary Material).

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Finally, when representative specimens were scaled to a similar body size (Figure 6), the anatomical 326 consequences of differences in allometric trends on body form became apparent. Here, larger 327 ground-dwelling *Pristurus* species displayed disproportionately larger heads and limbs as compared 328 with large *Pristurus* species utilizing other habitat types. Conversely, smaller rock-dwelling species 329 were found to have disproportionately larger heads and limbs than smaller ground-dwelling species. 330 These patterns corresponded closely with those identified in morphospace (Figure 5), where large 331 ground-dwelling species were similar in body form to small rock-dwelling species, while small 332 ground-dwelling species were similar in body form to large rock-dwelling species (Figure 6). Thus, 333 synthesizing the patterns revealed in the phylomorphospace with those from the other analyses 334 revealed that the same body proportions could be obtained in different ways, as determined by 335 subtle differences in allometric slope across habitats, combined with body size differences. As such, 336 species with similar body proportions displayed differing overall size, were found in distinct habitats, and exhibited different allometric trends. 338

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4. Discussion

Elucidating the selective forces that generate patterns of phenotypic diversity is a major goal in evolutionary biology. For species that utilize distinct habitats, disentangling the causes of phenotypic differentiation across those habitats is essential for our understanding of how natural selection operates and how evolution proceeds. In this study, we evaluated the role of potential drivers of body form differentiation in the geckos of the genus *Pristurus*. To this end, we compared allometric trends and levels of integration among *Pristurus* occupying distinct habitats, interrogated
allometric patterns at both the static intraspecific and evolutionary levels, and related these trends
to diversification in body form. Our findings have several important implications for how ecological
specialization, phenotypic integration, and body form evolution along allometric trajectories relate
to patterns of phenotypic diversity generally, and the evolution of phenotypic diversification in *Pristurus* in particular.

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First, our analyses revealed that patterns of allometry in body form and morphological integration 353 are relatively distinct in ground-dwelling *Pristurus* lizards, as compared with *Pristurus* occupying 354 other habitats. Specifically, we found that multivariate vectors of regression coefficients differed 355 significantly from what was expected under isometry (Table 23) for taxa utilizing all habitat 356 types (ground, rock, tree), indicating that in *Pristurus*, allometric scaling patterns predominate. 357 Further, our interrogation of allometric trends revealed differences between habitat types, where 358 ground-dwelling *Pristurus* displayed steeper allometric slopes for head traits as compared with 359 rock and tree-dwelling taxa. Biologically, these patterns revealed that not only does shape differ 360 between large and small *Pristurus*, but this pattern also differs across habitat types. Specifically, 361 large ground-dwelling *Pristurus* present disproportionately larger heads relative to large individuals 362 in other habitats, while small ground-dwelling *Pristurus* exhibit disproportionately smaller 363 heads (Figure 3). These findings are consistent with previous work at the macroevolutionary 364 level (Tejero-Cicuéndez et al. 2021a), where large ground species were also found to display disproportionately large heads. 366

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Second, our findings revealed that, within species, rock-dwelling *Pristurus* show a converse pattern,
where smaller individuals displayed relatively larger heads, while larger individuals have smaller
heads relative to their body size. These allometric patterns also corresponded with findings at
macroevolutionary scales (Tejero-Cicuéndez et al. 2021a), where similar patterns at the species
level were observed. Regarding relative limb proportions, we found a high variability among
small rock-dwelling species rather than a common pattern (Figure 3B). Indeed, earlier work in
the subclade comprising several of these species (the *P. rupestris* species complex) found two

well-differentiated phenotypes in populations of these lineages segregated by elevation (Garcia-Porta et al. 2017). These two ecotypes, defined as 'slender' and 'robust', differed in their head and limb characteristics. Our work is consistent with this, and extends these patterns to the allometric 377 realm. Tejero-Cicuéndez et al. (2021a) also performed habitat ancestral estimation, finding that the 378 rock habitat was the most likely ancestral condition in the group, with subsequent colonization by 379 Pristurus of ground habitats. When patterns of allometry are viewed through this lens, it suggests 380 the hypothesis that habitat shifts from rock-dwelling to ground-dwelling incurred a concomitant 381 evolutionary shift in allometric trajectories as well (Adams and Nistri 2010). Indeed, our analyses 382 are consistent with this hypothesis, as allometric trends are inferred to be more rock-like towards 383 the root of the *Pristurus* phylogeny are inferred to be more similar to those found in extant species 384 living in rocky habitats (Figure 3), with subsequent shifts along branches leading to ground-dwelling 385 species. This further suggests species exploiting different habitats (although it must be noted that all extant ground species have a single origin in the phylogeny and consitute a monophyletic 387 group). This might further indicate that the segregation in body size and shape through differential 388 allometric relationships across habitats responds to adaptive dynamics concerning the colonization 389 of new habitats, even though the fact that all ground species belong to the same clade hinders our 390 ability to draw stronger conclusions about adaptive dynamics involving the colonization of ground 391 habitats. Thus, in *Pristurus*, there is support for our results are consistent with the hypothesis 392 that colonization of ground habitats has been a trigger for morphological change (Tejero-Cicuéndez 393 et al. 2021a), as there appears to be a link between shifts in allometric trajectories as a result of 394 habitat-induced selection, and differential patterns of body form observed across taxa. Similar 395 patterns have been observed in other taxa, where differences in allometric trajectories are associated 396 with ecological differences across species (Esquerré et al. 2017; Patterson et al. 2022; Chatterji 397 et al. 2022). More broadly, these findings are consistent with prior discoveries in other lizards, 398 where the differential selective pressures imposed by rocky and ground habitats have resulted in the 399 differentiation of head and limb morphology (Goodman et al. 2008; Kaliontzopoulou et al. 2010; 400 Garcia-Porta et al. 2017; Foster et al. 2018). Indeed, such phenotypic differences resulting from the 401 effects of habitat-based ecological selection have been extensively documented in reptiles as well as in 402 other vertebrates (Losos 2009; Reynolds et al. 2016; Hipsley and Müller 2017; Samuels and Hopkins 403 2017; Watanabe et al. 2019; Busschau and Boissinot 2022; Navalón et al. 2022; Friedman et al. 2022), and our work in *Pristurus* thus contributes to this growing body of literature. Nonetheless, because the ecological shift to ground-dwelling habitats occurred only once on the phylogeny, it is also possible that some unmeasured feature that evolved on the same branch could have affected the observed patterns. Thus, some caution in interpreting the causal direction of this trend is warranted.

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However, considering the habitat-driven morphology perspective, the findings revealed here may 410 reflect ecological and behavioral changes linked to the adoption of a new lifestyle. For lizards, the 411 transition to utilizing ground habitats implies adopting an existence in more open environments 412 than in rocky substrates. As such, numerous aspects of daily existence (including activity patterns, 413 climatic factors, prey availability, abundance of predators, etc.) are expected to exert a differential 414 influence on an organism's phenotype when compared with life in their ancestral environment 415 (Fuentes and Cancino 1979). Indeed, the largest ground-dwelling Pristurus species (P. carteri, P. 416 ornitocephalus, and P. collaris) differ from the rest of the genus in having developed partially 417 nocturnal habits, which would presumably have major ecological consequences for their survival 418 and reproduction. In this sense, these species might have been subjected to evolutionary processes 419 selecting for larger relative head proportions, which would allow them to accommodate larger 420 or modified eyes, a clear advantage in animals with nocturnal and semi-nocturnal habits (Hall 421 and Ross 2006; Ross et al. 2007; Hall et al. 2012). Likewise, the large relative proportions found 422 in the limbs of large ground-dwelling species (Figure 3B) might be related to selective processes 423 favoring longer limbs in large species present in this new ecological context. Longer limbs in open habitats – particularly for large species – might be advantageous for rapidly running and hiding in 425 the sparse vegetation (Arnold 2009) and, in hyper-arid areas such as the Arabian Peninsula, this 426 morphology might contribute to thermoregulation separating the body from the ground (Huey 1974; 427 Arnold 1980; Avery et al. 1982). A-The lack of repeated events of colonization of ground habitats 428 in *Pristurus* makes it challenging to corroborate these adaptive explanations about phenotypic 429 changes, but a more detailed examination of behavioral and morphological traits (e.g., eye shape, 430 limb insertion) might shed light on the factors driving this pattern and serve to establish a stronger 431 adaptive link between habitat use and morphological and allometric trends in *Pristurus*. 432

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Another important finding of our study was the strong concordance between static intraspecific allometry across individuals and evolutionary allometry among *Pristurus* species. Our analyses 435 revealed small pairwise angles between static intraspecific and evolutionary allometry vectors, 436 indicating that allometric trends at these two levels were oriented in similar directions and 437 were largely concordant. As such, size-associated changes in body form among individuals 438 were predictive of evolutionary shifts across taxa at higher macroevolutionary scales. 439 in turn, suggests that body form evolution in Pristurus may follow an allometric line of 440 least resistance (Marroig and Cheverud 2005). In other empirical systems, a similarly tight 441 correspondence between static intraspecific and evolutionary allometry has also been observed (Marroig and Cheverud 2005; Firmat et al. 2014; Voje et al. 2014; Brombacher et al. 2017; 443 Marcy et al. 2020), though the trend is not universal across all taxa or traits (see Klingenberg 444 and Zimmermann 1992; Voje et al. 2022). Nonetheless, when such trends are present, they 445 imply that allometric trajectories impose a prevailing influence on the magnitude, direction, 446 and rate of phenotypic change across the phylogeny. Our work in *Pristurus* contributes to 447 the growing literature on this topic, and suggests that perhaps such patterns may be more widespread. 448

Given the observation that static-intraspecific and evolutionary allometry in *Pristurus* are largely 450 concordant, an obvious question is: why might this be the case? One possible explanation is 451 that when genetic covariation remains relatively constant, selection on body size will generate 452 an evolutionary allometric trajectory along the trend described by static-intraspecific allometry (Lande 1979, 1985). Here, allometry effectively acts as a constraint on evolutionary change, as 454 size-associated shape changes at one hierarchical level are linked to changes at another level (Voje 455 et al. 2014, 2022; Pélabon et al. 2014). Further, when this is the case, one may also expect 456 high levels of phenotypic integration in traits associated with body size changes. Indeed, our 457 analyses reveal precisely this pattern in *Pristurus*, with the highest levels of integration in the 458 group (ground-dwelling) whose static-intraspecific allometry is most similar to that of evolutionary 459 allometry. Thus, our results reveal that patterns of trait covariation are more constrained in ground-dwelling species, such that their differences in body form are most likely found along 461 the primary allometric axis. When viewed in this light, integration and allometry may thus be

449

interpreted as potential drivers that facilitate morphological change, as they provide a phenotypic pathway through adaptive lines of least resistance that enable rapid evolutionary changes in particular phenotypic directions but not in others (Felice et al. 2018; Navalón et al. 2020). The fact that ground-dwelling species in *Pristurus* have been found to have the widest phenotypic disparity, greatest range of body sizes, and highest rates of morphological evolution (Tejero-Cicuéndez et al. 2021a) are all consistent with this hypothesis, and suggest that in this group, integration describes the path of morphological evolution along allometric lines of least resistance.

470

Finally, interpreting the observed patterns of phenotypic integration and allometry relative to habitat-471 specific differences helps to shed light on the possible pathways by which phenotypic diversity in 472 Pristurus has evolved. For instance, prior work on this system (Tejero-Cicuéndez et al. 2021a) 473 revealed that the colonization of new ecological habitats might have elicited strong ecological selection 474 and phenotypic responses. This was particularly true of the invasion of ground habitats, where 475 ground-dwelling species displayed the largest variation in body size in the genus. This observation 476 implies-might be related to some level of ecological selection on body size. In lizards, the ecological 477 context in which species exist is known to play a pervasive role in body size evolution (James and 478 M'closkey 2004; Meiri 2008; Tamar et al. 2019), as it does in other animal groups (Bergmann 479 1847; Calder 1983; Peters 1983; LaBarbera 1989; Olson et al. 2009). While to date this has not 480 been thoroughly explored in *Pristurus*, the evolutionary patterns revealed by our analyses suggest 481 that the body size diversity in this clade conforms, at least in part, with patterns expected under 482 ecological selection on body size (although this perspective would be further supported if there had 483 been repeated instances of colonization of ground habitats in the genus). Intriguingly, such patterns 484 are not only observed in ground- and rock-dwelling taxa, but also in arboreal species -(which, unlike 485 ground-dwelling *Pristurus*, have evolved this lifestyle independently), whose restricted phenotypic 486 diversity in both size and shape (Figures 3 & 5) is consistent with strong ecological selection 487 in the arboreal habitat (Baken and Adams 2019; Baken et al. 2021b). If that is the case, this 488 contrasts with the evolutionary dynamics observed in other lizards such as the Anolis radiations, 489 where there are multiple morphotypes for different strata of the arboreal habitat (Losos 2009). 490 Furthermore, our study identified the presence of strong integration and allometric trajectories, such 491

that evolutionary changes in body size elicit corresponding changes in body form. However, these trends differed significantly across habitats, implying that, at evolutionary scales, they might serve to channel phenotypic responses to selection, but do so in differing directions for the different habitat groups. This, in turn, suggests that *Pristurus* species occupying different habitats display differing combinations of body size with body form. The From this adaptive perspective, the evolutionary consequence of ecological selection is that species have evolved similar shapes (Figure 6), but do so in differing habitats, and at different body sizes (Figure 5). Therefore, the phenotypic diversity observed in *Pristurus* is best explained as the result of a complex interplay between ecological selection, body size differentiation, and differing allometric trajectories across ecological habitats.

References

- Adams, D. C., and M. L. Collyer. 2009. A general framework for the analysis of phenotypic trajectories in evolutionary studies. Evolution 63:1143–1154.
- Adams, D. C., and A. Nistri. 2010. Ontogenetic convergence and evolution of foot morphology in
- European cave salamanders (Family: Plethodontidae). BMC Evolutionary Biology 10:1–10.
- BioMed Central.
- Albertson, R. C., and T. D. Kocher. 2001. Assessing morphological differences in an adaptive
- trait: A landmark-based morphometric approach. Journal of Experimental Zoology 289:385–
- 509 403.
- Arnold, E. N. 1993. Historical changes in the ecology and behaviour of semaphore geckos
- 511 (Pristurus, Gekkonidae) and their relatives. Journal of Zoology 229:353–384.
- Arnold, E. N. 2009. Relationships, evolution and biogeography of semaphore geckos, *Pristurus*
- (Squamata, Sphaerodactylidae) based on morphology. Zootaxa 2060:1–21.
- Arnold, E. N. 1980. The reptiles and amphibians of Dhofar, Soutern Arabia. Journal of Oman
- Studies Special Report:273–332.
- Arnold, S. J. 1983. Morphology, performance, fitness. American Zoologist 23:347–361.
- Avery, R. A., Bedford J. D., and C. P. Newcombe. 1982. The role of thermoregulation in lizard
- biology: Predatory efficiency in a temperate basker. Behavioral Ecology and Sociobiology
- 11:261-267.
- Baken, E. K., and D. C. Adams. 2019. Macroevolution of arboreality in salamanders. Ecology
- and Evolution 9:7005–7016.
- Baken, E. K., M. L. Collyer, A. Kaliontzopoulou, and D. C. Adams. 2021a. Geomorph
- 4.0 and gmShiny: Enhanced analytics and a new graphical interface for a comprehensive
- morphometric experience. Methods in Ecology and Evolution 12:2355–2363.
- Baken, E. K., L. E. Mellenthin, and D. C. Adams. 2021b. Is salamander arboreality limited by
- broad-scale climatic conditions? PLoS ONE 16:e0255393.
- Bardua, C., A.-C. Fabre, J. Clavel, M. Bon, K. Das, E. L. Stanley, D. C. Blackburn, and A.
- Goswami. 2021. Size, microhabitat, and loss of larval feeding drive cranial diversification in
- frogs. Nature Communications 12. Springer Science; Business Media LLC.

- Bergmann, C. 1847. Über die verhaltnisse der warmeokonomie der thiere zu ihrer grosse.
- Göttinger Studien 1:595–708.
- Berner, D., D. C. Adams, A.-C. Grandchamp, and A. P. Hendry. 2008. Natural selection
- drives patterns of lake-stream divergence in stickleback foraging morphology. Journal of
- Evolutionary Biology 21:1653–1665.
- Bookstein, F. L. 2022. Dimensions of morphological integration. Evolutionary Biology 49:342–
- 536 372.
- Bright, J. A., J. Marugán-Lobón, S. N. Cobb, and E. J. Rayfield. 2016. The shapes of bird
- beaks are highly controlled by nondietary factors. Proceedings of the National Academy of
- Sciences 113:5352–5357.
- Bright, J. A., J. Marugán-Lobón, E. J. Rayfield, and S. N. Cobb. 2019. The multifactorial
- nature of beak and skull shape evolution in parrots and cockatoos (psittaciformes). BMC
- Evolutionary Biology 19. Springer Science; Business Media LLC.
- Brombacher, A., P. A. Wilson, I. Bailey, and T. H. G. Ezard. 2017. The breakdown of static and
- evolutionary allometries during climatic upheaval. The American Naturalist 190:350–362.
- Busschau, T., and S. Boissinot. 2022. Habitat determines convergent evolution of cephalic horns
- in vipers. Biological Journal of the Linnean Society 135:652–664.
- 547 Calder, W. A. 1983. Ecological scaling: Mammals and birds. Annual Review of Ecology and
- 548 Systematics 14:213–230.
- Cardini, A., D. Polly, R. Dawson, and N. Milne. 2015. Why the long face? Kangaroos and
- wallabies follow the same "rule" of cranial evolutionary allometry (CREA) as placentals.
- Evolutionary Biology 42:169–176.
- ⁵⁵² Cardini, A., and P. D. Polly. 2013. Larger mammals have longer faces because of size-related
- constraints on skull form. Nature Communications 4.
- Chatterji, R. M., C. A. Hipsley, E. Sherratt, M. N. Hutchinson, and M. E. H. Jones. 2022.
- Ontogenetic allometry underlies trophic diversity in sea turtles (chelonioidea). Evolutionary
- Ecology 36:511–540.
- 557 Cock, A. G. 1966. Genetical aspects of metrical growth and form in animals. The Quarterly
- Review of Biology 41:131–190.
- Collar, D. C., J. A. Schulte, B. C. O'Meara, and J. B. Losos. 2010. Habitat use affects

- morphological diversification in dragon lizards. Journal of Evolutionary Biology 23:1033–
 1049.
- Collyer, M. L., and D. C. Adams. 2007. Analysis of two-state multivariate phenotypic change in ecological studies. Ecology 88:683–692.
- Collyer, M. L., and D. C. Adams. 2013. Phenotypic trajectory analysis: Comparison of shape change patterns in evolution and ecology. Hystrix 24:75–83.
- Collyer, M. L., and D. C. Adams. 2022. R: RRPP: Linear model evaluation with randomized
 residuals in a permutation procedure. Vsn. 1.3.2. R Foundation for Statistical Computing,
 Vienna, Austria.
- Collyer, M. L., and D. C. Adams. 2018. RRPP: An R package for fitting linear models to
 high-dimensional data using residual randomization. Methods in Ecology and Evolution
 9:1772–1779.
- Collyer, M. L., D. J. Sekora, and D. C. Adams. 2015. A method for analysis of phenotypic change for phenotypes described by high-dimensional data. Heredity 115:357–365.
- Conaway, M. A., and D. C. Adams. 2022. An effect size for comparing the strength of morphological integration across studies. Evolution 76:2244–2259.
- Drake, A. G., and C. P. Klingenberg. 2008. The pace of morphological change: Historical transformation of skull shape in St Bernard dogs. Proceedings of the Royal Society B:

 Biological Sciences 275:71–76.
- Durant, S. M., N. Pettorelli, S. Bashir, R. Woodroffe, T. Wacher, P. De Ornellas, C. Ransom,
- T. Abáigar, M. Abdelgadir, H. El Algamy, M. Beddiaf, F. Belbachir, A. Belbachir-Bazi, A.
- A. Berbash, R. Beudels-Jamar, L. Boitani, C. Breitenmoser, M. Cano, P. Chardonnet, B.
- Collen, W. A. Cornforth, F. Cuzin, P. Gerngross, B. Haddane, M. Hadjeloum, A. Jacobson,
- A. Jebali, F. Lamarque, D. Mallon, K. Minkowski, S. Monfort, B. Ndoassal, J. Newby, B.
- E. Ngakoutou, B. Niagate, G. Purchase, S. Samaïla, A. K. Samna, C. Sillero-Zubiri, A. E.
- Soultan, M. R. Stanley Price, and J. E. M. Baillie. 2012. Forgotten biodiversity in desert ecosystems. Science 336:1379–1380.
- Esquerré, D., E. Sherratt, and J. S. Keogh. 2017. Evolution of extreme ontogenetic allometric
- diversity and heterochrony in pythons, a clade of giant and dwarf snakes. Evolution 71:2829–
- 589 2844.

- Felice, R. N., D. Pol, and A. Goswami. 2021. Complex macroevolutionary dynamics underly the evolution of the crocodyliform skull. Proceedings of the Royal Society B: Biological Sciences 288:20210919.
- Felice, R. N., M. Randau, and A. Goswami. 2018. A fly in a tube: Macroevolutionary expectations for integrated phenotypes. Evolution 72:2580–2594.
- Firmat, C., I. Lozano-Fernández, J. Agustí, G. H. Bolstad, G. Cuenca-Bescós, T. F. Hansen, and
 C. Pélabon. 2014. Walk the line: 600000 years of molar evolution constrained by allometry
 in the fossil rodent *Mimomys savini*. Philosophical Transactions of the Royal Society B:
 Biological Sciences 369:20140057.
- Foster, K. L., T. Garland, L. Schmitz, and T. E. Higham. 2018. Skink ecomorphology: Forelimb and hind limb lengths, but not static stability, correlate with habitat use and demonstrate multiple solutions. Biological Journal of the Linnean Society 125:673–692.
- Freedman, D., and D. Lane. 1983. A nonstochastic interpretation of reported significance levels.

 Journal of Business & Economic Statistics 1:292–298.
- Friedman, S. T., M. L. Collyer, S. A. Price, and P. C. Wainwright. 2022. Divergent Processes
 Drive Parallel Evolution in Marine and Freshwater Fishes. Systematic biology 71:1319–1330.
- Fuentes, E. R., and J. Cancino. 1979. Rock-ground patchiness in a simple liolaemus lizard community (reptilia, lacertilia, iguanidae). Journal of Herpetology 13:343.
- Garcia-Porta, J., M. Simó-Riudalbas, M. Robinson, and S. Carranza. 2017. Diversification
 in arid mountains: Biogeography and cryptic diversity of *Pristurus rupestris rupestris* in
 Arabia. Journal of Biogeography 44:1694–1704.
- Goodman, B. A., D. B. Miles, and L. Schwarzkopf. 2008. Life on the rocks: Habitat use drives morphological and performance evolution in lizards. Ecology 89:3462–3471.
- Goswami, A., M. Randau, P. D. Polly, V. Weisbecker, C. Verity Bennett, L. Hautier, and M. R. Sánchez-Villagra. 2016. Do developmental constraints and high integration limit the evolution of the marsupial oral apparatus? Integrative and Comparative Biology 56:404–415.
- Goswami, A., J. B. Smaers, C. Soligo, and P. D. Polly. 2014. The macroevolutionary consequences of phenotypic integration: From development to deep time. Philosophical Transactions of the Royal Society B: Biological Sciences 369:20130254.
- Gould, S. J. 1966. Allometry and size in ontogeny and phylogeny. Biological Reviews 41:587–638.

- Grant, P. R., and B. R. Grant. 2006. Evolution of character displacement in darwin's finches.

 Science 313:224–226.
- Hall, M. I., J. M. Kamilar, and E. C. Kirk. 2012. Eye shape and the nocturnal bottleneck of mammals. Proceedings of the Royal Society B: Biological Sciences 279:4962–4968.
- Hall, M. I., and C. F. Ross. 2006. Eye shape and activity pattern in birds. Journal of Zoology 271:437–444.
- Hipsley, C. A., and J. Müller. 2017. Developmental dynamics of ecomorphological convergence in a transcontinental lizard radiation. Evolution 71:936–948.
- Huey, R. B. 1974. Behavioral thermoregulation in lizards: Importance of associated costs.

 Science 184:1001–1003.
- Huxley, J. S. 1932. The problems of relative growth. Johns Hopkins University Pres.
- James, S. E., and R. T. M'closkey. 2004. Patterns of body size and habitat use in a lizard assemblage. Ecoscience 11:160–167.
- Jastrebski, C. J., and B. W. Robinson. 2004. Natural selection and the evolution of replicated trophic polymorphisms in pumpkinseed sunfish (*Lepomis gibbosus*). Evolutionary Ecology Research 6:285–305.
- Jolicoeur, P. 1963. The multivariate generalization of the allometry equation. Biometrics 19:497–499.
- Kaliontzopoulou, A., M. A. Carretero, and D. C. Adams. 2015. Ecomorphological variation in male and female wall lizards and the macroevolution of sexual dimorphism in relation to habitat use. Journal of Evolutionary Biology 28:80–94.
- Kaliontzopoulou, A., M. A. Carretero, and G. A. Llorente. 2010. Intraspecific ecomorphological variation: Linear and geometric morphometrics reveal habitat-related patterns within *Podarcis bocagei* wall lizards. Journal of Evolutionary Biology 23:1234–1244.
- Kappelman, J., D. T. Rasmussen, W. J. Sanders, M. Feseha, T. Bown, P. Copeland, J. Crabaugh,
- J. Fleagle, M. Glantz, A. Gordon, B. Jacobs, M. Maga, K. Muldoon, A. Pan, L. Pyne, B.
- Richmond, T. Ryan, E. R. Seiffert, S. Sen, L. Todd, M. C. Wiemann, and A. Winkler. 2003.
- Oligocene mammals from Ethiopia and faunal exchange between Afro-Arabia and Eurasia.
- Nature 426:549–552.
- Klingenberg, C. P. 1996. Advances in morphometrics. Pp. 23–49 in L. F. Marcus, M. Corti, A.

- Loy, and G. J. P. N. D. E. Slice, eds. Plenum Press.
- Klingenberg, C. P. 2016. Size, shape, and form: Concepts of allometry in geometric morphometrics. Development Genes and Evolution 226:113–137.
- Klingenberg, C. P., and J. Marugán-Lobón. 2013. Evolutionary covariation in geometric morphometric data: Analyzing integration, modularity, and allometry in a phylogenetic context. Systematic Biology 62:591–610.
- Klingenberg, C. P., and M. Zimmermann. 1992. Static, ontogenetic, and evolutionary allometry:

 A multivariate comparison in nine species of water striders. American Naturalist 140:601–620.
- Kolmann, M. A., F. P. L. Marques, J. C. Weaver, M. N. Dean, J. P. Fontenelle, and N. R. Lovejoy. 2022. Ecological and phenotypic diversification after a continental invasion in neotropical freshwater stingrays. Integrative and Comparative Biology 62:424–440.
- LaBarbera, M. 1989. Analyzing body size as a factor in ecology and evolution. Annual Review of Ecology and Systematics 20:97–117.
- Lande, R. 1979. Quantitative genetic analysis of multivariate evolution, applied to brain-body size allometry. Evolution 33:402–416.
- Lande, R. 1985. Size and scaling in primate biology. Pp. 21–32 in W. L. Jungers, ed. Plenum
 Press.
- Losos, J. B. 2009. Lizards in an evolutionary tree: Ecology and adaptive radiation of anoles.

 University of California Press.
- Losos, J. B. 1992. The evolution of convergent structure in Caribbean *Anolis* communities.

 Systematic Biology 41:403–420.
- Losos, J. B., and R. E. Ricklefs. 2009. Adaptation and diversification on islands. Nature 457:830–836.
- Mahler, D. L., T. Ingram, L. J. Revell, and J. B. Losos. 2013. Exceptional convergence on the macroevolutionary landscape in island lizard radiations. Science 341:292–295.
- Marcy, A. E., T. Guillerme, E. Sherratt, K. C. Rowe, M. J. Phillips, and V. Weisbecker. 2020.

 Australian rodents reveal conserved cranial evolutionary allometry across 10 million years of
 murid evolution. The American Naturalist 196:755–768.
- Marroig, G., and J. M. Cheverud. 2005. Size as a line of least evolutionary resistance: Diet and adaptive morphological radiation in New World monkeys. Evolution 59:1128–1142.

- Martinez, C. M., S. T. Friedman, K. A. Corn, O. Larouche, S. A. Price, and P. C. Wainwright.
- 2021. The deep sea is a hot spot of fish body shape evolution. Ecology Letters 24:1788–1799.
- Meiri, S. 2008. Evolution and ecology of lizard body sizes. Global Ecology and Biogeography
 17:724–734.
- Mosimann, J. E. 1970. Size allometry: Size and shape variables with characterizations of the lognormal and generalized gamma distributions. Journal of the American Statistical Association 65:930–945.
- Navalón, G., A. Bjarnason, E. Griffiths, and R. B. J. Benson. 2022. Environmental signal in the evolutionary diversification of bird skeletons. Nature 611:306–311.
- Navalón, G., J. Marugán-Lobón, J. A. Bright, C. R. Cooney, and E. J. Rayfield. 2020. The consequences of craniofacial integration for the adaptive radiations of Darwin's finches and Hawaiian honeycreepers. Nature Ecology & Evolution 4:270–278. Nature Publishing Group.
- Olson, V. A., R. G. Davies, C. D. L. Orme, G. H. Thomas, S. Meiri, T. M. Blackburn, K. J. Gaston, I. P. F. Owens, and P. M. Bennett. 2009. Global biogeography and ecology of body size in birds. Ecology Letters 12:249–259.
- Patterson, M., A. K. Wolfe, P. A. Fleming, P. W. Bateman, M. L. Martin, E. Sherratt, and
 N. M. Warburton. 2022. Ontogenetic shift in diet of a large elapid snake is facilitated by
 allometric change in skull morphology. Evolutionary Ecology 36:489–509.
- Pavlicev, M., J. M. Cheverud, and G. P. Wagner. 2009. Measuring morphological integration using eigenvalue variance. Evolutionary Biology 36:157–170.
- Pélabon, C., G. H. Bolstad, C. K. Egset, J. M. Cheverud, M. Pavlicev, and G. Rosenqvist. 2014.

 On the relationship between ontogenetic and static allometry. The American Naturalist

 181:195–212.
- Peters, R. H. 1983. The ecological implications of body size. Cambridge University Press.
- Piras, P., P. Colangelo, D. C. Adams, A. Buscalioni, J. Cubo, T. Kotsakis, C. Meloro, and P. Raia. 2010. The gavialis-tomistoma debate: The contribution of skull ontogenetic allometry and growth trajectories to the study of crocodylian relationships. Evolution and Development 12:568–579.
- Price, S. A., S. T. Friedman, and P. C. Wainwright. 2015. How predation shaped fish: The impact of fin spines on body form evolution across teleosts. Proceedings of the Royal Society

- B: Biological Sciences 282:20151428.
- Pyron, R. A., and F. T. Burbrink. 2009. Body size as a primary determinant of ecomorphologi-
- cal diversification and the evolution of mimicry in the lampropeltinine snakes (serpentes:
- colubridae). Journal of Evolutionary Biology 22:2057–2067. Wiley.
- R Core Team. 2022. R: A language and environment for statistical computing. Version 4.2.1. R
- Foundation for Statistical Computing, Vienna, Austria.
- Reaney, A. M., Y. Bouchenak-Khelladi, J. A. Tobias, and A. Abzhanov. 2020. Ecological and
- morphological determinants of evolutionary diversification in Darwin's finches and their
- relatives. Ecology and Evolution 10:14020–14032.
- Revell, L. J. 2012. Phytools: An R package for phylogenetic comparative biology (and other
- things). Methods in Ecology and Evolution 3:217–223.
- Reynolds, R. G., D. C. Collar, S. A. Pasachnik, M. L. Niemiller, A. R. Puente-Rolón, and L. J.
- Revell. 2016. Ecological specialization and morphological diversification in Greater Antillean
- boas. Evolution 70:1882–1895.
- Ross, C. F., M. I. Hall, and C. P. Heesy. 2007. Were basal primates nocturnal? Evidence from
- eye and orbit shape. Pp. 233–256 in PRIMATE ORIGINS: Adaptations and evolution.
- Springer.
- Samuels, J. X., and S. S. B. Hopkins. 2017. The impacts of Cenozoic climate and habitat changes
- on small mammal diversity of North America. Global and Planetary Change 149:36–52.
- Schluter, D. 1996. Adaptive radiation along genetic lines of least resistance. Evolution 50:1766–
- 730 1774.
- Schluter, D., and P. R. Grant. 1984. Determinants of morphological patterns in communities of
- Darwin's finches. The American Naturalist 123:175–196.
- Schluter, D., and J. D. McPhail. 1992. Ecological character displacement and speciation in
- sticklebacks. The American Naturalist 140:85–108.
- Sherratt, E., D. J. Gower, C. P. Klingenberg, and M. Wilkinson. 2014. Evolution of cranial
- shape in caecilians (amphibia: gymnophiona). Evolutionary Biology 41:528–545.
- 537 Sidlauskas, B. 2008. Continuous and arrested morphological diversification in sister clades of
- characiform fishes: A phylomorphospace approach. Evolution 62:3135–3156.
- Stuart, Y. E., T. Veen, J. N. Weber, D. Hanson, M. Ravinet, B. K. Lohman, C. J. Thompson, T.

- Tasneem, A. Doggett, R. Izen, N. Ahmed, R. D. H. Barrett, A. P. Hendry, C. L. Peichel, and
 D. I. Bolnick. 2017. Contrasting effects of environment and genetics generate a continuum
 of parallel evolution. Nature Ecology and Evolution 1:158.
- Tamar, K., P. Mitsi, M. Simó-Riudalbas, H. Tejero-Cicuéndez, T. Al-Sariri, and S. Carranza.

 2019. Systematics, biogeography, and evolution of *Pristurus minimus* (Squamata, Sphaerodactylidae) with the discovery of the smallest Arabian vertebrate. Systematics and Biodiversity 17:349–366.
- Tejero-Cicuéndez, H., A. H. Patton, D. S. Caetano, J. Šmíd, L. J. Harmon, and S. Carranza.

 2022. Reconstructing squamate biogeography in Afro-Arabia reveals the influence of a

 complex and dynamic geologic past. Systematic Biology 71:261–272.
- Tejero-Cicuéndez, H., M. Simó-Riudalbas, I. Menéndez, and S. Carranza. 2021a. Ecological specialization, rather than the island effect, explains morphological diversification in an ancient radiation of geckos. Proceedings of the Royal Society B: Biological Sciences 288:20211821.
- Tejero-Cicuéndez, H., M. Simó-Riudalbas, I. Menéndez, and S. Carranza. 2021b. Ecological specialization, rather than the island effect, explains morphological diversification in an ancient radiation of geckos. Dryad digital repository. (Doi:10.5061/dryad.xwdbrv1f6).
- Urban, S., J. Gerwin, C. D. Hulsey, A. Meyer, and C. F. Kratochwil. 2022. The repeated evolution of stripe patterns is correlated with body morphology in the adaptive radiations of East African cichlid fishes. Ecology and Evolution 12:e8568.
- Voje, K. L., M. A. Bell, and Y. E. Stuart. 2022. Evolution of static allometry and constraint on evolutionary allometry in a fossil stickleback. Journal of Evolutionary Biology 35:423–438.
- Voje, K. L., T. F. Hansen, C. K. Egset, G. H. Bolstad, and C. Pélabon. 2014. Allometric constraints and the evolution of allometry. Evolution 68:866–885.
- Wagner, G. P., and J. Zhang. 2011. The pleiotropic structure of the genotype-phenotype map:
 The evolvability of complex organisms. Nature Reviews Genetics 12:204–213.
- Wagner, G., and L. Altenberg. 1996. Perspective: Complex adaptations and the evolution of evolvability. Evolution 50:967–976.
- Watanabe, A., A. C. Fabre, R. N. Felice, J. A. Maisano, J. Müller, A. Herrel, and A. Goswami.

 2019. Ecomorphological diversification in squamates from conserved pattern of cranial
 integration. Proceedings of the National Academy of Sciences of the United States of

- America 116:14688–14697.
- Wickham, H., M. Averick, J. Bryan, W. Chang, L. D. McGowan, R. François, G. Grolemund, A.
- Hayes, L. Henry, J. Hester, M. Kuhn, T. L. Pedersen, E. Miller, S. M. Bache, K. Müller, J.
- Ooms, D. Robinson, D. P. Seidel, V. Spinu, K. Takahashi, D. Vaughan, C. Wilke, K. Woo,
- and H. Yutani. 2019. Welcome to the tidyverse. Journal of Open Source Software 4:1686.
- Zelditch, M. L., and D. L. Swiderski. 2022. The predictable complexity of evolutionary allometry.
- Evolutionary Biology 50:56–77.

Acknowledgments: We are very grateful to J. Roca, M. Metallinou, K. Tamar, J. Šmíd, R. Vasconcelos, R. Sindaco, F. Amat, Ph. de Pous, L. Machado, J. Garcia-Porta, J. Els, T. Mazuch,
T. Papenfuss, and all the people from the Environment Authority, Oman, for their help in
different aspects of the work.

777

778

This work was funded in part by PGC2018-098290-B-I00 (MCI-Funding Statement: 783 U/AEI/FEDER, UE) and PID2021-128901NB-I00 (MCIN/AEI/10.13039/501100011033 and by 784 ERDF, A way of making Europe), Spain to SC. HT-C is supported by a "Juan de la Cierva -785 Formación" postdoctoral fellowship (FJC2021-046832-I). IM was funded by the Alexander von 786 Humboldt Foundation through a Humboldt Research Fellowship. AT is supported by the "la 787 Caixa" doctoral fellowship programme (LCF/BQ/DR20/11790007). GR was funded by an FPI 788 grant from the Ministerio de Ciencia, Innovación y Universidades, Spain (PRE2019-088729). 789 BB-C was funded by FPU grant from Ministerio de Ciencia, Innovación y Universidades, Spain 790 (FPU18/04742). DCA was funded in part by National Science Foundation Grant DBI-1902511. 791

Data availability statement: All the data used in this study are available on DRYAD from a previous study: https://doi.org/10.5061/dryad.xwdbrv1f6 (Tejero-Cicuéndez et al. 2021b). The scripts for implementing all analyses and generating the figures in this manuscript can be found in the Supplementary Material and in a GitHub repository (and on DRYAD upon acceptance).

Competing interests: The authors declare no competing interests.