Supplementary Material: Evolution along allometric lines of least resistance: Morphological differentiation in *Pristurus* geckos

# 1: R-script for Article Computations and Flow of Allometry Operations

Below is an R-script that may be used to reproduce all statistical analyses found in the paper. The data are found on DRYAD: (doi:10.5061/dryad.xwdbrv1f6 (Tejero-Cicuéndez et al. 2021)). Additional scripts used to generate publication-ready plots, and scripts for the additional plots, are found below.

```
libs <- c('geomorph', 'RRPP', 'phytools', 'geiger', 'tidyverse')</pre>
easypackages::libraries(libs)
# 0: Data Prep
data0 <- read.table('Analyses/data/morpho/morpho_pristurus.csv', sep = ';',</pre>
                dec = '.', header = TRUE, stringsAsFactors = TRUE)
  sp.to.keep <- names(which(table(data0$species) >= 5))
data <- data0[data0$species %in% sp.to.keep, ]</pre>
  data$species <- droplevels(data$species)</pre>
  data$SVL <- log(data$SVL)</pre>
shape <- as.matrix(log(data[, 8:ncol(data)]))</pre>
rdf <- rrpp.data.frame(svl = data$SVL, shape = shape,
            habitat = data$habitat_broad, species = data$species)
tree0 <- read.nexus('Analyses/data/phylogeny/pristurus_tree_final.nex')</pre>
LS.mns <- pairwise(lm.rrpp(shape~species, data = rdf, iter=0),
                    groups = rdf$species)$LS.means[[1]]
sz.mn <- tapply(rdf$svl,rdf$species,mean)</pre>
hab.mn <- as.factor(by(rdf$habitat,rdf$species,unique))</pre>
levels(hab.mn) <- levels(rdf$habitat)</pre>
tree <- treedata(phy = tree0, data = LS.mns)$phy</pre>
C <- vcv.phylo(tree)</pre>
SVL.resid <- resid(lm(data$SVL~data$species))</pre>
shape.resid <- resid(lm(shape~data$species))</pre>
rdf2 <- rrpp.data.frame(svl = SVL.resid, shape = shape.resid,
                          habitat = data$habitat_broad,
                          species = data$species)
```

```
# 1: Evolutionary Allometry & Static Allometry
allom.sp <- lm.rrpp(LS.mns~sz.mn, Cov = C)
allom.ind <- lm.rrpp(shape~svl, data = rdf2)</pre>
anova(allom.sp)
anova(allom.ind)
M <-rbind(coef.sp <- allom.sp$LM$gls.coefficients[2,],</pre>
        coef.ind <- allom.ind$LM$coefficients[2,])</pre>
acos(RRPP:::vec.cor.matrix(M))*180/pi
# 2: Comparison of multivariate allometry among habitat types
fit.hab <- lm.rrpp(shape~svl*habitat, data = rdf2)</pre>
  anova(fit.hab)
# 2A: Compare habitat vectors versus isometry and to each other
  #H_0: isometry as common slope model
mn.sz <- tapply(rdf2$svl,rdf2$habitat,mean)</pre>
mn.shape <- rowsum(rdf2$shape, rdf2$habitat)/as.vector(table(rdf2$habitat))</pre>
coef.iso \leftarrow c(1,1,1,1,1,1,1,1)
intercepts <- mn.shape - t(tcrossprod(coef.iso,mn.sz))</pre>
X <- model.matrix(~rdf2$svl+rdf2$habitat)</pre>
b <- rbind(intercepts[1,],coef.iso,intercepts[2,]-intercepts[1,],</pre>
           intercepts[3,]-intercepts[1,])
preds <- X%*%b
E.iso <- rdf2$shape - preds
perms <- RRPP:::perm.index(n = fit.hab$LM$n, iter = 999)</pre>
slopes <- list()</pre>
for(j in 1:1000){
  slopes[[j]] <- pairwise(lm.rrpp((preds+E.iso[perms[[j]],]) ~ rdf2$svl*rdf2$habitat,</pre>
          iter=0), groups = rdf2$habitat,covariate = rdf2$svl)$slopes[[1]]
slp.ang <- lapply(1:1000, function(j)</pre>
      acos(RRPP:::vec.cor.matrix(rbind(slopes[[j]],coef.iso)))*180/pi)
slp.hab.obs <- slp.ang[[1]]</pre>
slp.Z <- RRPP:::effect.list(slp.ang)</pre>
slp.P <- RRPP:::Pval.list(slp.ang)</pre>
slp.hab.obs
slp.Z
slp.P
# 2B: Compare evolutionary and static (habitat) allometry
  #H_0: common slope isometry
slp.ang.ev <- lapply(1:1000, function(j)</pre>
   acos(RRPP:::vec.cor.matrix(rbind(coef.evol,slopes[[j]])))*180/pi)
```

```
slp.hab.ev.obs <- slp.ang.ev[[1]]</pre>
slp.Z.ev <- RRPP:::effect.list(slp.ang.ev)</pre>
slp.P.ev <- RRPP:::Pval.list(slp.ang.ev)</pre>
slp.hab.ev.obs
slp.Z.ev
slp.P.ev
res \leftarrow cbind(slp.hab.ev.obs[-1,1],slp.Z.ev[-1,1],slp.P.ev[-1,1])
colnames(res) <- c("Angle", "Effect Size", "P-value")</pre>
rownames(res) <- c("Ev vs. Ground", "Ev vs. Rock", "Ev vs. Tree")
res
# 3: Map allometry slopes on phylogeny
head.scores <- plot(lm.rrpp(shape[, c(2:4)]~ rdf$svl),
      type = "regression", predictor = rdf$svl, reg.type = "RegScore")$RegScore
limb.scores <- plot(lm.rrpp(shape[, c(5:8)]~ rdf$svl),</pre>
      type = "regression", predictor = rdf$svl, reg.type = "RegScore")$RegScore
coef.head <- lm.rrpp(head.scores ~ rdf$svl*rdf$species)$LM$coefficients</pre>
coef.limb <- lm.rrpp(limb.scores ~ rdf$svl*rdf$species)$LM$coefficients</pre>
head.slp <- coef.head[grep('svl', rownames(coef.head)), ]</pre>
  head.slp[-1] \leftarrow head.slp[-1] + head.slp[1]
limb.slp <- coef.limb[grep('svl', rownames(coef.limb)), ]</pre>
  limb.slp[-1] \leftarrow limb.slp[-1] + limb.slp[1]
names(limb.slp) <- names(head.slp) <- levels(rdf$species)</pre>
cor(head.slp,limb.slp)
plot(head.slp,limb.slp)
contMap(tree = tree, x = head.slp, outline = FALSE)
cm.limb <- contMap(tree = tree, x = limb.slp, outline = FALSE)</pre>
# 4: Compare Integration
lindims.gp <- lapply( split( shape.resid[,1:ncol(shape.resid)],</pre>
                 rdf$habitat), matrix, ncol=ncol(shape.resid))
Vrel.gp <- Map(function(x) integration.Vrel(x), lindims.gp)</pre>
c(Vrel.gp$ground$ZR, Vrel.gp$rock$ZR, Vrel.gp$tree$ZR)
out <- compare.ZVrel(Vrel.gp$ground, Vrel.gp$rock, Vrel.gp$tree)</pre>
summary(out)
# 5: phylomorphospace of size-standardized data (residuals)
shape.res <- residuals(allom.sp)</pre>
pca.w.phylo <- gm.prcomp(shape.res, phy = tree)</pre>
plot(pca.w.phylo, phylo = TRUE, pch = 21, bg = 'black',
```

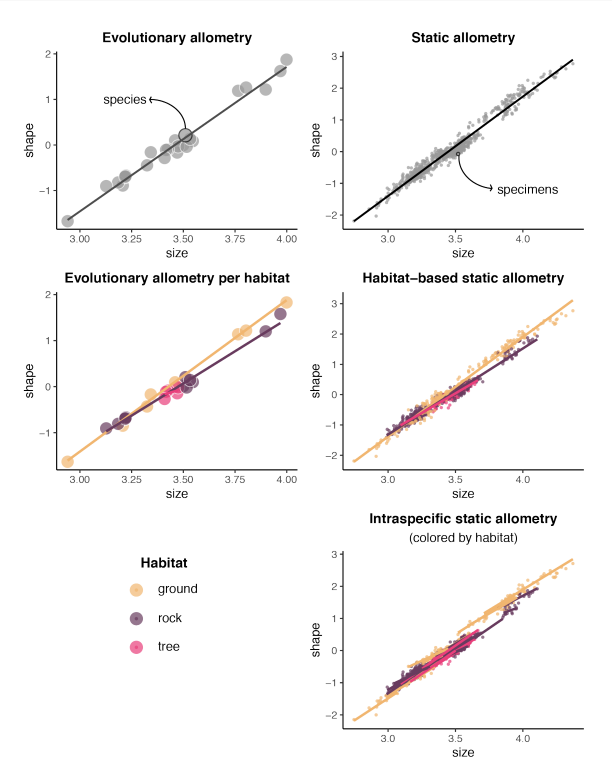


Figure 1: Visual depiction of flow of operations for interrogating levels of allometry. Colors designate the different habitat groups.

# 2: Additional Analyses and Visualizations

Here we provide additional analyses which complement those found in the article.

### Inspection of Regression Coefficients (Slopes)

Here are the regression slopes for each habitat group, found from our linear model. These display differences in allometry among groups, variable by variable.

```
## Warning in treedata(phy = tree0, data = LS.mns): The following tips were not found in 'data
## Pristurus_adrarensis
## Pristurus_flavipunctatus
## Pristurus_sp12
## Pristurus_sp4
## Pristurus_sp9
fit.hab <- lm.rrpp(shape~svl*habitat, data = rdf2)</pre>
pw.hab1 <- pairwise(fit.hab, groups = rdf2$habitat, covariate = rdf2$svl)</pre>
  slp.hab <- pw.hab1$slopes[[1]] #slopes by habitat</pre>
slp.hab
##
                          HL
                                     HW
                                                                             Lfe
               TrL
                                               HH
                                                        Lhu
                                                                   Lun
## ground 1.106798 0.7871177 0.8312497 0.8447515 0.8925517 0.8899040 0.9737474
## rock
          1.103346 0.7551718 0.7883580 0.6385472 1.0113416 0.9793085 0.9503882
          1.099800 0.7296915 0.8185439 0.7595467 1.1163738 1.0655536 0.9351170
## tree
##
               Ltb
## ground 0.893884
## rock
          1.009734
## tree
          1.012326
```

### Traitgrams of Individual Trait Allometry

In the main article we provided traitgrams of allometric slopes for composites of head traits (Figure 3A) and limb traits (Figure 3B). Here we provide traitgrams for the allometric relationship of each body trait separately. As in the main article, traitgrams are visualized from an evolutionary mapping of body size (SVL), and the color represents changes in the allometric slope for each phenotypic trait, found from an evolutionary mapping of the species-level slopes under a Brownian motion model of evolution.

Here we see that there are differential allometric dynamics for some morphological variables. Most notably, large species inhabiting rocky habitats show an increase in the allometric slope for the length of the tibia (Ltb), while the rest of the limb segments (Lhu, Lun, and Lfe) have a decreasing trend. Conversely, these species show virtually identical trends for the head shape variables (HL, HW, and HH), where the size increase has occurred together with a reduction of the allometric slope. Overall, large ground-dwelling species show opposite allometric trends to those of the large rock dwellers in most variables. However, it might be interesting to notice that one of these large ground species, *Pristurus ornithocephalus* (the fourth largest species of the genus, and the second largest ground species) presents unique allometric tendencies relative to the other large ground species, both for all head variables and some of the limb variables (e.g., Lhu and Lfe). This might be reflecting species-specific ecological dynamics, and more detailed data and analyses (e.g., geometric morphometrics) could shed light on this morphological pattern.

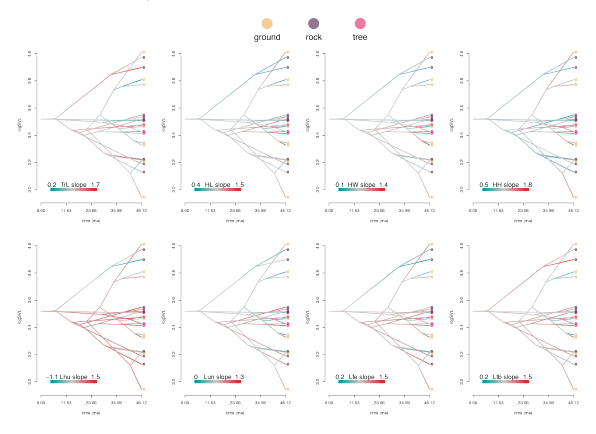


Figure 2: Traitgrams of regression slopes for each phenotypic variable. Colors at the tips designate habitat groups.

## Evolutionary Mapping of Head & Limb Allometry

In the main article, allometric trends in both head dimensions were mapped onto the phylogeny under a Brownian motion model of evolution to discern macroevolutionary changes across the phylogeny. These were visualized on traitgrams, where body size differences were optimized (main article: Fig. 3). Here we present evolutionary mappings of allometric trends individually, so that increases and decreases in allometric slopes across the phylogeny are more readily interpreted. A summary of these patterns was described in the main article.

Briefly, these plots show that changes in allometry were not concentrated to particular regions of the phylogeny, but rather displayed both increases and decreases in allometry of both the head traits and the limb traits occurred repeatedly in this group.

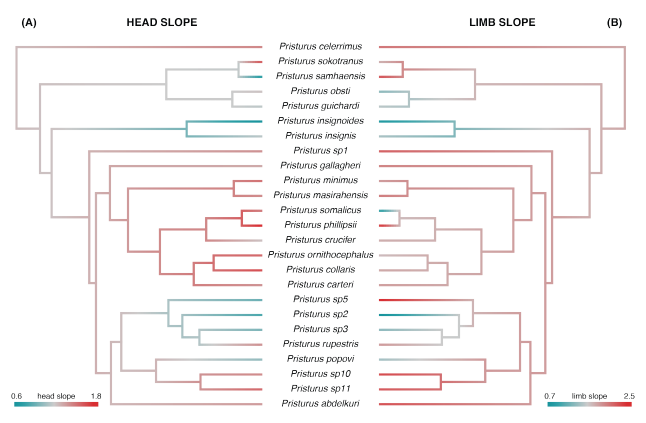


Figure 3: Evolutionary mapping of regression slopes describing the relationship of (A) head morphology versus body size, and (B) limb proportions versus body size.

### Morphological Integration

In the main article, we performed analyses of morphological integration on the set of body traits representing body form. Here we perform the same analysis, using size-standardized data.

```
shape.2 <- shape - rdf$svl</pre>
shape.2 <- resid(lm(shape.2~data$species))</pre>
shape.gp <- lapply( split( shape.2[,1:ncol(shape.2)], rdf$habitat),</pre>
                     matrix, ncol=ncol(shape.2))
Vrel.gp.shp <- Map(function(x) integration.Vrel(x), shape.gp)</pre>
c(Vrel.gp.shp$ground$ZR,Vrel.gp.shp$rock$ZR,Vrel.gp.shp$tree$ZR)
## [1] 1.626647 1.599207 1.042223
out.shp <- compare.ZVrel(Vrel.gp.shp$ground, Vrel.gp.shp$rock, Vrel.gp.shp$tree)
summary(out.shp)
##
## Effect sizes
##
## Vrel.gp.shp$ground
                         Vrel.gp.shp$rock
                                             Vrel.gp.shp$tree
##
           -0.9232867
                               -1.2658429
                                                   -1.1016353
##
## Effect sizes for pairwise differences in rel.eig effect size
##
##
                       Vrel.gp.shp$ground Vrel.gp.shp$rock Vrel.gp.shp$tree
## Vrel.gp.shp$ground
                                 0.000000
                                                    3.953883
                                                                      1.409394
## Vrel.gp.shp$rock
                                  3.953883
                                                    0.000000
                                                                      1.402138
## Vrel.gp.shp$tree
                                  1.409394
                                                    1.402138
                                                                     0.000000
```

## Phylomorphospace and Disparity

##

## ##

## P-values

## Vrel.gp.shp\$ground

## Vrel.gp.shp\$rock

## Vrel.gp.shp\$tree

In the main article we presented a representation of phylomorphospace using size-standardized shape variables obtained from a non-phylogenetic regression. Here we present the phylomorphospace generated using size-standardized species means obtained from a phylogenetic regression (which shows very similar results) along with a test of disparity of these size-standardized data. Additionally, we present a similar plot for the unadjusted species means. In addition, we calculate the phenotypic disparity among species in each habitat group, and compare these using permutation.

1.000000e+00

7.689287e-05

1.587186e-01

Vrel.gp.shp\$ground Vrel.gp.shp\$rock Vrel.gp.shp\$tree

7.689287e-05

1.000000e+00

1.608740e-01

0.1587186

0.1608740

1.000000

This is the phylomorphospace with size-corrected species means after a phylogenetic regression (very similar to the one resulting from the non-phylogenetic regression, see Fig. 5 in the main text).

```
shape.res <- residuals(allom.sp)
pca.w.phylo <- gm.prcomp(shape.res, phy = tree)
# see main script for the code that generates this figure:</pre>
```

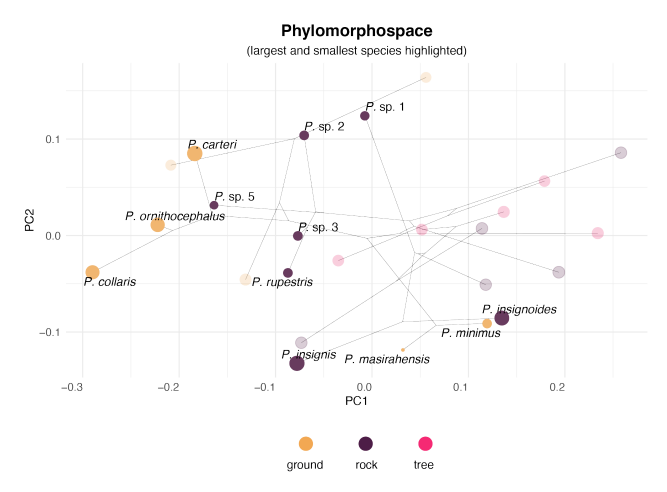


Figure 4: Phylomorphospace of Pristurus, based on residuals from a phylogenetic regression of body measurements on size (SVL). Species means are colored by habitat use: ground (beige), rock (dark purple), and tree (magenta). Large and small rock-dwelling and ground-dwelling are highlighted with darker colors to highlight their differentiation and relative positions in morphospace. Point size is proportional to mean species body size. 79% of the total variation is displayed in the first two PC axes (PC1 = 62.8%; PC2 = 16.3%).

First, we show that in the size-standardized morphospace, morphological disparity does not differ across habitat groups. There are no differences.

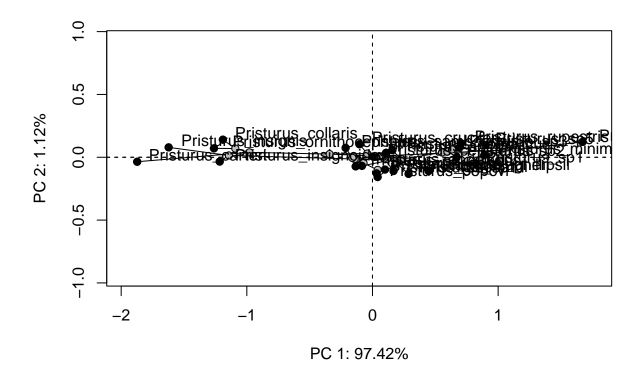
```
fit.lm <- lm.rrpp(shape.res~hab.mn, Cov =C)
PW.lm <- pairwise(fit.lm,groups = hab.mn)
summary(PW.lm, test.type = "var")</pre>
```

```
##
## Pairwise comparisons
##
## Groups: ground rock tree
##
## RRPP: 1000 permutations
##
##
## Observed variances by group
##
##
      ground
                  rock
                            tree
## 0.2938297 0.3015873 0.2260537
##
## Pairwise distances between variances, plus statistics
                         d UCL (95%)
                                              Z Pr > d
## ground:rock 0.007757614 0.1477325 -1.4732960 0.921
## ground:tree 0.067776058 0.1503602 0.2855938 0.421
## rock:tree
               0.075533672 0.1020809 1.0336714 0.158
```

Next we generate a phylomorphospace for the unadusted species means. Not surprisingly, PC1 of the morphospace represents size, and in fact scores on PC1 are highly correlated with SVL ( $\rho = 0.987$ ).

```
pca.w.phylo2 <- gm.prcomp(LS.mns, phy = tree)
cor(sz.mn,-1*pca.w.phylo2$x[,1])</pre>
```

```
## [1] 0.9879806
```



Disparity was estimated from a phylogenetic anova, obtained using RRPP. Here we observed that the ground-dwelling species display greater disparity then do the other two habitat groups.

```
fit.lm <- lm.rrpp(LS.mns~hab.mn, Cov =C)
PW.lm <- pairwise(fit.lm,groups = hab.mn)
summary(PW.lm, test.type = "var")</pre>
```

```
##
## Pairwise comparisons
##
## Groups: ground rock tree
##
##
  RRPP: 1000 permutations
##
##
  Observed variances by group
##
##
##
      ground
                  rock
  18.385078
              7.666641
                        5.635931
##
## Pairwise distances between variances, plus statistics
##
                      d UCL (95%)
                                             Z Pr > d
## ground:rock 10.71844 7.924989
                                    2.12714952 0.013
```

## ground:tree 12.74915 8.616215 2.41223861 0.002 ## rock:tree 2.03071 5.745950 -0.03422601 0.521

# References

Tejero-Cicuéndez, H., M. Simó-Riudalbas, I. Menéndez, and S. Carranza. 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).