- The role of habitat and evolutionary allometry in the morphological
- differentiation of *Pristurus* geckos (OR: Evolution along allometric
- lines of least resistance: Morphological differentiation in *Pristurus*
- 4 geckos)
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22 Short Title: XXX

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- et al. 2021b)). R-scripts are available at XXX.

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# 35 Abstract

asdf

#### 37 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 habitats (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 50 (e.g., Losos 1992; Schluter and McPhail 1992), this convergent evolution is treated as strong 51 evidence of adaptation. Indeed the ecomorphological paradigm (sensu Arnold 1983) is predicated, in part, on such cases, which emphasize the strong association between the phenotypic traits that 53 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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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
the covariance between body parts resulting from body size variation (i.e., allometry). It has
long been recognized that the interrelationships among traits can have a strong influence on how
phenotypic evolution proceeds, as 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 (Jolicoeur 1963; Bookstein
2022), allometric trends could be considered the quintessential measure 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 (Tejero-Cicuéndez et al. 2021a) 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 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 exploiting rocky habitats (the most common ecological feature in *Pristurus*), 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 shape is related to differences in body size and whether habitat specialization has an impact in this shape-size relationship.

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In this study, we employed a combination of multivariate morphometric and phylogenetic comparative analysis 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 allometric patterns differed across species occupying distinct ecological habitats. 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. Overall our results demonstrate that the interplay between ecological specialization and differing allometric trajectories in species with disparate body size may have a determinant role in shaping the phenotypic evolution and hence in adaptive dynamics in this clade.

#### Materials and Methods

#### 102 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 104 work on this system (Tejero-Cicuéndez et al. 2021a, 2022), and are briefly described here. First 105 we used a time-dated, molecular phylogeny that included all members of the genus Pristurus, 106 including several currently undescribed taxa. The tree was estimated in a Bayesian framework, 107 using five mitochondrial markers, six nuclear markers, and 21 calibration points (for details see 108 Tejero-Cicuéndez et al. 2022). Next we categorized each species as belonging to one of three 109 ecological groups (ground, rock, or tree), based on descriptions of habitat use found in the literature 110 (see Tejero-Cicuéndez et al. 2021a). Finally, we obtained a phenotypic data set containing body 111 size (snout-vent length: SVL) and eight linear measurements (Figure 1) that described overall 112 body form: trunk length (TrL), head length (HL), head width (HW), head height (HH), humerus 113 length (Lhu), ulna length (Lun), femur length (Lfe), and tibia length (Ltb) (Tejero-Cicuéndez et 114 al. 2021a). We restricted our study to those species represented by nine or more individuals; 115 resulting in a dataset of 687 individuals from 25 species (invidivuals per species:  $\mu = 27$ ; min = 116 9, max = 56). Species in the phenotypic dataset were then matched to the phylogeny, which was 117 subsequently pruned to arrive at the final topology. All measurements were log-transformed prior 118 to statistical analyses. Additional details regarding data collection and formal descriptions of each 119 linear measurement may be found in the original sources (see Tejero-Cicuéndez et al. 2021a, 2022). 120

The data are found on DRYAD: https://doi.org/10.5061/dryad.xwdbrv1f6 (Tejero-Cicuéndez et al. 2021b).

#### 123 Statistical and Comparative Analyses

We conducted a series of analyses to interrogate allometric trends, patterns of integration, 124 and macroevolutionary changes in allometry, relative to differentiation in body form. First we 125 characterized evolutionary allometry in the genus by performing a phylogenetic multivariate regression of body form on size, using the species means as data. We then performed an analogous 127 procedure at the individual level, regressing body form on size using our entire dataset. From both 128 the species-level (phylogenetic) and the individual-level regression models, we obtained the set of 129 regression coefficients, and calculated the difference between them to describe the extent to which 130 patterns of allometry at the individual level were concordant with evolutionary allometric trends 131 across species. 132

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Next we used the individual dataset to determine whether allometric trends in body form differed 134 across habitat groups. This was accomplished by performing a multivariate analysis of covariance, 135 with body size (SVL), habitat, and  $SVL \times habitat$  as model effects. Significance was evaluated 136 using 999 iterations of a permutation procedure, where residuals from a reduced model were 137 randomly permuted in each permutation (RRPP), model statistics were recalculated, and used to 138 generate empirical null sampling distributions to evaluate the observed test statistics (following 139 Freedman and Lane 1983; Collver and Adams 2007; Collver et al. 2015). We then compared the 140 multivariate allometric vectors for each habitat group by calculating pairwise differences in their 141 angular direction in morphospace, and evaluating these relative to empirical sampling distributions obtained through RRPP (Collver and Adams 2007, 2013; Adams and Collver 2009). Patterns 143 of multivariate allometry relative to body size were visualized via regression scores (Drake and 144 Klingenberg 2008) and predicted lines (Adams and Nistri 2010), based on the coefficients and fitted values from the linear model described above. 146

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We then examined changes in static allometry across the phylogeny. Here we treated the head

dimensions and limb dimensions separately, as static allometric trends could potentially differ 149 between these body regions due to differential functional or selective constraints (Kaliontzopoulou et al. 2010). Because both the head and limb data were multivariate, we first performed a partial 151 least squares (PLS) analysis (Rohlf and Corti 2000) of the head traits versus SVL, and the limb 152 traits versus SVL, to describe the direction of maximal covaration between each body region 153 and size. PLS scores from each analysis were obtained, and species-specific slopes describing 154 the extent of head and limb allometry within each species were extracted from an analysis of 155 covariance modeled as:  $PLS1_{head} \sim SVL * species$  and  $PLS1_{limb} \sim SVL * species$  respectively. 156 The species-specific allometric slopes were then mapped on the phylogeny of *Pristurus* using 157 a Brownian motion model of evolution, to qualitatively evaluate shifts in allometry across the 158 phylogeny for the group (for a similar approach see Adams and Nistri 2010). 159

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Next, because allometry describes the extent to which traits covary with size and with each other 161 (i.e., integration), we conducted an analysis of integration. Here we characterized the extent of 162 morphological integration in body form for individuals within each habitat group. Integration 163 was estimated by summarizing he dispersion of eigenvalues of the trait covariance matrix (sensu 164 Pavlicev et al. 2009). This measure  $(V_{rel})$  was subsequently converted to an effect size (a Z-score), 165 which quantified the strength of morphological integration (see Conaway and Adams 2022). We 166 then performed a series of two-sample tests to compare the strength of morphological integration 167 across habitat groups. Additionally and for comparison, we repeated these analyses on the set of 168 size-standardized trait data, found as a set of shape ratios (sensu Mosimann 1970) where each trait was divided by body size (Supplemental Information). 170

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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 the size-standardized species means obtained from a phylogenetic regression (see Tejero-Cicuéndez et al. 2021a). Here, phenotypic similarities among species, relative to their phylogenetic relationships and habitat affiliations, were observed. A similar phylomorphospace was constructed for species means not corrected for body size, and the phenotypic disparity among species means in each habitat was calculated and subsequently

compared (Supplemental Information). All analyses were conducted in R 4.2.1 (R Core Team 2022), using RRPP version 1.3.1 (Collyer and Adams 2018, 2022) and geomorph 4.0.4 (Baken et al. 2021), and scripts written by the authors (available at **XXX**).

#### $_{\scriptscriptstyle{181}}$ Results

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Using phylogenetic regression, we found significant evolutionary allometry in body form across 182 species ( $N_{sp}=25;\;F=217.9;\;Z=5.53;\;P<0.001$ ). Likewise, when allometry in body form 183 was examined across individuals, a similar pattern was observed ( $N_{ind}=687;\ F=7910.8;$ 184 Z = 9.20; P < 0.001). Further, the vectors of regression coefficients between the two analyses 185 were highly correlated ( $\rho = 0.94$ ) and were oriented in nearly parallel directions in morphospace 186  $(\theta = 1.49^{\circ})$ . This revealed that the pattern of multivariate allometry across individuals was 187 concordant with macroevolutionary trends of interspecific allometry among species of Pristurus 188 across the phylogeny. 189

Our analyses also exposed significant differences in the allometry of body form among Pristurus 191 utilizing distinct habitats (Table 1). Here, comparisons of multivariate allometric vectors identified 192 that ground-dwelling *Pristurus* displayed a distinct allometric trend as compared with *Pristurus* 193 occupying both the rock and tree habitats (Table 2). In addition, allometric patterns in both 194 rock and tree habitats were similar to the multivariate line of isometry (Figure 2), while patterns 195 of multivariate allometry in ground-dwelling *Pristurus* was decidely steeper. Inspection of the regression coefficients for each trait (Supplemental Information) confirmed the steeper allometric 197 coefficients for all head and limb traits in ground-dwelling *Pristurus* as compared with rock and 198 tree-dwelling taxa, corroborating this result. Taken together, these findings implied that larger 199 individuals of ground-dwelling *Pristurus* species displayed proportionately larger heads and limbs, 200 as compared with large individuals in taxa utilizing other habitat types. A visualization of 201 multivariate allometric trends (Figure 2) confirmed these statistical findings, and indicated that 202 the allometric trajectory in ground-dwelling animals was more extreme as compared with either 203 rock or tree-dwelling *Pristurus*. 204

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When static allometric patterns were mapped on the phylogeny, traitgrams elucidated that 206 changes in allometric trends were not concentrated to specific regions of the phylogeny (Figure 3). 207 Rather, increases and decreases in allometry of both the head traits and the limb traits occurred repeatedly (see also Supplemental Information). When these patterns were viewed relative to body 209 size, large ground-dwelling species tended to display steeper head allometry as compared to large 210 rock-dwelling species, who displayed shallower head allometry (Figure 3A). Likewise, a similar 211 pattern was observed when comparing small species utilizing these habitats. In contrast, large 212 ground-dwelling species exhibited intermediate patterns of limb allometry, and small species in gen-213 eral were more variable in such patterns (Figure 3B). Inspection of traitgrams thus revealed some 214 degree of discordance in patterns of allometry across body regions; a pattern evidenced by the rela-215 tively low correlation between slopes representing head and limb allometry respectively ( $\rho = 0.42$ ). 216

Examination of patterns of trait covariation revealed strong levels of morphological integration 218 within each habitat type ( $Z_{ground} = 3.97$ ;  $Z_{rock} = 3.72$ ;  $Z_{tree} = 2.15$ ). Further, two-sample tests 219 revealed that the strength of morphological integration was significantly greater in ground-dwelling 220 Pristurus than either those utilizing rock ( $Z_{Groung-Rock} = 6.59$ ;  $P \ll 0.001$ ) or tree habitats  $(Z_{Groung-Tree} = 11.17; P \ll 0.001)$ . Pristurus utilizing tree habitats displayed the lowest levels 222 of integration, which were also significantly less than in the rock habitat ( $Z_{Rock-Tree} = 7.19$ ; 223  $P \ll 0.001$ ). When size was accounted for in the data, levels of integration dropped considerably, though the overall pattern and differences among habitat groups remained the same (Supplemental 225 Information). 226

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Finally, when body shape differentiation in *Pristurus* was viewed in phylomorphospace (Figure 4), we found 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 (Supplemental Information). 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 smaller rock-dwelling species were found close to large ground-dwelling species in morphospace, indicating they displayed similar body shapes as well. Thus, synthesizing the patterns revealed in the phylomorphospace with those identified in our earlier analyses revealed a complex interplay between body shape, body size, habitat use, morphological integration, and multivariate allometry; where species with similar body shapes displayed differing overall size, were found in distinct habitats, and exhibited different allometric trends.

#### Discussion

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- First paragraph; restate topic/questions (see last par of Intro); summarize general findings
  ...
  - something about linking allometry and phenotypic diversification?

The relationship between certain phenotypic traits and the organisms' environment is a central 246 paradigm in evolutionary biology. In this context, disentangling the causes of phenotypic differen-247 tiation is essential to understand how natural selection operates. In this study, we evaluated the 248 role of potential drivers of body shape differentiation in the geckos of the genus Pristurus. In par-249 ticular, we investigated how the interplay of ecological specialization, phenotypic integration and 250 allometric trends have shaped the morphological evolution in this radiation of Afro-Arabian geckos. 251 Our results show that allometric trends and integration patterns are different across habitats, with 252 ground-dwelling species having the steepest multivariate allometric slope and also the strongest 253 morphological integration. These patterns are also different across body parts, with decoupled trends between head and limb proportions. Additionally, we found that changes in static allomet-255 ric trends are not restricted to specific regions of the phylogeny, but rather they show multiple independent increases and decreases following common dynamics within habitat groups. Overall, 257 these results suggest that the interplay between allometric and integration patterns is a fundamental factor to explain the morphological evolution across a variety of habitats, which is consistent with the theoretical expectation that different ecological contexts impose distinct selective pressures triggering phenotypic change.

- result 1: allometry; overall trend among species nearly identical to that among individuals.

  Thus, patterns of static and evolutionary allometry are consistent in *Pristurus...* Something else about this
- Patterns of multivariate allometry in body form calculated from individuals were found to be nearly identical to those calculated from per-species means in Pristurus geckos. Specifically, the vectors of regression coefficients of the two analyses are virtually parallel ( $\theta = 1.49^{\circ}$ ), indicating that the evolutionary allometry is not substantially different whether measured with individual measurements or with species means in this genus.
- 270 ??We also explored patterns of static allometry to compare them among species and with general trends of evolutionary allometry.
- result 2: Allometry differs among habitat groups: 'steeper' allometry in Ground-dwelling (implication: proportionately larger heads and longer limbs in species at larger body sizes).
- When we compared multivariate allometric slopes of species occupying different habitats, we found 274 that, while rock-dwelling and arboreal species do not significantly differ from the isometric trend, 275 ground-dwelling species have a steeper slope which is statistically different from isometry. This 276 means that large ground-dwelling *Pristurus* present proportionally larger heads and longer limbs 277 relative to other large species, while small species in the ground have proportionally smaller heads 278 and shorter limbs (Figure residuals traitgrams). This is consistent with previous results on the 279 morphological evolution of Pristurus (Tejero-Cicuéndez et al. 2021), where large ground species 280 were indeed found to have also proportionally large heads and long limbs. This suggests that 281 the segregation in body size and shape through differential allometric relationships across habitats 282 responds to adaptive dynamics concerning the colonization of ground habitats, and perhaps with 283 a particular interest of hard ground environments inhabited by the largest ground-dwelling species 284 (including the largest of the genus, P. carteri), which has already been suggested to be the main 285 driver of the morphological evolution in this genus (Tejero-Cicuéndez et al. 2021). This points

toward the existence of a specialized form of *Pristurus* geckos adapted to hard grounds (e.g., **some**definition of hard ground vs. soft grounds??), illustrating the ecomorphological relationships
in the genus with a rather conspicuous 'ecomorph' (see Figure X for an example of the hard-ground
ecomorph, *P. carteri*).

- result 3: Morphological integration differs among habitat groups. Strongest in ground-dwelling; weakest in tree-dwelling. SOME MEANING (combined with allometric trend implies that patterns of trait covariation are more constrained within ground-dwelling.... Thus, differences in body form are most likely found along this primary axis... (harken to Schluter evolution along lines of least resistance)
  - Additionally, rank-order of magnitude of integration across habitat groups corresponds with the range of body sizes in each: ground-dwelling display the largest size-range, while tree-dwelling the least (Supp. Information). On the one hand this matches the expectation that much of the integration observed in *Pristurus* is the result of allometric trends.... And the fact that levels of integration drop so precipitously when data are size-standardized are in accord with this interpretation. Nevertheless, when size is accounted for, the rank-order of magnitudes of integration remain the same, implying that ground-dwelling *Pristurus* are still relatively constrained in patterns of trait covariation as compared with the other two groups.
  - This notion was further supported when viewing the phylomorphospace of the species means not adjusted for size (SI). Here (and not surprisingly), PC1 is dominated by size, with small species at one end and larger species at the other. More importantly however, is that the disparity among species utilizing different habitats differed significantly in this space. Here, ground-dwelling displayed significantly greater phenotypic disparity than did the other groups (SI).

Similarly, when analyzing patterns of morphological integration, we found important differences among habitat groups: ground-dwelling species present the strongest integration, which in turn is weakest in arboreal species. Morphological integration occurs when different body parts coevolve,

and has been suggested as an evolutionary constraint, since it restricts the specific lines along which 315 integrated structures are allowed to vary (**REF**). Weaker integration levels (i.e., modularity), on the contrary, might facilitate morphological evolution by allowing a less constrained exploration of 317 the morphospace (**REF**). However, integration might also be interpreted as a potential driver of 318 morphological change, since it may provide a phenotypic pathway through adaptive lines of least 319 resistance that enable rapid evolutionary processes (Navalón et al. 2020). In this context, our 320 results on allometry and integration suggest that patterns of trait covariation are more constrained 321 in ground-dwelling species, such that their differences in body form are most likely found along 322 this primary axis. The fact that ground species in Pristurus have been found to have the widest 323 phenotypic disparity and highest rates of morphological evolution (Tejero-Cicuéndez et al. 2021) is 324 consistent with the idea that integration patterns are acting to facilitate morphological evolution 325 along lines of least resistance. 326

• result 4: morphospace: Thus there was a reciprocal relationship between body shape and body size across ground-dwelling and rock-dwelling species. SOMEHOW TIE THIS TO integration (DCA pondering this one)

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• one interesting... head vs. (correlation of head vs. limb slopes: 0.42. Pretty low. Implies some sort of differential something here, resulting in distinct allometric patterns for these two body regions. SImilar to Antigoni's work (and refs therein). IMPLICATION: tie this into integration/modularity. Less integrated across the whole organism, and more modular...

Future studies should examine this.

Another insightful result is the low correlation between head and limbs in their allometric slopes,
which implies different evolutionary trajectories for these two body regions. This is likely to happen
when different parts of an organism are subjected to different functional pressures (e.g., head
evolution might be mainly influenced by diet while limb evolution might respond more tightly
to the substrate used by the species), resulting in a decoupling of their respective morphological
change. Ultimately, the combination of selective pressures upon which organisms evolve may lead
to differential levels of integration across different body parts, with certain structures coevolving
in a similar (i.e., integrated) manner and others in a segregated (i.e., modular) way. This, in turn,

may have fundamental implications for the extent of morphological diversification within clades, and can be key to describe the phenotypic divergence observed across the tree of life. Future and more in-depth studies on the evolution of different body parts in *Pristurus* and other lizards, including for instance finer phenotypic data and comprehensive ecological information, may allow for discerning the functional drivers of head and limb evolution.

- In conclusion... -Synthesizing these patterns together ... (summarize: steeper allometry, higher integration, greater disparity in body size and body form all in ground-dwelling species). TOgether the patterns uncovered in our study imply that phenotypic diversification among ground-dwelling *Pristurus* follows tightly along its allometric trajectory, as evidenced by the higher disparity and stronger morphological integration.... some reference back to Goswami 'fly in a tube' paper. Thus, *Pristurus body forms appear to diversify along* allometric\* lines of least resistance.... (Schluter ref again)
  - On the relationship between static and evolutionary allometry

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There is no general consensus about the relationship between the three types of allometry: on-356 togenetic (allometry during the development), static (allometry among individuals at the same 357 developmental stage), and evolutionary (allometry across populations or species). This, in turn, 358 is reflected in the broadly ambiguous interpretation of allometric patterns in the literature, with 359 an often uncertain distinction between allometry as an evolutionary constraint and allometry as functional optimization resulting from natural selection (Pélabon et al. 2014, Voje et al. 2014). 361 Even though testing these alternative hypotheses is beyond the scope of this work, our results do 362 suggest that static and evolutionary allometry are very similar in *Pristurus* geckos, which could be 363 explained by low evolvability of allometry, but also by the effect of relatively homogeneous selective pressures at different scales. Further analyses, for instance including a broader phylogenetic context 365 and developmental assessments, are needed to illuminate the relationships between different levels 366 of allometric trends.

### 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 289:385–403.
- Arnold, S. J. 1983. Morphology, performance, fitness. American Zoologist 23:347–361.
- Baken, E. K., M. L. Collyer, A. Kaliontzopoulou, and D. C. Adams. 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.
- 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 49:342–372.
- <sup>384</sup> 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.
- <sup>390</sup> 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.

- <sup>392</sup> Collyer, M. L., and D. C. Adams. 2022. R: RRPP: Linear model evaluation with randomized
- residuals in a permutation procedure. Vsn. 1.3.1. R Foundation for Statistical Computing,
- Vienna, Austria.
- 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:(In Press).
- Drake, A. G., and C. P. Klingenberg. 2008. The pace of morphological change: Historical trans-
- formation of skull shape in st bernard dogs. Proceedings of the Royal Society B: Biological
- sciences 275:71–76.
- Felice, R. N., M. Randau, and A. Goswami. 2018. A fly in a tube: Macroevolutionary expectations
- for integrated phenotypes. Evolution 72:2580–2594.
- Freedman, D., and D. Lane. 1983. A nonstochastic interpretation of reported significance levels.
- Journal of Business & Economic Statistics 1:292–298.
- 406 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.
- 412 Grant, P. R., and B. R. Grant. 2006. Evolution of character displacement in darwin's finches.
- 413 Science 313:224–226.
- 414 Jastrebski, C. J., and B. W. Robinson. 2004. Natural selection and the evolution of replicated
- trophic polymorphisms in pumpkinseed sunfish (*Lepomis qibbosus*). Evolutionary Ecology Re-
- search 6:285–305.
- Jolicoeur, P. 1963. The multivariate generalization of the allometry equation. Biometrics 19:497–

- 418 499.
- 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.
- 422 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.
- 425 Klingenberg, C. P., and J. Marugán-Lobón. 2013. Evolutionary covariation in geometric mor-
- phometric data: Analyzing integration, modularity, and allometry in a phylogenetic context.
- Systematic Biology 62:591–610.
- Kolmann, M. A., F. P. L. Marques, J. C. Weaver, M. N. Dean, J. P. Fontenelle, and N. R. Love-
- joy. 2022. Ecological and phenotypic diversification after a continental invasion in neotropical
- freshwater stingrays. Integrative and Comparative Biology 62:424–440.
- 431 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. Sys-
- tematic Biology 41:403–420.
- 435 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.
- 437 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.
- 439 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
- 441 65:930-945.
- 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.
- Pavlicev, M., J. M. Cheverud, and G. P. Wagner. 2009. Measuring morphological integration using
   eigenvalue variance. Evolutionary Biology 36:157–170.
- 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.
- 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 darwins finches and their relatives.
- 454 Ecology and Evolution 10:14020–14032.
- Rohlf, F. J., and M. Corti. 2000. Use of two-block partial least-squares to study covariation in shape. Systematic Biology 49:740–753.
- Schluter, D. 1996. Adaptive radiation along genetic lines of least resistance. Evolution 50:1766–458 1774.
- Schluter, D., and P. R. Grant. 1984. Determinants of morphological patterns in communities of darwins 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.
- 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.
- 467 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.
- Wagner, G., and L. Altenberg. 1996. Perspective: Complex adaptations and the evolution of evolvability. Evolution 50:967–976.
- Wagner, G. P., and J. Zhang. 2011. The pleiotropic structure of the genotypephenotype map: The evolvability of complex organisms. Nature Reviews Genetics 12:204–213.

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

	Df	SS	MS	Rsq	F	Z	Pr(>F)
svl	1	516.036559	516.0365588	0.9203096	10188.69842	9.490057	0.001
habitat	2	6.218510	3.1092552	0.0110902	61.38957	9.322480	0.001
svl:habitat	2	3.974307	1.9871536	0.0070879	39.23464	7.077264	0.001
Residuals	681	34.491245	0.0506479	0.0615124			
Total	686	560.720622					

Table 2: Pairwise comparisons of multivariate allometry vectors. Effect sizes  $(Z_{\theta_{12}})$  based on pairwise differences in angular direction are below the diagonal, and their corresponding significance levels are above diagonal. Significant values in bold.

	Ground	Rock	Tree
Ground	0	0.001	0.001
Rock	6.872	0	0.261
Tree	3.657	0.649	0

## **Figures**

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).

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 regression slopes
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. 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.

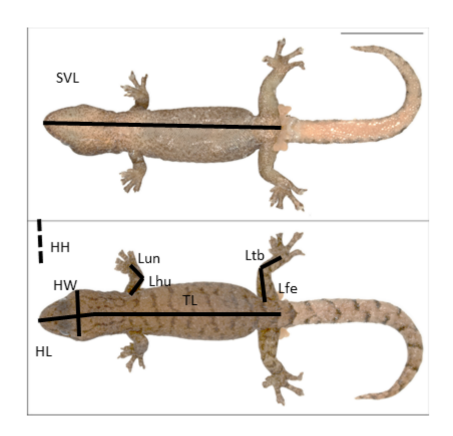


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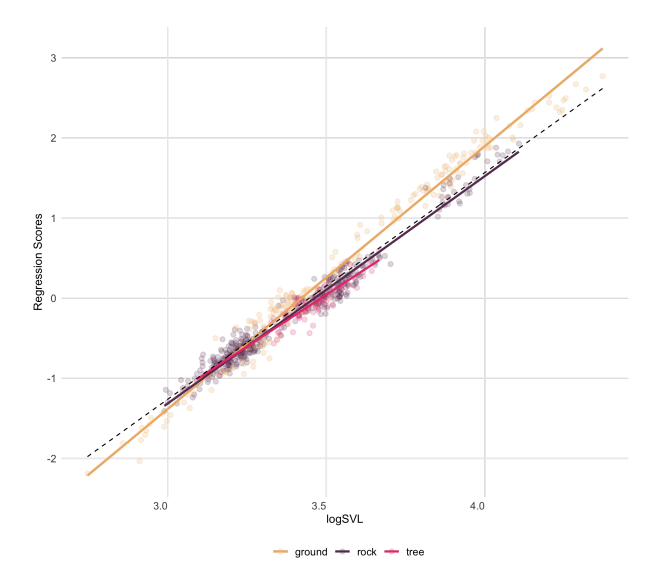


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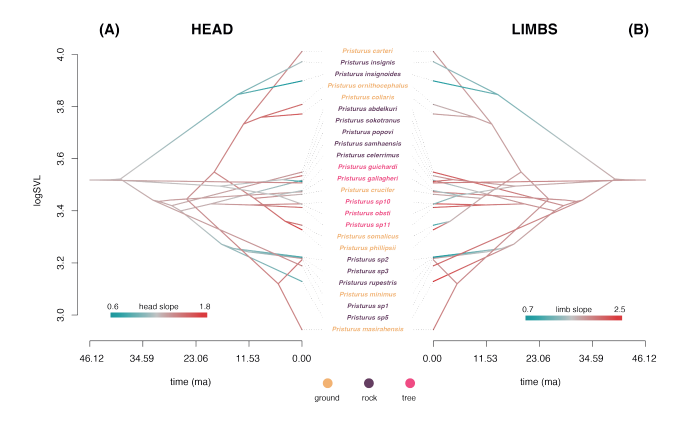


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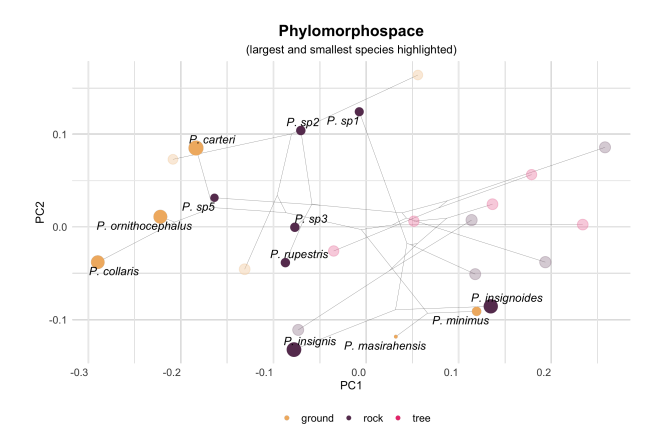


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