- Evolution along allometric lines of least resistance: Morphological
- differentiation in *Pristurus* geckos
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3 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 15 trends and trait integration levels is less well understood. Here we evaluate the role of allometry in 16 shaping body size and shape diversity in *Pristurus* geckos utilizing differing habitats. We found that patterns of body shape allometry and integration were distinct in species with different habitat 18 preferences, with ground-dwelling *Pristurus* displaying the most divergent allometric trend and 19 the strongest integration. There was also strong concordance between static allometry across individuals and evolutionary allometry among species, revealing that body shape differences among 21 individuals were predictive of evolutionary changes across the phylogeny at macroevolutionary scales. 22 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 shape 24 changes across the phylogeny. When viewed in phylomorphospace, the largest rock-dwelling species 25 were most similar in body shape to the smallest ground-dwelling species, and vice versa. Thus, in 26 Pristurus, phenotypic evolution along the differing habitat-based allometric trajectories resulted in similar body shapes at differing body sizes in distinct ecological habitats.

29 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 in their habitat. Evolutionary theory 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 [1–5].

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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 [6,7], this convergent evolution is treated as strong evidence of adaptation. Indeed the ecomorphological paradigm [8] 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 [6,9,10], differential beak morphology in species of Darwin's finches [11–13], the recurring phenotypes of African lake cichlids across ecological regimes [14,15], and the distinct body forms of freshwater fishes in benthic and limnetic habitats [16–18], among others.

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However, while the patterns of morphological differences in distinct ecological contexts have been well documented, less-well understood is how this differentiation has been influenced by trait covariation associated with body size differences (i.e., allometry). Evaluating allometric trends across hierarchical levels (e.g., comparing allometry at the individual level, or static allometry, and among species, or evolutionary allometry) may aid in our understanding of how adaptive

morphological change occurs at macroevolutionary scales [19]. It has long been recognized that
the interrelationships among traits can exert a strong influence on how phenotypic evolution
proceeds, as trait correlations influence the degree to which phenotypic variation is exposed to
selection [20]. Thus, the integration among traits can constrain phenotypic change in certain
directions, or enhance variation along other phenotypic axes [20–27]. Further, because nearly all
linear traits covary strongly with overall body size [28,29], allometric trends could be considered
the quintessential expression of phenotypic integration. Thus, identifying whether allometric
patterns differ across habitats, and how such patterns of trait covariation affect ecomorphological
trends among species utilizing those habitats, remains an important question worthy of investigation.

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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 diversity. Prior work on this system [30] revealed that the colonization of ground habitats has been a trigger of morphological change, specifically reflected in an increase in body size and shape disparity. Interestingly, some ground-dwelling species are among the largest of the genus and also show 71 increased relative head sizes and limb proportions, while some other species with this ecological specialization have evolved to be among the smallest of the group. Additionally, among the species 73 exploiting rocky habitats (the most common ecological feature in *Pristurus*), there are also species 74 with both considerably large and small body sizes [30]. What remains unexplored, however, is how 75 the evolution of body shape is related to differences in body size and whether habitat specialization has an impact in this shape-size relationship. 77

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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 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 within habitats, demonstrating that the interplay between ecological specialization and allometric trajectories in species with disparate body size may play a determinant role in shaping the phenotypic evolution and hence in adaptive dynamics in this clade.

90 2. Materials and Methods

91 (a) Data

We used a combination of phenotypic, phylogenetic, and ecological data to characterize and evaluate intra- and interspecific allometric trends. The data utilized here were obtained from our prior work on this system [30,31], and are briefly described here. First we used a time-dated, molecular phylogeny of squamates that included all members of the genus *Pristurus*, including several currently undescribed taxa. The tree was estimated in a Bayesian framework, using five mitochondrial markers, six nuclear markers, and 21 calibration points [31]. Next we categorized each species as belonging to one of three ecological groups (ground, rock, or tree), based on descriptions of habitat use found in the literature [30]. Finally, we obtained a phenotypic data set containing body size (snout-vent length: SVL) and eight linear measurements (Figure 1) that described overall body form: trunk 100 length (TL), head length (HL), head width (HW), head height (HH), humerus length (Lhu), ulna 101 length (Lun), femur length (Lfe), and tibia length (Ltb) [30]. We restricted our study to those 102 species represented by nine or more individuals; resulting in a dataset of 687 individuals from 25 103 species (invidivuals per species: $\mu = 27$; min = 9, max = 56). Species in the phenotypic dataset 104 were then matched to the phylogeny, which was subsequently pruned to the final topology. All 105 measurements were log-transformed prior to statistical analyses. Additional details regarding data 106 collection and formal descriptions of each linear measurement may be found in the original sources [30,31]. The data are available on DRYAD: https://doi.org/10.5061/dryad.xwdbrv1f6 [32]. 108

(b) Statistical and Comparative Analyses

We conducted a series of analyses to interrogate allometric trends, patterns of integration, and macroevolutionary changes in allometry, relative to differentiation in body form. First we characterized evolutionary allometry in the genus by performing a phylogenetic multivariate regression of body form on body size (i.e., SVL), using the species means as data. We then
performed an analogous procedure at the individual level, regressing body form on body size using
our entire dataset. From both the species-level (phylogenetic) and the individual-level analyses, we
obtained the set of regression coefficients, and calculated the difference in their angular direction to
describe the extent to which patterns of allometry at the individual level were concordant with
evolutionary allometric trends across species.

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Next we used the dataset containing all individuals to determine whether trends in static allometry 120 differed across habitat groups. This was accomplished by performing a multivariate analysis of 121 covariance, with body size (SVL), habitat, and $SVL \times habitat$ as model effects. Significance was 122 evaluated using 999 iterations of a permutation procedure, where residuals from a reduced model 123 were randomly permuted in each permutation (RRPP), model statistics were recalculated, and used 124 to generate empirical null sampling distributions to evaluate the observed test statistics [following 125 33,34,35]. We then compared the multivariate allometric vectors for each habitat group to one 126 another, and to a vector representing multivariate isometry, by calculating pairwise differences 127 in their angular direction in morphospace, and evaluating these relative to empirical sampling 128 distributions obtained through RRPP [34,36,37]. Here, residuals were obtained from a common 129 isometry reduced model, whose common slope component described a pattern of multivariate 130 isometry, and whose intercepts allowed for differences in least-squares means among groups. 131 Patterns of multivariate allometry relative to body size were visualized via regression scores [38] and 132 predicted lines [39], based on the coefficients and fitted values from the linear model described above. 133

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Additionally, because allometry describes the extent to which traits covary with body size and with each other (i.e., integration), we conducted an analysis of integration. Here we characterized the extent of morphological integration in body form for individuals within each habitat group by summarizing the dispersion of eigenvalues of their respective trait covariance matrix [40]. This measure (V_{rel}) was subsequently converted to an effect size (a Z-score), which quantified the strength of morphological integration [41]. We then performed a series of two-sample tests to compare the strength of morphological integration across habitat groups, following the procedures of [41].

Additionally and for comparison, we repeated these analyses on the set of size-standardized trait data, found as a set of shape ratios [42] where each trait was divided by body size (Supplementary Material).

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To determine the extent to which static and evolutionary allometry were concordant, we evaluated
the directions in morphospace of both the evolutionary (species-level) and static (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 differences in angular direction between the evolutionary trajectory and the
static 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 155 examined changes in the body shape proportions across the phylogeny. Here we treated the 156 head dimensions and limb dimensions separately, as allometric trends could potentially differ 157 between these body regions due to differential functional or selective constraints [43]. Because 158 both the head and limb data were multivariate, we first performed a partial least squares (PLS) analysis [44] of the head traits versus SVL, and the limb traits versus SVL, to describe the 160 direction of maximal covariation between each body region and size. We then measured the 161 mean residuals of each species to the inferred allometric trend, which described the extent to which head and limb proportions of species were greater or smaller than expected for their body 163 size. The species residuals were then mapped on the phylogeny of *Pristurus* using a Brownian 164 motion model of evolution, to qualitatively evaluate shifts in head and limbs proportional-165 ity across the phylogeny for the group. Similarly, within-species patterns of static allometry 166 were visualized by plotting regressions of PLS scores on SVL for both head and limb traits separately. 167

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Finally, to relate within-species allometric trends with patterns of phenotypic diversification in the group we generated a phylomorphospace, based on a phylogenetic principal component analyses

(PCA) on the size-standardized species means obtained from a phylogenetic regression [see 30]. 171 Here, phenotypic similarities among species, relative to their phylogenetic relationships and habitat affiliations, were observed. Additionally, representative specimens (scaled to unit size) were also 173 visually compared to aid in describing these trends. A similar phylomorphospace was constructed 174 for species means not corrected for body size, and the phenotypic disparity among species means in 175 each habitat was calculated and subsequently compared (Supplementary Material). All analyses 176 were conducted in R 4.2.1 [45], using RRPP version 1.3.1 [46,47] and geomorph 4.0.4 [48] for statistical 177 analyses and the tidyverse version 1.3.0 [49], phytools version 0.7-77 [50], and a modified version 178 of the function ggphylomorpho [https://github.com/wabarr/ggphylomorpho] for data manipulation 179 and visualization, as well as scripts written by the authors (Supplementary Material). 180

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 allometry in body form was examined across individuals, a similar pattern was observed ($N_{ind}=687; F=7910.8; Z=9.20;$ P<0.001). Further, the vectors of regression coefficients between the two analyses were highly correlated ($\rho=0.94$) and were oriented in nearly parallel directions in morphospace ($\theta=1.49^{\circ}$). This revealed that the pattern of multivariate allometry across individuals was concordant with macroevolutionary trends of interspecific allometry among species of *Pristurus* across the phylogeny.

Our analyses also exposed significant differences in the allometry of body form among Pristurus 190 utilizing distinct habitats (Table 1). Further, pairwise comparisons of multivariate allometric 191 vectors revealed that patterns of static allometry in each habitat differed significantly from isometry, 192 indicating the presence of multivariate allometry in each (Table 2). Additionally, comparisons 193 identified that ground-dwelling *Pristurus* displayed the most distinct allometric trend as compared 194 with *Pristurus* occupying both the rock and tree habitats (Table 2; Figure 2). Here, regression 195 coefficients of each trait versus size (Supplementary Material) revealed that ground-dwelling 196 Pristurus exhibited strong positive allometry for all head and limb traits (i.e., $\beta > 1.0$). By contrast, 197 rock and tree-dwelling *Pristurus* displayed negative allometry (i.e., $\beta < 1.0$) for head traits, and 198

were more varied for limb traits; with rock-dwelling *Pristurus* displaying positive limb allometry 199 (though less extreme than that of ground-dwelling taxa), whereas most limb traits in tree-dwelling 200 taxa showed negative allometry or near-isometry (Supplementary Material). Thus, these findings 201 implied that larger individuals of ground-dwelling *Pristurus* species displayed disproportionately 202 larger heads and limbs, as compared with large individuals in taxa utilizing other habitat types. 203 Multivariate visualizations of these multivariate allometric trends (Figure 2) confirmed these 204 statistical findings, and indicated that the allometric trajectory in ground-dwelling *Pristurus* was 205 more extreme as compared with either rock- or tree-dwelling *Pristurus*. 206

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Examination of patterns of trait covariation revealed strong levels of morphological integration 208 within each habitat type ($Z_{ground} = 3.97$; $Z_{rock} = 3.72$; $Z_{tree} = 2.15$). Further, two-sample tests 209 revealed that the strength of morphological integration was significantly greater in ground-dwelling 210 Pristurus than either those utilizing rock ($Z_{ground-rock} = 6.59$; $P \ll 0.001$) or tree habitats 211 $(Z_{ground-tree} = 11.17; P \ll 0.001)$. Arboreal Pristurus displayed the lowest levels of integration, 212 which were also significantly lower than in the rock habitat ($Z_{rock-tree} = 7.19$; P << 0.001). When 213 size was accounted for in the data, levels of integration dropped considerably, though the overall 214 pattern and differences among habitat groups remained the same (Supplementary Material). 215

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Comparisons of evolutionary allometry with static allometry in each habitat revealed substantial 217 concordance between allometric trends at these hierarchical levels. Here, vectors of regression 218 coefficients representing static allometry within habitat groups were oriented in very similar 219 directions with the regression vector representing evolutionary allometry, with small pairwise angles 220 between them $(\theta: 2.3^{\circ} \to 5.9^{\circ})$. Subsequent permutation tests indicated no differences between the 221 static allometry vectors and the regression vector representing evolutionary allometry, indicating 222 strong congruence between them (Table 3). Notably, static allometry in ground-dwelling Pristurus was most similar to trends of evolutionary allometry, displaying the smallest angular difference and 224 largest effect size. Thus, static and evolutionary allometry trends were essentially parallel in this 225 group, indicating a direct correspondence between the two. This result implied that phenotypic evolution across species aligned closely with directions of allometric variation within habitat groups 227

228 at the individual level; namely that larger individuals and larger ground-dwelling species exhibited 229 disproportionately larger heads and limbs, while smaller individuals and smaller ground-dwelling 230 species displayed disproportionately smaller heads and limbs.

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Mapping the residuals of species into the phylogeny showed that large ground-dwelling species 232 displayed greater head proportions than large rock-dwelling species, who exhibited smaller heads 233 relative to body size (Figure 3A). Conversely, the opposite pattern was observed when comparing small species utilizing these habitats: ground-dwelling species showed small relative head proportions 235 while rock-dwelling species displayed generally larger head proportions. In contrast, limb shape 236 showed more variable patterns. Although all large ground-dwelling species consistently displayed 237 large relative limb proportions, large rock-dwelling species were more variable in this trait, with 238 P. insignis exhibiting large and P. insignoides small limb proportions. For small species, shifts 239 in relative limb proportions seemed more independent of habitat utilization, since there were 240 differences in limb residuals both within rock- and ground-dwelling species (Figure 3B). Visual 241 inspection of static allometry trends within species (Figure 4) largely confirmed these patterns, 242 illustrating that ground-dwelling species generally displayed steeper allometric patterns in head 243 proportions as compared with rock-dwelling species. Overall there was general concordance across 244 taxa in terms of trends of multivariate allometry, affirming that the association between evolutionary 245 allometry and habitat-based static allometry was robust. 246

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Viewing body shape differentiation in *Pristurus* in phylomorphospace (Figure 5) revealed broad overlap among habitat groups, though arboreal (tree-dwelling) species were somewhat more separated 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 overall body shape. Likewise, the smallest rock-dwelling species were found close to
large ground-dwelling species in morphospace, indicating they displayed similar body shapes as well.

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Finally, when representative specimens were scaled to a similar body size (Figure 6), the 258 consequences of differences in allometric trends on body proportions became apparent. Here, 259 larger ground-dwelling *Pristurus* species displayed disproportionately larger heads and limbs 260 as compared with large *Pristurus* species utilizing other habitat types. Conversely, smaller 261 rock-dwelling species were found to have disproportionately larger heads and limbs as compared 262 with smaller ground-dwelling species. These patterns corresponded closely with those identified 263 in morphospace (Figure 5), where large ground-dwelling species were similar in body form 264 to small rock-dwelling species, while small ground-dwelling species were similar in body form 265 to large rock-dwelling species (Figure 6). Thus, synthesizing the patterns revealed in the phylomorphospace with those from the other analyses revealed that the same body shape could 267 be obtained in different ways, as determined by subtle differences in allometric slope across 268 habitats, combined with body size differences. As such, species with similar body shapes displayed differing overall size, were found in distinct habitats, and exhibited different allometric trends. 270

4. Discussion

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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 274 differentiation across those habitats is essential for our understanding of how natural selection 275 operates and how evolution proceeds. In this study, we evaluated the role of potential drivers of 276 body shape differentiation in the geckos of the genus *Pristurus*. To this end, we compared allometric 277 trends and levels of integration among *Pristurus* occupying distinct habitats, interrogated allometric 278 patterns at both the static and evolutionary levels, and related these trends to diversification in body form. Our findings have several important implications for how ecological specialization, phenotypic 280 integration, and body form evolution along allometric trajectories relate to patterns of phe-281 notypic diversity generally, and the evolution of phenotypic diversification in *Pristurus* in particular. 282

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First, our analyses revealed that patterns of body shape allometry and morphological integration are relatively distinct in ground-dwelling *Pristurus* lizards, as compared with *Pristurus* occupying 285 other habitats. Specifically, we found that multivariate vectors of regression coefficients differed 286 significantly from what was expected under isometry (Table 2) for taxa utilizing all habitat 287 types (ground, rock, tree), indicating that in *Pristurus*, allometric scaling patterns predominate. 288 Further, our interrogation of allometric trends revealed differences between habitat types, where 280 ground-dwelling *Pristurus* displayed steeper (i.e., positively allometric) trends for both head and 290 limb traits, while rock and tree-dwelling taxa displayed shallower (negatively allometric) trends for 291 head traits and more varied patterns for limb proportions. Biologically, these patterns revealed that not only does shape differ between large and small *Pristurus*, but this pattern differs across habitat 293 types. Specifically, large ground-dwelling *Pristurus* present disproportionately larger heads and 294 longer limbs relative to large individuals in other habitats, while small ground-dwelling Pristurus 295 exhibit disproportionately smaller heads and shorter limbs (Figure 3). These findings are consistent 296 with previous work at the macroevolutionary level [30], where large ground species were also found 297 to display disproportionately large heads and long limbs. 298

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Second, our findings revealed that rock-dwelling *Pristurus* show a converse pattern, where smaller 300 individuals displayed relatively larger heads, while larger individuals have smaller heads relative to 301 their body size. These allometric patterns also corresponded with findings at macroevolutionary 302 scales [30], where similar patterns at the species level were observed. Tejero-Cicuéndez et al. [30] 303 also performed habitat ancestral estimation, finding that the rock habitat was the most likely 304 ancestral condition in the group, with subsequent colonization by *Pristurus* of ground habitats. 305 When patterns of allometry are viewed through this lens, it suggests the hypothesis that habitat 306 shifts from rock-dwelling to ground-dwelling incurred a concomitant evolutionary shift in allometric 307 trajectories as well [39]. Indeed, our analyses are consistent with this hypothesis, as allometric 308 trends are inferred to be more rock-like towards the root of the *Pristurus* phylogeny (Figure 3), 300 with subsequent shifts along branches leading to ground-dwelling species. This further suggests 310 that the segregation in body size and shape through differential allometric relationships across 311 habitats responds to adaptive dynamics concerning the colonization of new habitats. Thus, in 312

Pristurus, there is support for the hypothesis that colonization of ground habitats has been a 313 trigger for morphological change [30], as there appears to be a link between shifts in allometric trajectories as a result of habitat-induced selection, and differential patterns of body shape observed 315 across taxa. More broadly, these findings are consistent with prior discoveries in other lizards, 316 where the differential selective pressures imposed by rocky and ground habitats have resulted in 317 the differentiation of head and limb morphology [43,51–53]. Indeed, such phenotypic differences 318 resulting from the effects of habitat-based ecological selection have been extensively documented 319 in reptiles as well as in other vertebrates [60], and our work in *Pristurus* thus contributes to this 320 growing body of literature. 321

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Another important finding of our study was the strong concordance between static allometry across 323 individuals and evolutionary allometry among *Pristurus* species. Our analyses revealed small 324 pairwise angles between static and evolutionary allometry vectors, indicating that allometric trends 325 at these two hierarchical levels were oriented in similar directions and were essentially parallel. As 326 such, size-associated changes in body shape among individuals were predictive of evolutionary shifts 327 across taxa at higher macroevolutionary scales. This in turn, suggests that body shape evolution in 328 Pristurus follows an allometric line of least resistance [61]. In other empirical systems, a similarly 329 tight correspondence between static and evolutionary allometry has also been observed [61–65]. 330 though the trend is not universal across all taxa or traits [see 19.66]. Nonetheless, when such trends 331 are present, they imply that allometric trajectories impose a prevailing influence on the magnitude, 332 direction, and rate of phenotypic change across the phylogeny. Our work in *Pristurus* contributes to 333 the growing literature on this topic, and suggests that perhaps such patterns may be more widespread. 334

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Given the observation that static and evolutionary allometry in *Pristurus* are so concordant, an obvious question is: why might this be the case? One possible explanation is that when genetic covariation remains relatively constant, selection on body size will generate an evolutionary allometric trajectory along the trend described by static allometry [67,68]. Here, allometry effectively acts as a constraint on evolutionary change, as size-associated shape changes at one hierarchical level are linked to changes at another level [63,66,69]. Further, when this is the case, one may also

expect high levels of phenotypic integration in traits associated with body size changes. Indeed, our 342 analyses reveal precisely this pattern in *Pristurus*, with the highest levels of integration in the group (ground-dwelling) whose static allometry is most similar to that of evolutionary allometry. Thus, 344 our results reveal that patterns of trait covariation are more constrained in ground-dwelling species, 345 such that their differences in body form are most likely found along the primary allometric axis. 346 When viewed in this light, integration and allometry may thus be interpreted as potential drivers 347 that facilitate morphological change, as they provide a phenotypic pathway through adaptive lines 348 of least resistance that enable rapid evolutionary changes in particular phenotypic directions but 349 not in others [22,27]. The fact that ground-dwelling species in Pristurus have been found to have 350 the widest phenotypic disparity, greatest range of body sizes, and highest rates of morphological 351 evolution [30] are all consistent with this hypothesis, and suggest that in this group, integration 352 describes the path of morphological evolution along allometric lines of least resistance. 353

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Finally, interpreting the observed patterns of phenotypic integration and allometry relative to habitat-355 specific differences helps to shed light on the possible pathways by which phenotypic diversity in 356 Pristurus has evolved. For instance, prior work on this system [30] revealed that the colonization 357 of new ecological habitats elicited strong ecological selection and phenotypic responses. This was 358 particularly true of the invasion of ground habitats, where ground-dwelling species displayed the 359 largest variation in body size in the genus. This observation implies some level of ecological selection 360 on body size. In lizards, the ecological context in which species exist is known to play a pervasive role 361 in body size evolution [70–72], as it does in other animal groups [73–77]. While to date this has not 362 been thoroughly explored in *Pristurus*, the evolutionary patterns revealed by our analyses suggest 363 that the body size diversity in this clade conforms, at least in part, with patterns expected under ecological selection on body size. Intriguingly, such patterns are not only observed in ground- and 365 rock-dwelling taxa, but also in arboreal species, whose restricted phenotypic diversity in both size 366 and shape (Figures 3 & 5) is consistent with strong ecological selection in the arboreal habit [78,79]. 367 Furthermore, our study identified the presence of strong integration and allometric trajectories, 368 such that evolutionary changes in body size elicit corresponding changes in body shape. However, 369 these trends differed significantly across habitats, implying that, at evolutionary scales, these trends 370

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 shape. 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

- 1. Collar DC, Schulte JA, O'Meara BC, Losos JB. 2010 Habitat use affects morphological diversification in dragon lizards. *Journal of Evolutionary Biology* **23**, 1033–1049. (doi:10.1111/j.1420-9101.2010.01971.x)
- 2. Kaliontzopoulou A, Carretero MA, Adams DC. 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. (doi:10.1111/jeb.12540)
- 33. Price SA, Friedman ST, Wainwright PC. 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. (doi:10.1098/rspb.2015.1428)
- Martinez CM, Friedman ST, Corn KA, Larouche O, Price SA, Wainwright PC. 2021
 The deep sea is a hot spot of fish body shape evolution. *Ecology Letters* **24**, 1788–1799.

 (doi:10.1111/ele.13785)
- 5. Kolmann MA, Marques FPL, Weaver JC, Dean MN, Fontenelle JP, Lovejoy NR. 2022 Ecological and phenotypic diversification after a continental invasion in neotropical freshwater stingrays. *Integrative and Comparative Biology* **62**, 424–440. (doi:10.1093/icb/icac019)
- 6. Losos JB. 1992 The evolution of convergent structure in Caribbean *Anolis* communities.

 Systematic Biology 41, 403–420. (doi:10.1093/sysbio/41.4.403)
- 7. Schluter D, McPhail JD. 1992 Ecological character displacement and speciation in sticklebacks.

 The American Naturalist 140, 85–108. (doi:10.1086/285404)
- 8. Arnold SJ. 1983 Morphology, performance, fitness. *American Zoologist* **23**, 347–361. (doi:10.1093/icb/23.2.347)
- 9. Losos JB. 2009 Lizards in an evolutionary tree: Ecology and adaptive radiation of anoles.

 University of California Press.
- 10. Mahler DL, Ingram T, Revell LJ, Losos JB. 2013 Exceptional convergence on the macroevolutionary landscape in island lizard radiations. Science **341**, 292–295. (doi:10.1126/science.1232392)

- Schluter D, Grant PR. 1984 Determinants of morphological patterns in communities of Darwin's finches. *The American Naturalist* **123**, 175–196. (doi:10.1086/284196)
- 12. Grant PR, Grant BR. 2006 Evolution of character displacement in darwin's finches. Science 313, 224–226. (doi:10.1126/science.1128374)
- 13. Reaney AM, Bouchenak-Khelladi Y, Tobias JA, Abzhanov A. 2020 Ecological and morphological determinants of evolutionary diversification in Darwin's finches and their relatives.

 Ecology and Evolution 10, 14020–14032. (doi:10.1002/ece3.6994)
- 405 14. Albertson RC, Kocher TD. 2001 Assessing morphological differences in an adaptive trait: A landmark-based morphometric approach. *Journal of Experimental Zoology* **289**, 385–403.

 (doi:10.1002/jez.1020)
- Urban S, Gerwin J, Hulsey CD, Meyer A, Kratochwil CF. 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. (doi:10.1002/ece3.8568)
- Jastrebski CJ, Robinson BW. 2004 Natural selection and the evolution of replicated trophic polymorphisms in pumpkinseed sunfish (*Lepomis gibbosus*). Evolutionary Ecology Research
 6, 285–305.
- Berner D, Adams DC, Grandchamp A-C, Hendry AP. 2008 Natural selection drives patterns of lake-stream divergence in stickleback foraging morphology. *Journal of Evolutionary Biology* 21, 1653–1665. (doi:10.1111/j.1420-9101.2008.01583.x)
- 18. Stuart YE et al. 2017 Contrasting effects of environment and genetics generate a continuum of parallel evolution. Nature Ecology and Evolution 1, 158. (doi:10.1038/s41559-017-0158)
- 19. Klingenberg CP, Zimmermann M. 1992 Static, ontogenetic, and evolutionary allometry: A multivariate comparison in nine species of water striders. American Naturalist 140, 601–620.
 (doi:10.1086/285430)
- Wagner G, Altenberg L. 1996 Perspective: Complex adaptations and the evolution of evolvability. Evolution 50, 967–976. (doi:10.1111/j.1558-5646.1996.tb02339.x)
- 21. Schluter D. 1996 Adaptive radiation along genetic lines of least resistance. *Evolution* **50**, 1766–1774. (doi:10.1111/j.1558-5646.1996.tb03563.x)

- Felice RN, Randau M, Goswami A. 2018 A fly in a tube: Macroevolutionary expectations for integrated phenotypes. *Evolution* **72**, 2580–2594. (doi:10.1111/evo.13608)
- Wagner GP, Zhang J. 2011 The pleiotropic structure of the genotype—phenotype map: The evolvability of complex organisms. Nature Reviews Genetics 12, 204–213. (doi:10.1038/nrg2949)
- 425 24. Klingenberg CP, Marugán-Lobón J. 2013 Evolutionary covariation in geometric morphometric data: Analyzing integration, modularity, and allometry in a phylogenetic context. Systematic Biology 62, 591–610. (doi:10.1093/sysbio/syt025)
- 25. Goswami A, Smaers JB, Soligo C, Polly PD. 2014 The macroevolutionary consequences of phenotypic integration: From development to deep time. *Philosophical Transactions of the Royal Society B: Biological Sciences* 369, 20130254. (doi:10.1098/rstb.2013.0254)
- 26. Goswami A, Randau M, Polly PD, Weisbecker V, Verity Bennett C, Hautier L, Sánchez-Villagra MR. 2016 Do developmental constraints and high integration limit the evolution of the marsupial oral apparatus? *Integrative and Comparative Biology* **56**, 404–415. (doi:10.1093/icb/icw039)
- 27. Navalón G, Marugán-Lobón J, Bright JA, Cooney CR, Rayfield EJ. 2020 The consequences of craniofacial integration for the adaptive radiations of Darwin's finches and Hawaiian honeycreepers. *Nature Ecology & Evolution* 4, 270–278. (doi:10.1038/s41559-019-1092-y)
- Jolicoeur P. 1963 The multivariate generalization of the allometry equation. *Biometrics* **19**, 497–499. (doi:10.2307/2527939)
- 29. Bookstein FL. 2022 Dimensions of morphological integration. *Evolutionary Biology* **49**, 342–372. (doi:10.1007/s11692-022-09574-0)
- 30. 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. *Proceedings of the Royal Society B: Biological Sciences* 288, 20211821.

 (doi:10.1098/rspb.2021.1821)

- Tejero-Cicuéndez H, Patton AH, Caetano DS, Šmíd J, Harmon LJ, Carranza S. 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, 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).
- 33. Freedman D, Lane D. 1983 A nonstochastic interpretation of reported significance levels.

 Journal of Business & Economic Statistics 1, 292–298. (doi:10.2307/1391660)
- Collyer ML, Adams DC. 2007 Analysis of two-state multivariate phenotypic change in ecological studies. *Ecology* 88, 683–692. (doi:10.1890/06-0727)
- 35. Collyer ML, Sekora DJ, Adams DC. 2015 A method for analysis of phenotypic change for phenotypes described by high-dimensional data. *Heredity* 115, 357–365. (doi:10.1038/hdy.2014.75)
- Adams DC, Collyer ML. 2009 A general framework for the analysis of phenotypic trajectories in evolutionary studies. *Evolution* **63**, 1143–1154. (doi:10.1111/j.1558-5646.2009.00649.x)
- The state of the s
- 38. Drake AG, Klingenberg CP. 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. (doi:10.1098/rspb.2007.1169)
- 39. Adams DC, Nistri A. 2010 Ontogenetic convergence and evolution of foot morphology in European cave salamanders (Family: Plethodontidae). *BMC Evolutionary Biology* **10**, 1–10. (doi:10.1186/1471-2148-10-216)
- 40. Pavlicev M, Cheverud JM, Wagner GP. 2009 Measuring morphological integration using eigenvalue variance. *Evolutionary Biology* **36**, 157–170. (doi:10.1007/s11692-008-9042-7)
- 41. Conaway MA, Adams DC. 2022 An effect size for comparing the strength of morphological integration across studies. *Evolution* **76**, 2244–2259. (doi:10.1111/evo.14595)

- 42. Mosimann JE. 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. (doi:10.1080/01621459.1970.10481136)
- 43. Kaliontzopoulou A, Carretero MA, Llorente GA. 2010 Intraspecific ecomorphological variation: Linear and geometric morphometrics reveal habitat-related patterns within *Podarcis bocagei* wall lizards. *Journal of Evolutionary Biology* 23, 1234–1244. (doi:10.1111/j.1420-9101.2010.01984.x)
- 44. Rohlf FJ, Corti M. 2000 Use of two-block partial least-squares to study covariation in shape.

 Systematic Biology 49, 740–753. (doi:10.1080/106351500750049806)
- 45. R Core Team. 2022 R: A language and environment for statistical computing. Version 4.2.1.

 Vienna, Austria: R Foundation for Statistical Computing. See https://www.R-project.org/.
- 46. Collyer ML, Adams DC. 2018 RRPP: An R package for fitting linear models to high-dimensional data using residual randomization. *Methods in Ecology and Evolution* **9**, 1772–1779. (doi:10.1111/2041-210X.13029)
- 47. Collyer ML, Adams DC. 2022 R: RRPP: Linear model evaluation with randomized residuals in a permutation procedure. Vsn. 1.3.1. Vienna, Austria: R Foundation for Statistical Computing. See https://CRAN.R-project.org/package=RRPP.
- 48. Baken EK, Collyer ML, Kaliontzopoulou A, Adams DC. 2021 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. (doi:10.1111/2041-210X.13723)
- 49. Wickham H et al. 2019 Welcome to the tidyverse. Journal of Open Source Software 4, 1686.

 (doi:10.21105/joss.01686)
- 50. Revell LJ. 2012 Phytools: An R package for phylogenetic comparative biology (and other things). Methods in Ecology and Evolution 3, 217–223. (doi:10.1111/j.2041-210X.2011.00169.x)
- 51. 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)

- 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.

 Journal of Biogeography 44, 1694–1704. (doi:10.1111/jbi.12929)
- 53. Foster KL, Garland T, Schmitz L, Higham TE. 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. (doi:10.1093/biolinnean/bly146)
- 54. Busschau T, Boissinot S. 2022 Habitat determines convergent evolution of cephalic horns in vipers. Biological Journal of the Linnean Society 135, 652–664. (doi:10.1093/biolinnean/blac008)
- Hipsley CA, Müller J. 2017 Developmental dynamics of ecomorphological convergence in a transcontinental lizard radiation. *Evolution* **71**, 936–948. (doi:10.1111/evo.13186)
- Watanabe A, Fabre AC, Felice RN, Maisano JA, Müller J, Herrel A, Goswami A. 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. (doi:10.1073/pnas.1820967116)
- 57. 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)
- 58. Samuels JX, Hopkins SSB. 2017 The impacts of Cenozoic climate and habitat changes on small mammal diversity of North America. Global and Planetary Change 149, 36–52. (doi:10.1016/j.gloplacha.2016.12.014)
- 59. Friedman ST, Collyer ML, Price SA, Wainwright PC. 2022 Divergent Processes Drive Parallel Evolution in Marine and Freshwater Fishes. Systematic biology 71, 1319–1330. (doi:10.1093/sysbio/syab080)
- Navalón G, Bjarnason A, Griffiths E, Benson RBJ. 2022 Environmental signal in the evolutionary diversification of bird skeletons. *Nature* **611**, 306–311. (doi:10.1038/s41586-022-05372-y)

- Marroig G, Cheverud JM. 2005 Size as a line of least evolutionary resistance: Diet and adaptive morphological radiation in New World monkeys. *Evolution* **59**, 1128–1142. (doi:10.1111/j.0014-3820.2005.tb01049.x)
- 62. Firmat C, Lozano-Fernández I, Agustí J, Bolstad GH, Cuenca-Bescós G, Hansen TF, Pélabon C. 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. (doi:10.1098/rstb.2014.0057)
- 503 Voje KL, Hansen TF, Egset CK, Bolstad GH, Pélabon C. 2014 Allometric constraints and the evolution of allometry. *Evolution* **68**, 866–885. (doi:10.1111/evo.12312)
- 505 64. Brombacher A, Wilson PA, Bailey I, Ezard THG. 2017 The breakdown of static and evolutionary allometries during climatic upheaval. *The American Naturalist* **190**, 350–362. (doi:10.1086/692570)
- 65. Marcy AE, Guillerme T, Sherratt E, Rowe KC, Phillips MJ, Weisbecker V. 2020 Australian rodents reveal conserved cranial evolutionary allometry across 10 million years of murid evolution. The American Naturalist 196, 755–768. (doi:10.1086/711398)
- Voje KL, Bell MA, Stuart YE. 2022 Evolution of static allometry and constraint on evolutionary allometry in a fossil stickleback. *Journal of Evolutionary Biology* 35, 423–438.
 (doi:10.1111/jeb.13984)
- 511 67. Lande R. 1979 Quantitative genetic analysis of multivariate evolution, applied to brain-body size allometry. *Evolution* **33**, 402–416. (doi:10.2307/2407630)
- 513 68. Lande R. 1985 Size and scaling in primate biology. In (ed WL Jungers), pp. 21–32. Plenum

 Press.
- Pélabon C, Bolstad GH, Egset CK, Cheverud JM, Pavlicev M, Rosenqvist G. 2014 On the relationship between ontogenetic and static allometry. The American Naturalist 181,
 195–212. (doi:10.1086/668820)
- 70. Meiri S. 2008 Evolution and ecology of lizard body sizes. *Global Ecology and Biogeography*17, 724–734. (doi:10.1111/j.1466-8238.2008.00414.x)

- 519 71. James SE, M'closkey RT. 2004 Patterns of body size and habitat use in a lizard assemblage.

 Ecoscience 11, 160–167. (doi:10.1080/11956860.2004.11682820)
- 72. Tamar K, Mitsi P, Simó-Riudalbas M, Tejero-Cicuéndez H, Al-Sariri T, Carranza S. 2019
 Systematics, biogeography, and evolution of *Pristurus minimus* (Squamata, Sphaerodactylidae) with the discovery of the smallest Arabian vertebrate. *Systematics and Biodiversity* 17,
 349–366. (doi:10.1080/14772000.2019.1614694)
- 523 73. Bergmann C. 1847 Über die verhaltnisse der warmeokonomie der thiere zu ihrer grosse.

 Göttinger Studien 1, 595–708.
- 74. Calder WA. 1983 Ecological scaling: Mammals and birds. Annual Review of Ecology and

 Systematics 14, 213–230. (doi:10.1146/annurev.es.14.110183.001241)
- 75. Peters RH. 1983 The ecological implications of body size. Cambridge University Press.

528

- 529 76. LaBarbera M. 1989 Analyzing body size as a factor in ecology and evolution. Annual Review of Ecology and Systematics 20, 97–117. (doi:10.1146/annurev.es.20.110189.000525)
- 77. Olson VA, Davies RG, Orme CDL, Thomas GH, Meiri S, Blackburn TM, Gaston KJ, Owens IPF, Bennett PM. 2009 Global biogeography and ecology of body size in birds. *Ecology Letters* 12, 249–259. (doi:10.1111/j.1461-0248.2009.01281.x)
- 533 78. Baken EK, Adams DC. 2019 Macroevolution of arboreality in salamanders. *Ecology and Evolution* 9, 7005–7016. (doi:10.1002/ece3.5267)
- 535 79. Baken EK, Mellenthin LE, Adams DC. 2021 Is salamander arboreality limited by broad-scale climatic conditions? *PLoS ONE* **16**, e0255393. (doi:10.1371/journal.pone.0255393)

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Table 1: Multivariate analysis of covariance describing variation in body form in Pristurus.

	Df	SS	MS	Rsq	F	Z	Pr(>F)
SVL	1	516.04	516.04	0.92	10188.70	9.49	0.001
habitat	2	6.22	3.11	0.01	61.39	9.32	0.001
SVL:habitat	2	3.97	1.99	0.01	39.23	7.08	0.001
Residuals	681	34.49	0.05	0.06			
Total	686	560.72					

Table 2: Pairwise comparisons of multivariate static allometry for each habitat group. Comparisons with the vector of multivariate isometry are included. Displayed are: pairwise angular differences (θ_{12}) , their associated effect sizes $(Z_{\theta_{12}})$, and significance levels obtained via permutation (RRPP).

	Ground	Rock	Tree	Isometry	
Angle					
Ground	0				
Rock	6.629	0			
Tree	8.095	3.628	0		
Isometry	5.034	5.901	7.189	0	
Effect Size					
Ground	0				
Rock	7.004	0			
Tree	2.1	-0.408	0		
Isometry	7.673	7.357	1.779	0	
P-value					
Ground	1				
Rock	0.001	1			
Tree	0.027	0.673	1		
Isometry	0.001	0.001	0.042	1	

Table 3: Pairwise comparisons of multivariate evolutionary allometry versus static allometry for each habitat group. Pairwise angular differences between evolutionary and static allometry (θ_{ES}) , their associated effect sizes $(Z_{\theta_{ES}})$, and significance levels are displayed.

	θ_{ES}	$Z_{\theta_{ES}}$	P-value
Evol. vs. Ground	2.37	-4.26	1.000
Evol. vs. Rock	4.55	0.87	0.191
Evol. vs. Tree	5.96	0.21	0.405

Figures

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- Figure 1. Linear Measurements used in this study. SVL = snout-vent length, TL = trunk length,

 HL = head length, HW = head width, HH = head height, Lhu = humerus length, Lun = ulna

 length, Lfe = femur length, Ltb = tibia length [for details see 30].
- Figure 2. Plot of regression scores and predicted lines representing the relationship between linear body measurements and size (SVL). Individuals are colored by habitat use: ground (beige), rock (dark purple), and tree (magenta). Isometric trend represented by the dashed line.
- Figure 3. Traitgrams showing the evolution of body size (SVL) through time based on the
 phylogenetic tree of *Pristurus*. Colors represent an evolutionary mapping of residuals from
 phylogenetic regressions describing the relationship of (A) head morphology versus body size,
 and (B) limb proportions versus body size (see text for descriptions). Species names are colored
 by habitat use: ground (beige), rock (dark purple), and tree (magenta).
- Figure 4. Patterns of static allometry for each species for head traits (upper panel) and limb traits

 (lower panel). Species are separated by their habitat groups and colored by the magnitude of

 their regression slope (purple: steeper slopes, yellow: shallower slopes).
- Figure 5. Phylomorphospace of *Pristurus*, based on residuals from a phylogenetic regression of body measurements on size (SVL). Species means are colored by habitat use: ground (beige), rock (dark purple), and tree (magenta). Large and small rock-dwelling and ground-dwelling are highlighted with darker colors to highlight their differentiation and relative positions in morphospace.
- Figure 6. Representative specimens (based on real specimens) from large and small *Pristurus*species, colored by habitat use: ground (beige) and rock (dark purple). Specimens are scaled
 to a common body size (SVL, gray rectangles) to emphasize the relative differences in limb
 and head proportions. Relatively slender-headed and short-limbed species shown on the left.
 Original scale shown as the gray bar.



Figure 1: Linear Measurements used in this study. SVL = snout-vent length, TL = trunk length, HL = head length, HW = head width, HH = head height, Lhu = humerus length, Lun = ulna length, Lfe = femur length, Ltb = tibia length (for details see Tejero-Cicuéndez et al. 2021a).

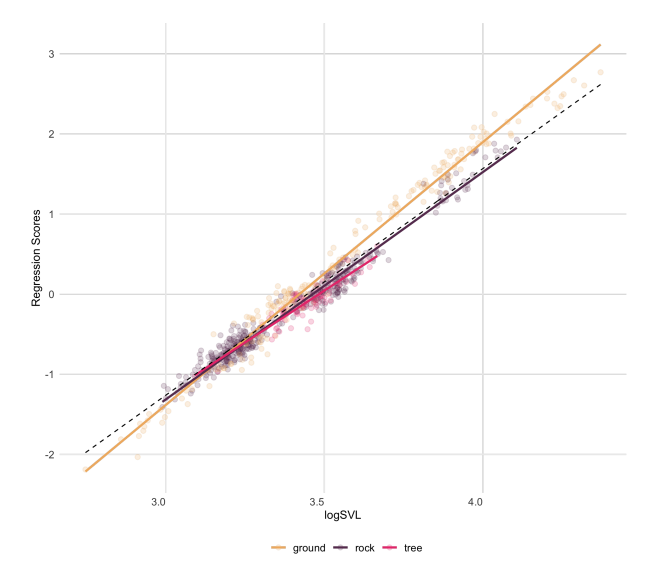


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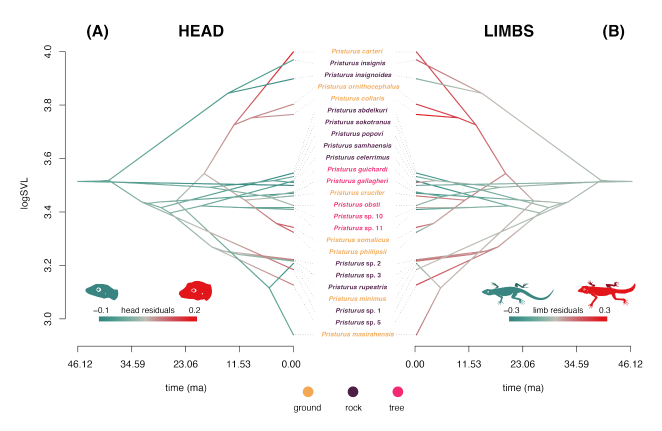


Figure 3: Traitgrams showing the evolution of body size (SVL) through time based on the phylogenetic tree of *Pristurus*. Colors represent an evolutionary mapping of residuals from phylogenetic regressions describing the relationship of (A) head morphology versus body size, and (B) limb proportions versus body size (see text for descriptions). Species names are colored by habitat use: ground (beige), rock (dark purple), and tree (magenta).

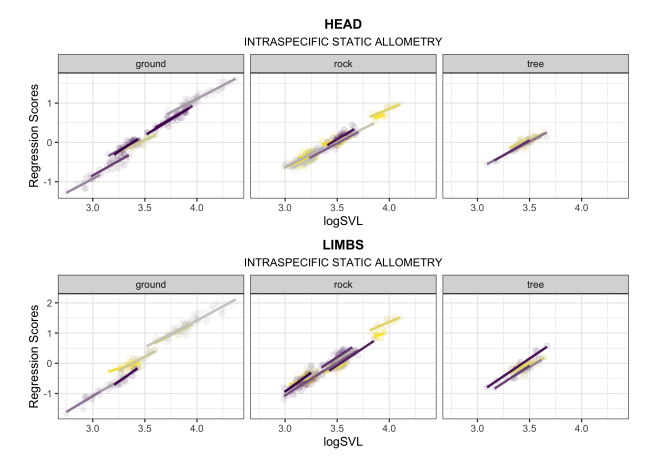


Figure 4: Patterns of static allometry for each species for head traits (upper panel) and limb traits (lower panel). Species are separated by their habitat groups and colored by the magnitude of their regression slope (purple: steeper slopes, yellow: shallower slopes).

Phylomorphospace (largest and smallest species highlighted) *P.* sp. 1 P. sp. 2 0.1 P. carteri 0 *P.* sp. 5 P. ornithocephalus *P.* sp. 3 0.0 P. collaris P. rupestris P. insignoides -0.1P. minimus P. insignis P. masirahensis -0.3-0.2-0.10.0 0.1 0.2 PC1 ground rock tree

Figure 5: Phylomorphospace of *Pristurus*, based on residuals from a phylogenetic regression of body measurements on size (SVL). Species means are colored by habitat use: ground (beige), rock (dark purple), and tree (magenta). Large and small rock-dwelling and ground-dwelling are highlighted with darker colors to highlight their differentiation and relative positions in morphospace.



Figure 6: Representative specimens (based on real specimens) from large and small *Pristurus* species, colored by habitat use: ground (beige) and rock (dark purple). Specimens are scaled to a common body size (SVL, gray rectangles) to emphasize the relative differences in limb and head proportions. Relatively slender-headed and short-limbed species shown on the left. Original scale shown as the gray bar.