Hominoid intraspecific cranial variation mirrors neutral genetic diversity

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Natural selection, developmental constraint, and plasticity have all been invoked as explanations for intraspecific cranial variation in humans and apes. However, global patterns of human cranial variation are congruent with patterns of genetic variation, demonstrating that population history has influenced cranial variation in humans. Here we show that this finding is not unique to Homo sapiens but is also broadly evident across extant ape species. Specifically, taxa that exhibit greater intraspecific cranial shape variation also exhibit greater genetic diversity at neutral autosomal loci. Thus, cranial shape variation within hominoid taxa reflects the population history of each species. Our results suggest that neutral evolutionary processes such as mutation, gene flow, and genetic drift have played an important role in generating cranial variation within species. These findings are consistent with previous work on human cranial morphology and improve our understanding of the evolutionary processes that generate intraspecific cranial shape diversity within hominoids. This work has implications for the analysis of selective and developmental pressures on the cranium and for interpreting shape variation in fossil hominin crania.

Significance

In humans, patterns of cranial variation mirror genetic diversity globally, implicating population history as a key driver of cranial disparity. Here, we demonstrate that the magnitude of genetic diversity within 12 extant ape taxa explains a large proportion of cranial shape variation. Taxa that are more genetically diverse tend to be more cranially diverse also. Our results suggest that neutral evolutionary processes such as mutation, genetic drift, and gene flow are reflected in both genetic and cranial diversity in apes. This work provides a perspective on intraspecific cranial variation in apes which has important implications for interpreting selective and developmental pressures on the cranium and for understanding shape variation in fossil hominin crania.

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Results and Discussion

Using 396 adult crania from 12 hominoid taxa, we measured cranial variation with two different metrics, the average pairwise Procrustes distance (PPD) and the sum of eigenvalues (SEV) (SI Appendix, Table S3). The first measure, PPD, reflects cranial shape differences among members of the same taxon, while SEV measures total shape variation within a taxon. Morphological data consisted of 34 homologous cranial landmarks divided into three units: (i) whole cranium, (ii) cranial vault (neurocranium), and (iii) face. Genetic data included 11 homologous noncoding autosomal loci across the same 12 taxa. Genetic variation was measured with pairwise nucleotide diversity (θ), the number of segregating sites (θs), and the effective population size (Ne) (SI Appendix, Table S4).

We performed 72 ordinary least squares regressions to determine the strength of the relationship between genetic and morphological diversity (SI Appendix, Figs. S4 and S5 and Table S5). Twenty-four regressions were performed for each of the three landmark sets. Within each landmark set, the data were divided into a mixed-sex sample, females only, and males only. For each landmark set, within each sex category, eight regressions were performed to test all combinations of the different genetic and morphological metrics of variation: two with θ (SEV, PPD), two with θs (SEV, PPD), two with Ne from θ (SEV, PPD), and two with Ne from θs (SEV, PPD).

With a significance threshold of P value < 0.05, 57 of 72 regressions are statistically significant. If we apply a more conservative significance threshold of P value < 0.01, 42 of 72 regressions are significant. If we apply the Bonferroni correction to account for the multiple regressions that were performed (0.05/72) P value < 0.00069, 12 of 72 regressions are significant. Results reported within the text below use the P value < 0.01 significance threshold; results from all 72 regressions can be found in SI Appendix, Figs. S4 and S5 and Table S5.

In the mixed-sex sample for the whole cranium, θ accounted for 80% of the variance with SEV (P = 0.0001) (Fig. 1A). In the female whole-cranium dataset, no results were significant at the 0.01 P value threshold. In the male whole-cranium dataset, θ accounted for 77% of cranial variance with SEV (P = 0.0002).

For the cranial vault, all regressions were statistically significant (P value < 0.01) for the mixed-sex sample and for males. For the mixed-sex cranial vault results, θ accounted for 61% of the variance with SEV (P = 0.0028) (Fig. 1B). For females, θ accounted for only 47% of the variance with SEV (P = 0.0143). In the male cranial vault dataset θ accounted for 61% of variance with SEV (P = 0.0026).

For the face, θ accounted for 80% of the variance with SEV (P = 0.0001) in the mixed-sex sample (Fig. 1C). None of the eight regressions of the female-only facial dataset were significant. For males, only two of eight regressions with the facial dataset were significant, θ and θs with SEV.

The cranial vault shows a statistically significant relationship with genetic data in 20 of 24 regression analyses (P value < 0.01), while the face does so only in 6 of 24 regression analyses (SI Appendix, Table S5).
The male cranial datasets were statistically significant in 18 of 24 regressions, and the female cranial datasets were statistically significant in only 4 of 24 regressions.

Between the two metrics of morphological variation, SEV and PPD, we see mostly agreement in which regressions are statistically significant, with two exceptions. In the cranial vault, females across all genetic metrics are significant with PPD but are not significant with SEV. In the face mixed-sex sample, all results are significant with SEV but are not significant with PPD (SI Appendix, Table S5).

We find no differences in which regressions are statistically significant between the two different measures of nucleotide variation (\(\pi\), \(\theta_w\)). The same combinations of variables (sex, landmark set, and morphological metric) are statistically significant for both \(\pi\) and \(\theta_w\) within males, with only one exception: SEV in the male faces dataset which is significant with \(\pi\) and \(\theta_w\) but is not significant with \(\pi\).

**Natural Selection or Neutrality in Hominoid Cranial Evolution?** A number of important implications arise from these results. First, our results suggest that the population processes that generate genetic variation at neutral loci explain a portion of the magnitude of cranial variation within each taxon, especially in the cranial vault. This result is consistent with patterns found in humans and indicates that neutral cranial evolution may not be unique to humans but rather may be part of a broader pattern in extant hominoids. In modern humans, the face tracks population history less closely than the rest of the cranium. The face may be more influenced by environmental pressures such as climate and bone remodeling due to masticatory strains. In humans, populations that live in extremely cold environments show departures from neutrality in aspects of nasal morphology, cranial breadth, and vault size and shape (2, 7, 30). In the primate cranium, mechanical strain has been shown to inflate variation in phenotypically plastic regions, especially the mandible and the, face (12). Thus, the less consistent relationship between facial shape and genetic variation here may be partially driven by mechanical stress and phenotypic plasticity in hominoids. The cranial vault, however, is not subject to the same functional strains from mastication, is less variable than the face and mandible, and shows a statistically significant relationship with genetic data in the majority of the analyses here.

Despite the congruence between genetic and cranial data here, it is unlikely that only neutral evolutionary forces are acting to impact intraspecific cranial variation within hominoid taxa. Natural processes such as mutation and genetic drift act in concert with developmental and selective pressures, and disentangling the differential effects of these processes on morphological and genetic evolution remains a long-standing challenge in evolutionary biology (31). It is important to note that the intraspecific focus of our study differs from recent work looking at how drift and selection impact variation and diversification between species (22, 23). It is possible that neutral population processes and levels of genetic variation explain a portion of cranial variation observed within the taxa included here but that directionally or stabilizing selection were the dominant forces in driving or constraining diversification between these taxa. For example, Weaver and Stringer (22) show that between subspecies of *Pan* cranial differentiation is constrained relative to their genetic divergence. This suggests that cranial divergence between subspecies of *Pan* may be under stabilizing selection or that there is less variation available for genetic drift to act on because of developmental or genetic constraints. Here, subspecies of *Pan* fit the overall pattern across hominoids, that cranial shape variation scales with genetic diversity, but they consistently fall below the regression line, indicating that their cranial diversity is lower than expected given their genetic diversity (Fig. 1). Further analysis, which incorporates morphological variation within and between hominoids in a quantitative framework, could clarify rates of cranial versus genetic change at different taxonomic scales.

**Sexual Dimorphism.** All the male-only regressions show a statistically significant relationship with genetic data for the whole cranium and cranial vault, but not for the face. In females, however, none of the regressions were significant for the whole cranium or the face, and only half were significant for the cranial vault. In total, 18 of 24 regressions were significant in males, and only 4 of 24 were significant in females (SI Appendix, Table S5). This result is interesting in the context of other work that demonstrates that morphology reflects phylogeny more in males than females, especially in highly dimorphic taxa (32). Here, males show a more consistent relationship with genetic data than do females due to the differences in male and female cranial variation within the same taxon. For example, in highly dimorphic taxa (*Pongo* and *Gorilla*), females show less cranial shape variation than males, but this is not the case for less dimorphic taxa such as *Homo sapiens* and *Pan paniscus* (Fig. 2). The stronger relationship between genetic and cranial data in males, coupled with females showing less cranial shape variation than males, suggests that selection may be playing a role in constraining variation in female cranial shape relative to males in certain taxa.

Additionally, *Gorilla* and *Pongo* are not only the most sexually dimorphic apes; they also show some of the highest levels of genetic diversity. Therefore, it is key to note the single-sex results in these taxa particularly. If we look only at the single-sex
whole-cranium analyses (SEV), Pongo and Gorilla are highly variable compared with other taxa, especially in the male-only datasets. These species also show the highest $\pi$ values of all of the apes. If cranial dimorphism were inflating cranial variation in the mixed-sex sample and driving the relationship between genetic and cranial variation in these taxa, we would not expect single-sex data to show the statistically significant relationship with genetic data that they do. Additionally, the mixed-sex gorilla data (SEV) show more cranial shape variation than would be expected, given their genetic diversity; this is due to marked sexual dimorphism in this species, especially in the cranial vault (Fig. 1B).

Hominoid Population History. If population history has impacted both genetic and morphological intraspecific variation in apes, the central question then becomes: What demographic and ecological factors drove this parallel change in molecular and skeletal diversity? Fluctuations in population size and structure through time, which are gleaned from genetic data, together with biogeographical information, provide baseline explanations for our findings in cranial data. All living apes have smaller population sizes than humans, but genetic diversity has been maintained in many species as a vestige of large ancestral population sizes, population substructuring, and older lineage ages (25). The reverse is true in modern humans, in which a recent origin and a population expansion without evidence of a bottleneck (36). Bonobos and common chimpanzees were separated by the formation of the Congo River ~1.5–2 Ma. This barrier inhibited gene flow and restricted bonobos to a small area south of the river. Periodic contractions of forest cover in this region may have forced bonobos into a bottleneck, which is consistent with their low genetic diversity and low cranial shape diversity (37). Nucleotide diversity within Western lowland gorillas (G. gorilla gorilla) is close to estimates within Central chimpanzees and is higher than in other members of the genus Pan (Fig. 3). During the Last Glacial Maximum, African rainforests became fragmented, a process that was reversed postglacially when forest patches expanded and rejoined (38, 39). Gorilla populations may have become discontinuous during this time, creating separate reservoirs of diversity. This population structure would have resulted in the maintenance of genetic (and phenotypic) diversity by providing novel mutations when groups resumed gene flow.

Orangutans show the highest levels of nucleotide diversity among the great apes, with the Sumatran species (Pongo abelii) being more variable than the Bornean species (Pongo pygmaeus) (40, 41). Sumatran orangutans have three deeply structured genetic clusters, indicating long-term separation of these groups (42). Our results support a complex population history for orangutans that is marked by high intraspecific morphological and genetic diversity among hominoids despite small population sizes. In comparison with orangutans, hylobatids are more species rich and geographically continuous. There is evidence of a recent radiation of hylobatid species less than 2 Ma, followed by continued gene flow between certain species (43, 44). These processes have reduced variation between species but may have acted to maintain variation within certain species. Here, nucleotide diversity is higher within Symphalangus syndactylus and Hylobates moloch than within members of the genus Pan. S. syndactylus also shows greater cranial shape diversity than Pan. S. syndactylus still maintains a geographic distribution on Sumatra and a central section of the Malay Peninsula and has a larger census size relative to other species.

Table 1. Cranial shape variation: SEV and PPD in mixed-sex, whole-cranium datasets, pairwise nucleotide diversity ($\pi$), geographic range, and population history for extant hominoids

<table>
<thead>
<tr>
<th>Hominoid taxon</th>
<th>N</th>
<th>n</th>
<th>SEV</th>
<th>PPD</th>
<th>$\pi$ (%)</th>
<th>Census size</th>
<th>Geographic distribution</th>
<th>Population history inference</th>
</tr>
</thead>
<tbody>
<tr>
<td>P. abelii</td>
<td>8</td>
<td>10</td>
<td>0.0200</td>
<td>0.0166</td>
<td>0.42</td>
<td>~7,300 Sumatra</td>
<td>Long-term fragmented range, recent range reduction (25, 40)</td>
<td></td>
</tr>
<tr>
<td>P. pygmaeus</td>
<td>20</td>
<td>15</td>
<td>0.0142</td>
<td>0.0091</td>
<td>0.35</td>
<td>~50,000 Borneo</td>
<td>Long-term fragmented range, recent range reduction (25, 40, 41)</td>
<td></td>
</tr>
<tr>
<td>G. gorilla gorilla</td>
<td>29</td>
<td>41</td>
<td>0.0141</td>
<td>0.0102</td>
<td>0.19</td>
<td>~95,000 Central Africa</td>
<td>Constant population size, almost continuous distribution, recent range reduction (25, 38, 39)</td>
<td></td>
</tr>
<tr>
<td>S. syndactylus</td>
<td>17</td>
<td>22</td>
<td>0.0090</td>
<td>0.0103</td>
<td>0.21</td>
<td>~190,000 Sumatra, Malay Peninsula</td>
<td>Long-term widespread, shrinking but continuous populations (43)</td>
<td></td>
</tr>
<tr>
<td>H. sapiens</td>
<td>18</td>
<td>20</td>
<td>0.0065</td>
<td>0.0098</td>
<td>0.10</td>
<td>~7 billion Cosmopolitan</td>
<td>Bottleneck followed by recent massive range expansion (33, 34)</td>
<td></td>
</tr>
<tr>
<td>P. troglodytes schweinfurthii</td>
<td>3</td>
<td>8</td>
<td>0.0057</td>
<td>0.0062</td>
<td>0.15</td>
<td>~89,000 Congo River to W. Uganda, Rwanda, W. Tanzania</td>
<td>Bottleneck, expansion, and recent range reduction (25, 35, 36)</td>
<td></td>
</tr>
<tr>
<td>P. troglodytes troglodytes</td>
<td>50</td>
<td>26</td>
<td>0.0056</td>
<td>0.0064</td>
<td>0.18</td>
<td>~90,000 Central Africa Sanaga River to Congo River</td>
<td>Constant population size, recent range reduction (25, 35, 36)</td>
<td></td>
</tr>
<tr>
<td>H. moloch</td>
<td>6</td>
<td>8</td>
<td>0.0050</td>
<td>0.0080</td>
<td>0.21</td>
<td>~2,500 Java</td>
<td>Constant island population, recent range reduction (43)</td>
<td></td>
</tr>
<tr>
<td>P. troglodytes verus</td>
<td>12</td>
<td>10</td>
<td>0.0044</td>
<td>0.0064</td>
<td>0.12</td>
<td>~55,000 West Africa, Senegal to Nigeria</td>
<td>Bottleneck, expansion and recent range reduction (25, 35, 36)</td>
<td></td>
</tr>
<tr>
<td>P. paniscus</td>
<td>21</td>
<td>17</td>
<td>0.0037</td>
<td>0.0057</td>
<td>0.09</td>
<td>~50,000 Central Africa, South of Congo River</td>
<td>Bottleneck and continuous restricted range (25, 36)</td>
<td></td>
</tr>
<tr>
<td>H. klossii</td>
<td>10</td>
<td>11</td>
<td>0.0035</td>
<td>0.0062</td>
<td>0.08</td>
<td>~25,000 Mentawai Islands</td>
<td>Long-isolated island populations (44)</td>
<td></td>
</tr>
<tr>
<td>H. pileatus</td>
<td>3</td>
<td>3</td>
<td>0.0031</td>
<td>0.0052</td>
<td>0.05</td>
<td>~40,000 SE Thailand, Cambodia, SW Laos</td>
<td>Restricted range (44)</td>
<td></td>
</tr>
</tbody>
</table>

Taxa are ordered by SEV. N = number of cranial samples in females (f) and males (m); n = number of genetic samples. All population census size estimates are from the International Union for the Conservation of Nature Red List.
to other hylobatids. Genetic variation in *H. moloch* is higher than might be expected given its critically low census size of 2,500 individuals, although its current distribution in forest fragments in Western and Central Java is not representative of the historical range of the species. *Hylobates klossii* lives exclusively on the Mentawai islands and has the smallest geographic range of the hylobatids in this study; its genetic and cranial variation are among the lowest presented here. *Hylobates pileatus*, in southeastern Thailand, Cambodia, and southwestern Laos, shows the lowest genetic diversity and cranial variation of all hominoid species here.

**Implications for Fossil Hominins.** The finding that cranial morphology preserves signals of past population expansion and bottleneck history in hominoids can also guide our understanding of variation in fossil hominin crania. For example, extant hominoids often serve as modern analogs of variation which inform inferences of intraspecific variation in extinct groups. Our results suggest that the population history of extant apes should be considered when choosing analogs of intraspecific variation for fossil hominins. Accordingly, modern *H. sapiens* may be a suboptimal analog for variation in fossil hominins, despite their close phylogenetic relatedness. Modern humans have a unique population history featuring at least one severe bottleneck followed by rapid expansion and repeated founder events (33, 34). If these features of human population history have a major impact on cranial variation, then modern human variation provides a limited model of cranial variation for extinct hominins. For example, for fossil species such as *Homo erectus*—a cosmopolitan hominin species with a temporally longer lineage than modern humans—we might expect more variation than we see in humans today if this species did not experience an equivalent bottleneck and rapid expansion. Additionally, for fossil hominin populations from the same species and time horizon (e.g., *Homo naledi* from the Dinaledi chamber, South Africa, *H. erectus* from Dmanisi, Georgia, and the fossil hominins *Simi de los Huesos*, Spain), the amount of cranial variation in adults may broadly reflect population genetic structure (45–47). Overall, results here suggest that intraspecific morphological variation in hominin crania can be viewed through a population genetics framework—with consideration of how multiple different population models could explain the observed levels of variation.

**Future Research.** The results we report here open possibilities for future analyses within the primates, including those at different geographic sampling scales and with the inclusion of additional skeletal elements such as postcrania and dentition. This work also may have implications for developmental analyses of the cranium. For example, if neutral genetic diversity explains a portion of intraspecific cranial shape variation, then studies assessing variation in different developmental modules of the cranium could account for different population histories when comparing taxa.

More broadly, this work provides preliminary empirical support that neutral population processes have impacted extant hominid cranial morphology and evolution, and that this pattern may be relevant to other nonhuman groups.

**Materials and Methods**

**Morphological Data.** A total of 396 adult crania from 12 taxa (species and subspecies) were included in this analysis: *H. sapiens*, *P. paniscus*, *P. troglodytes* *troglobytes* *troglodytes* *verus*, *P. troglodytes* *schweinfurthii*, *G. gorilla gorilla*, *P. pygmaeus*, *P. abelii*, *S. syndactylus*, *H. moloch*, *H. klossii*, and *H. pileatus* (Dataset S1).

Morphological data consisted of 34 homologous cranial landmarks (each consisting of a set of x, y, z coordinates) capturing cranial shape differences (SI Appendix, Table S2) (48, 49). All landmark data were subjected to a generalized Procrustes Analysis (GPA) to project them into a common shape space. The GPA superimposes the centroids of each individual’s landmark configuration, then scales all landmark configurations to unit size and rotates all specimens around that centroid. Differences in translation, size, and orientation are eliminated during this step, so that only differences in shape remain among specimens. Landmark data were divided into three analytic units: (i) whole cranium, (ii) cranial vault, and (iii) face, consisting of 34, 12, and 22 landmarks, respectively. Three separate GPs were performed for each of the three landmark units.

We applied two types of morphological analyses: average PPD and SEV. The average PPD for each taxon was calculated as the mean squared difference between all pairs of individual landmarks belonging to the same taxon and optimally aligned in Kendall’s shape space (SI Appendix, Fig. S2 and Table S3) (50, 51). The SEV for each taxon was calculated from the variance–covariance matrix of the superimposed coordinates calculated separately for each taxon. This is a symmetric square matrix in which the diagonal elements are the variances of the individual landmark coordinates and the off-diagonal elements are the covariances among coordinates subsequent to the GPA superimposition described above. This value equals the cumulative variance that is captured across all landmarks (Fig. 2 and SI Appendix, Table S3) and is also equivalent to the group’s Procrustes variance, which measures the mean squared Procrustes distance of each specimen to the average shape (52). Quantitative genetic theory predicts a linear relationship between SEV and neutrally evolving population genetic data; however, this is not the case for PPD (53). We chose to include both measures of morphological variation here to provide evidence that the patterns of cranial shape variation within taxa are similarly robust when different methods are used.

Tests were performed to determine how sample size influenced the mean PPD within each group. The largest sample size was 76 individuals for *P. troglodytes* *troglobytes*. Two individuals from the same species were randomly sampled, and the PPD between them was calculated. The resampling procedure was repeated 10,000 times with replacement for each taxon, and the average pairwise distance was recorded for random subsets of 75, 50, 20, 10, and 5 individuals. For *P. troglodytes* *schweinfurthii*, with 10 individuals, the average pairwise distance was recorded for random subsets of 75, 50, 20, 10, and 5 individuals. For *H. pileatus*, the resampling procedure was repeated 10 separate times and then was averaged. Across the different sample sizes, the average PPD values were 0.00641 for all three of the largest sample sizes (*n* = 75, *n* = 50, and *n* = 20), 0.00645 for *n* = 10, and 0.00642 for *n* = 5. Varying the sample sizes did not yield appreciably different average PPD values, thus confirming that sample sizes used here were adequate for capturing intraspecific cranial shape diversity that reflects a larger taxon-wide trend. This is especially relevant for estimation of PPD for single-sex, within-taxa samples, which were represented by the smallest number of individuals.

Sample size tests for SEV were also performed. Overall, the smallest mixed-sex cranial dataset was six individuals (*H. pileatus*). All taxa were sampled down to six individuals (three females, three males) for the whole-cranium dataset, and SEV was calculated (SI Appendix, Table S6). Even with sample sizes of six individuals, we see similar trends in the magnitude of variation within taxa. *Pongo, Gorilla,* and *Symphalangus* remain the most variable taxa, and *P. paniscus*, *H. klossii*, and *H. pileatus* are the least variable. We see the largest difference in SEV values between the sample size of six and the larger sample size of 76 in *P. troglodytes* *troglobytes*. This taxon represents the largest sample size in our analysis.

**Genetic Data.** Genetic diversity was summarized by pairwise nucleotide diversity (π) and number of segregating sites (w). Sequence data from 11 homologous noncoding autosomal loci across all 12 taxa were downloaded from GenBank (SI Appendix, Table S1). The genetic estimator of π is calculated by randomly sampling two individuals within...
each taxon, with replacement, and then taking the average nucleotide difference between pairs. This was performed for each locus separately and then averaged across all loci to arrive at a single $\pi$ value for each taxon (Fig. 3 and SI Appendix, Table S4). Nucleotide diversity estimates ($\pi$, $\theta$) from the autosomal loci chosen here reflect neutral evolutionary processes and serve as a proxy for genome-wide impacts of population history. Effective population size was calculated using the standard equations: $N_e = \pi(4\mu)$ and $N_e = \theta(4\mu)$. The mutation rate ($\mu$) was derived from the average pairwise differences between two species (for all loci) divided by the number of generations between the estimated divergence of the two species (SI Appendix, Table S4) (55). The following generation times were used in the $N_e$ calculations: H. sapiens, 29 y; Pan, 25 y; Gorilla, 19.3 y; Pongo, 26.7 y; and Hylobates and Symphalangus, 20 y. The following divergence times were used: Homo–Pan, 7 Ma; Pan–Gorilla, 13.5 Ma; Gorilla–Pongo, 16 Ma; Pongo–Symphalangus, 20 Ma; and Pongo–Hylobates, 20 Ma (S5).

**Regression Analyses.** Finally, 72 ordinary least squares regressions were performed to determine the strength of the relationship between genetic and morphological diversity (SI Appendix, Table S5). Ordinary least squares (OLS) regressions were performed with raw $\pi$ and $\theta$ values as well as $N_e = \pi(4\mu)$ and $N_e = \theta(4\mu)$. Each of these four genetic values was separately regressed against the corresponding SEV and PPD values. This was done for each landmark set and for both sexes together and separately. Additionally, phylogenetic generalized least squares (PGLS) regressions were performed to determine if the results from the OLS regressions were the result of close evolutionary relationships between taxa. A tree file of all species used in this analysis was generated from the website [https://10ktrees.nunn-lab.org/index.html](https://10ktrees.nunn-lab.org/index.html). This file was loaded into R, and PGLS regressions were performed using the packages (caper) and (ape). All results showed that the relationship between genetic and cranial variation was not a result of close evolutionary relationships between the taxa sampled here, with lambda values at or close to 0 (indicating that the data are not compatible with a Brownian motion model of evolution).

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