Skull diversity and evolution in miniaturized amphibians, genus Brachycephalus (Anura: Brachycephalidae)

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Abstract
Miniaturized amphibians of the genus Brachycephalus are phenotypically diverse. The species of Brachycephalus have bufoniform or leptodactytilform Baupläne and any of three skeletal states: nonhyperossified, hyperossified without dorsal shield, and hyperossified with dorsal shield. We integrate high-resolution microcomputed tomography, geometric morphometrics, and an estimate of molecular phylogenetic relationships to investigate skull diversity in shape and size-shape space in selected species of Brachycephalus. Skull diversity amongst species of Brachycephalus can be partitioned into shape and size-shape space according to the four conditions of skeletal states-Baupläne, namely, nonhyperossified leptodactytilform, nonhyperossified bufoniform, hyperossified bufoniform without dorsal shield, and hyperossified bufoniform with dorsal shield. Skull diversity in shape and size-shape space in nonhyperossified leptodactytilform species of Brachycephalus is markedly larger, when compared to skull diversity in species of the three other conditions of skeletal states-Baupläne. Variation in skull shape scales with size across Brachycephalus and, therefore, can be explained by allometry. Skull diversity, Baupläne, and skeletal states covary to a large extent with monophyletic lineages of Brachycephalus, as revealed by a mitochondrial DNA species tree. Nonhyperossified bufoniform species and hyperossified bufoniform species with or without dorsal shield are monophyletic lineages, as inferred from a mitochondrial DNA species tree. Nonhyperossified leptodactyliform species of Brachycephalus do not share, however, a most recent common ancestor. The nonhyperossified leptodactyliform species of Brachycephalus, due to their marked skull diversity and lack of monophyly, emerge as evolutionarily complex. Therefore, further sampling of the nonhyperossified leptodactyliform condition of skeletal states-Baupläne will be necessary to further understand the evolutionary history of Brachycephalus.

KEYWORDS
Allometry, Brazilian Atlantic Forest, geometric morphometrics, hyperossification, mitochondrial DNA species tree, shape space, size-shape space, skull
Amphibians of the genus *Brachycephalus* are amongst the smallest vertebrates in the world. Currently, 36 species are described for the genus *Brachycephalus*, which occurs in the Atlantic Forest from Northeast to Southern Brazil (Condez, Haddad, & Zamudio, 2020; Frost, 2020). Externally, the species of *Brachycephalus* have one of two Baupläne (Handigran & Wassersug, 2007), that is, a common, basic organizational plan: bufoniform (toad-like) or leptodactyliform (frog-like, Figure 1). The bufoniform/leptodactyliform Baupläne are defined comparatively as follows (Condez et al., 2014, 2016): body robust/slender, pectoral girdle robust/slender, head as wide as long/wider than long, snout short/long. The bufoniform species of *Brachycephalus* amount to 32 taxa, with several additional species still unnamed (Condez et al., 2020). Four species of leptodactyliform *Brachycephalus* have been described and several others remain undescribed (Condez et al., 2020; Napoli, Caramaschi, Cruz, & Dias, 2011). Bufoniform species vary markedly in body coloration, whereas leptodactyliform species are brownish in color (Figure 1). *Brachycephalus* is at the lowest limit of vertebrate size, ranging from 8.6 to 19.0 mm in body size. The smallest species of *Brachycephalus* are near in size to the smallest known vertebrate, the frog *Paedophryne amauensis* from New Guinea, which attains an average body size of 7.7 mm (Rittmeyer, Allison, Gründler, Thompson, & Austin, 2012).

The dramatic reduction in body size in *Brachycephalus* is the outcome of miniaturization, an evolutionary process by which extremely small body size in a lineage arises from a larger ancestor (Hanken & Wake, 1993). Whereas there is not a critical size that defines miniaturization on theoretical or biological criteria (Hanken & Wake, 1993), it is widely accepted that amphibians <20 mm in body size are miniaturized (Clarke, 1996). *Brachycephalus* is therefore miniaturized given this threshold in body size. An evolutionary novelty, a hyperossified dorsal shield, did arise in some bufoniform species of *Brachycephalus* (Clemente-Carvalho et al., 2009; Hanken, 1993). Hyperossification, an increased mineralization and excessive ossification, is a prominent feature of the skeleton of some bufoniform species (Clemente-Carvalho et al., 2009; Hanken, 1993; Trueb, 1973; Trueb & Alberch, 1985). Three skeletal states evolved in *Brachycephalus*, namely, nonhyperossified, hyperossified without dorsal shield, and hyperossified with dorsal shield (Clemente-Carvalho et al., 2009). Nonhyperossified species lack bone sculpturing in the skull, spinal processes of sacral, and presacral vertebrae (Clemente-Carvalho et al., 2009). Hyperossified species have bone sculpturing, which appears macroscopically as ridges and crests inducing a reticulated or pitted pattern in the skull, spinal processes of sacral, and presacral vertebrae (Clemente-Carvalho et al., 2009). The dorsal shield is also sculptured (Clemente-Carvalho et al., 2009).

Morphological diversity such as that observed in complex structures like the skull can be measured qualitatively in terms of variation in phenotypic states and, quantitatively, in terms of the association between size and shape within the context of allometry, which explains how

**Figure 1** Images of *Brachycephalus*. From left to right (top to bottom). Leptodactyliform species: *B. hermogenesi*. Bufoniform species: *B. mariaeterezae*, *B. albolineatus*, *B. garbeanus*, *B. pitanga*, and *B. toby*. Images not to scale.
changes in size extrapolate to changes in shape (Klingenberg, 2016). A fundamental task is therefore to determine the extent to which morphological diversity is associated with molecular phylogenetic relationships. Here, we apply these concepts to investigate skull diversity in selected species of *Brachycephalus*. We chose the skull because this structure is a vertebrate novelty that houses the brain and specialized sensory organs (Fish, 2017; Yang & Ornitz, 2019). Specifically, our objectives were the following. First, we use virtual surfaces derived from high-resolution microcomputed tomography to characterize the three skeletal states and their distribution across the species of *Brachycephalus* sampled here. Second, we use threedimensional geometric morphometrics (3DGM; Cardini, 2020; Mitteroecker, 2020) to characterize skull diversity in *Brachycephalus* in ordinated shape space (Mitteroecker, 2020). From a geometric perspective, the scale of description of shape is determined by the density of points, registered as Cartesian coordinates, sampled from individual phenotypes. We used the virtual surfaces to manually register points, defined as landmarks and curve and surface semilandmarks, which yield a limited number of coordinates (Gao, Yapuncich, Daubechies, Mukherjee, & Boyer, 2018). We also used a fully automated method of registration of coordinates, which yields tens of thousands of coordinates (Pomidor, Makedonska, & Slice, 2016). Third, we further quantify the diversity in the skull of *Brachycephalus* in ordinated size-shape space within the framework of allometry (Mitteroecker, Gunz, Fernhard, Schaefer, & Bookstein, 2004). Fourth, we use mitochondrial DNA sequences to compute a species tree to estimate ancestor–descendant relationships in *Brachycephalus* under the formalism of the multi-species coalescent model (Rannala, Edwards, Leaché, & Yang, 2020).

2 | MATERIAL AND METHODS

2.1 | Taxon sampling

The species of *Brachycephalus* have one of two Baupläne, bufoniform or leptodactyliform. Each species of *Brachycephalus* examined here was classified as either bufoniform or leptodactyliform based on definitions of these Baupläne and/or information in the original descriptions. Each species was also classified to a skeletal state, namely, nonhyperossified, hyperossified without dorsal shield, or hyperossified with dorsal shield, based on the reconstructed tomographic surfaces. Given the Baupläne and the skeletal states there are four conditions to which a species of *Brachycephalus* can be assigned to, namely, nonhyperossified leptodactyliform, nonhyperossified bufoniform, hyperossified bufoniform without dorsal shield, and hyperossified bufoniform with dorsal shield. Taxon sampling of *Brachycephalus* aimed at morphological and molecular analyses should include species representative of each of these four conditions, and we sampled 18 species spanning the four conditions of skeletal states-Baupläne. For the morphological analyses we sampled 55 alcohol-preserved specimens of 18 species of *Brachycephalus* of each three skeletal states. For the molecular analyses we had access to tissue samples of the following species of the four skeletal states-Baupläne conditions. Nonhyperossified leptodactyliform: *B. didactylus*, *B. hermogenesi*, and *B. pulex*; nonhyperossified bufoniform: *B. brunneus*, *B. ferruginus*, *B. izecksohni*, *B. pernix*, and *B. pombali*; hyperossified bufoniform without dorsal shield: *B. alipi**, *B. crispus*, *B. guarani*, *B. nodoterga*, *B. pitanga*, *B. vertebralis*, and hyperossified bufoniform with dorsal shield: *B. toby*, and *B. ephippium*, *B. garbeaus*, and *B. margaritatus*. Information on samples sizes, localities, and geographic coordinates are given in Table 1. Taxonomic identification of species was based on morphological information provided in the original descriptions. The specimens analyzed here are deposited in the collections of the Museu Nacional, Rio de Janeiro; Universidade Estadual de Santa Cruz, Bahia; Coleção Célio Haddad, Universidade Estadual Paulista, Rio Claro, São Paulo; and Museu de Zoologia da Universidade Estadual de Campinas “Adão José Cardoso”, São Paulo, Brazil.

2.2 | Virtual reconstruction of skeletal diversity from 3D microcomputed tomography

High resolution three-dimensional (3D) microcomputed tomography images of museum preserved skeletons of *Brachycephalus* were made using the Phoenix V	omex300 M GE tomography system at the Laboratório de Instrumentação Nuclear at COPPE, Universidade Federal do Rio de Janeiro, Rio de Janeiro, Brazil. The parameters for image acquisition were as follows: 55 kV voltage and 250 μA current for each frame. We made an average of five frames (skipping 2), with 250 ms exposure time and a total of 1,200 projections with pixel size within the range of 13 μm to 18 μm per scan. Three-dimensional reconstructions of individual *Brachycephalus* skeletons were made using the software Phoenix Datos/X v. 2.2 (GE). The threshold-based and manual segmentation was performed with the AVIZO software (AVIZO Fire 9.1). Images were saved as polygon (.ply) files. According to Pomidor et al. (2016), to generate surfaces from tomographic images individual specimens must be the most complete, the most representative, and the most morphologically atypical in the sample. We made tomographic images of several specimens per
TABLE 1  Species, sample sizes, sampling localities, and geographic coordinates of the specimens of *Brachycephalus* for which mitochondrial genes 12S RNA, 16S RNA, COI, and Cyt b were sequenced

| Species            | Sample size | Locality                                      | Coordinates               |
|--------------------|-------------|-----------------------------------------------|---------------------------|
| *B. alipoi*        | 5           | Vargem Alu, Espírito Santo, Brazil            | 20° 28' S, 41° 00' W     |
| *B. brunnneus*     | 4           | Campina Grande do Sul, Paraná, Brazil         | 25° 23' S, 48° 83' W     |
| *B. crispus*       | 5           | Cunha, São Paulo, Brazil                      | 23° 15' S, 45° 01' W     |
| *B. didactylus*    | 6           | Ilha Grande, Rio de Janeiro, Brazil           | 23° 05' S, 44° 16' W     |
| *B. ephippium*     | 9           | São Francisco Xavier, Rio de Janeiro, Brazil  | 23° 10' S, 45° 53' W     |
| *B. ferruginus*    | 3           | Morretes, Paraná, Brazil                      | 25° 43' S, 48° 92' W     |
| *B. garbeanus*     | 12          | Macac de Cima, Rio de Janeiro, Brazil         | 22° 28' S, 42° 12' W     |
| *B. guarani*       | 4           | Ubatuba, São Paulo, Brazil                    | 23° 20' S, 45° 01' W     |
| *B. hermogenesi*   | 5           | Piedade, São Paulo, Brazil                    | 23° 42' S, 47° 25' W     |
| *B. izecksohni*    | 4           | Paranaguá, Paraná, Brazil                     | 25° 78' S, 49° 38' W     |
| *B. margaritatus*  | 4           | Engenheiro Paulo de Frontin, Rio de Janeiro, Brazil | 22° 30' S, 43° 35' W |
| *B. nodoterga*     | 8           | Boracéia, São Paulo, Brazil                   | 23° 31' S, 45° 50' W     |
| *B. pernix*        | 4           | Quatro Barras, Paraná, Brazil                 | 25° 38' S, 49° 08' W     |
| *B. pitanga*       | 2           | São Luís do Paraitinga, São Paulo, Brazil     | 23° 20' S, 45° 08' W     |
| *B. pombali*       | 4           | Guaratuba, Paraná, Brazil                     | 25° 65' S, 48° 85' W     |
| *B. pulex*         | 2           | Camacan, Bahia, Brazil                        | 15° 23' S, 39° 33' W     |
| *B. toby*          | 5           | Ubatuba, São Paulo, Brazil                    | 23° 27' S, 45° 11' W     |
| *B. vertebralis*   | 4           | Parati, Rio de Janeiro, Brazil                | 23° 09' S, 23° 13' W     |

species, digitally reconstructed the skull, and retained only individuals that met the three criteria defined by Pomidor et al. (2016). At the end of this process, we were able to select one specimen per species of *Brachycephalus* for which we computed clean surfaces needed for the automated procedures employed downstream in the geometric morphometric analyses (Pomidor et al., 2016).

2.3 Three-dimensional geometric morphometric analysis of skull diversity in shape space

We used three methods to analyze skull diversity between the species of *Brachycephalus* in ordinated shape space. In the first method, we used a landmark-free method, the Generalized Procrustes Surface Analysis (GPSA; Pomidor et al., 2016), with the 3D surfaces reconstructed from high-resolution computed tomography. Generalized Procrustes Surface Analysis optimally superimposes multiple surfaces associating each point in one surface with its nearest neighbor on another surface (Pomidor et al., 2016). Multiple surfaces are superimposed to a designated mean (prototype) surface, which should be the most representative, least atypical species in the sample (Pomidor et al., 2016). We chose *B. ferruginus* as the designated mean (prototype) surface because the specimen was extremely well preserved, most representative in the sense that the majority of species of *Brachycephalus* are not hyperossified, and had no atypical morphometric features. The optimal superimposition for multiple surfaces is computed recursively by minimizing the Procrustes surface metric (PSM) between surfaces. PSM is a distance analogous to the Procrustes distance derived from Generalized Procrustes Analysis (Pomidor et al., 2016), which is used to quantify shape differences between species. Diversity in skull shape between the species of *Brachycephalus* was visualized as a three-dimensional heat map. A heat map associates colors with the magnitude of variance measured at every point on the mean (prototype) surface with respect to the sample (Pomidor et al., 2016). The heat map is an effective way to graphically display shape variation in the sample (Pomidor et al., 2016). The color values are computed from the covariance matrix of the set of nearest neighbor points for each point on the mean (prototype) surface (Pomidor et al., 2016). Variance values are color coded, with blue and red indicating low and high values, respectively (Pomidor et al., 2016). All computations were carried out with the GPSA software package (Pomidor et al., 2016).

In the second method, we used Generalized Procrustes Analysis (GPA) with landmarks and curve and surface semilandmarks (hereafter, reference points [Reyment, 2010]),
which were manually registered on the three-dimensional images of skulls of Brachycephalus generated with AVIZO. Reference points are depicted in Figure 2 and described in Table 2. The coordinates for the reference points for the 18 species of Brachycephalus sampled here are available in the Supplementary Information S1. To establish geometric correspondence between semilandmarks across the samples, the semilandmarks were allowed to slide along tangents in order to minimize the thin-plate spline bending energy between each individual skull and the sample average (Bookstein, 1997; Gunz & Mitteroecker, 2013; Gunz, Mitteroecker, & Bookstein, 2005). Generalized Procrustes Analysis superimposes the original configurations of the reference points onto a single, global consensus configuration by carrying out three operations (Mitteroecker, 2020). First, differences in the position of the original configurations of the reference points are normalized by translation to a common centroid, computed as the average $x, y,$ and $z$ coordinates of all reference points of a configuration. Second, differences in size in the original configurations of reference points are normalized by scaling to an identical size, the unitary centroid size, defined as the square root of the summed squared distances between the reference points and their centroid. Third, the normalized and scaled reference points are optimally superimposed onto a mean (consensus) configuration so as to minimize the Procrustes distance, computed as the summed squared distances between corresponding reference points over all configurations. The size of each configuration of reference points was calculated as the centroid size, defined as the square root of the sum of squared distances of a set of reference points from their centroid. The Procrustes distances generated by GPA are global distances, which were used to compute differences in shape between the species of Brachycephalus. The sliding of semilandmarks was performed using the slider3d function from the R package Morpho version 2.8 (Schlager, 2017), whereas the GPA was computed using the procSym function in the same package.

In the third method, we relaxed the restriction of superimposition based on the Procrustes distances to a single, global consensus by using multiple, local consensuses (Von Zuben, Reis, & Perez, 2020). By using multiple, local consensuses the algorithm performs the superimposition taking into account the pattern of diversification in the skulls of Brachycephalus. The algorithm starts by computing the matrix of pairwise Procrustes distances among the reference points of the 18 species of Brachycephalus using ordinary Procrustes analyses (Dryden & Mardia, 2016). Based on these distances, we computed a tree using the neighbor-joining (NJ) algorithm (Saitou & Nei, 1987). The NJ tree has $n = 18$ leaf nodes which correspond to the 18 species of Brachycephalus and $n - 2 = 16$ internal nodes. We estimated the hypothetical landmarks for the unobserved, internal nodes with the squared-change parsimony (SCP) algorithm (McArdle & Rodrigo, 1994), which minimizes the sum of squared internode differences, for all the coordinates of landmarks and semilandmarks across the entire tree. The NJ and SCP algorithms are
computationally efficient and did not pose a significant impact on the whole computational burden. We still needed to perform \( n \) local Procrustes analyses, given the NJ tree composed of \( n \) leaf nodes. Therefore, given a reference node \( i \) for \( i = 1, \ldots, n \), we simply extracted from the NJ tree the local star tree having the \( i \)-th leaf node as one of its leaf nodes. This local star tree will always have three leaf nodes, one or two which will be internal nodes of the full tree, allowing for a very fast local Procrustes analysis. Although all the leaf nodes in the current local star tree have their corresponding reference points superimposed, only the reference node \( i \) will have the landmarks updated. Therefore, the proposal is iterative. Given the original skull reference points to be superimposed, we (a) obtained the distance matrix between them using the pairwise Procrustes distance; (b) applied the NJ algorithm to compute the binary tree for the current configuration of landmarks for the leaf nodes; (c) ran the SCP algorithm to compute the hypothetical landmarks and semilandmarks for the internal nodes of the tree; and finally (d) explored the hierarchical structure of the NJ tree to perform local Procrustes analysis for each one of the \( n \) leaf nodes. This sequence of steps is then repeated until convergence, measured by the amount of change in the landmarks and semilandmarks at the leaf nodes given the previous and the current iteration. The output of our algorithm includes the locally optimized landmarks and semilandmarks for the skull of the 18 species.

| Number and definition of landmarks | Position |
|-----------------------------------|----------|
| **Dorsal view of the skull**      |          |
| 1. Anteriormost point of nasal bone | Middle line |
| 2/9. Anteriormost projection of nasal bones | Bilateral |
| 3/10. Latero-anterior projection of nasal bones | Bilateral |
| 4/11. Lateral point at the border between nasal and frontoparietal bones | Bilateral |
| 5/12. Lateral point at the border between frontoparietal bones and anteriormost part of the postorbital crest | Bilateral |
| 6/13. Posteriormost point of squamosals | Bilateral |
| 7/14. Posteriormost point of the parotic plates | Bilateral |
| 8/15. Midlateral point of the occipital condyles | Bilateral |
| 16. Medial point of the os basale | Middle line |
| **Ventral view of the skull**     |          |
| 17. Medial point of the os basale | Middle line |
| 18/22. Posterolateral point at the prootic-exoccipital | Bilateral |
| 19/23. Mediolateral point at the prootic-exoccipital | Bilateral |
| 20/24. Anterolateral point at the prootic-exoccipital | Bilateral |
| 21/25. Latero-medial projection of nasal bones | Bilateral |
| 26. Anteriormost point of vomer | Middle line |
| **Mandible**                      |          |
| 27. Midpoint between the premaxillae | Middle line |
| 28/31. Point at tip of premaxillae | Bilateral |
| 29/32. Anteriormost point of maxillae | Bilateral |
| 30/33. Point at the tip of process of the maxillae | Bilateral |
| **Surface semilandmarks**         |          |
| Semilandmarks (\( n = 33 \)) equally spaced on the surface of frontoparietal, exoccipital, and squamosal bones | Bilateral |
| **Contour semilandmarks**         |          |
| Semilandmarks roughly equidistant at the margin of frontoparietal bones and on the squamosal, and prootic bones (\( n = 14 \)), and on the medial line of the parasphenoid and premaxilla bones (\( n = 6 \)) | Bilateral for frontoparietal, squamosal, and prootic bones |
the local Procrustes distances (l-Procrustes distances), and the corresponding NJ tree computed from the distances. Our method has been implemented in MATLAB (Von Zuben et al., 2020) and the convergence was always achieved with less than 20 iterations, considering a wide range of problems with a distinct number of leaf nodes (from tens to hundreds).

The PSM distances, the global Procrustes distances (g-Procrustes distances), and the local Procrustes distances (l-Procrustes distances) were then subjected to nonmetric multidimensional analysis (nmMDS; Mead, 1992). nmMDS optimally projects distances and shape coordinates onto a lower dimensional space for visualization of the ordinated shape space and was computed using PAST 4.0 (Hammer, Harper, & Ryan, 2001).

2.4 Three-dimensional geometric morphometric analysis of skull diversity in size-shape space

The covariance between shape and size in the skull of *Brachycephalus* was quantified using the landmarks, curve semilandmarks, and surface semilandmarks under the concept of allometry within the size-shape space formalism of geometric morphometrics (Dryden & Mardia, 2016; Mitteroecker et al., 2004). Under this formalism, interspecific allometric trajectories are calculated taking into account intraspecific allometries by computing a common allometric component (CAC), defined as $\mathbf{a} = (\mathbf{X}'s)/(\mathbf{s}'s)$. Here, $\mathbf{X}$ is the $n \times n$ matrix of shape coordinates derived from GPA and $\mathbf{s}$ is the $n \times 1$ vector of centroid sizes on a logarithmic scale. The vector $\mathbf{a}$ is normalized as $\mathbf{a}' = \mathbf{a} / \sqrt{\mathbf{a}'a}$. Visualization of allometric patterns is achieved by plotting scores of CAC against centroid size on a logarithmic scale (Mitteroecker et al., 2004). Residual shape components (RSC) are calculated by projecting out the common allometric component, computed as $\mathbf{W} = (\mathbf{I} - \mathbf{a}'(\mathbf{a}')'$. Residual shape components are the nonallometric shape components (Mitteroecker et al., 2004). Ordination of species in size-shape space was examined by plotting scores of the first residual component (RSC–1) against CAC (Mitteroecker et al., 2004). Diversity in skull morphology between species of *Brachycephalus* in size-shape space was visualized as regionalized shape deformations representing spatial patterning (Bookstein, 1989; Mitteroecker et al., 2004). Deformation is computed with the thin-plate splines that interpolates surfaces over the reference points, minimizing the bending energy in analogy with the bending of an infinitely thin metal plate in continuum mechanics (Batra, 2006). All computations were performed with the CAC function in the R package Morpho (Schlager, 2017).

2.5 Species tree estimation

Species tree estimation was performed using four mitochondrial genes, rRNA 12S, rRNA 16S, cytochrome c oxidase subunit I [COI], and cytochrome b [Cyt b]. We used 90 specimens of the 18 species of *Brachycephalus* sampled here (Table 1). Genomic DNA was extracted from liver (or muscle) tissue preserved in 100% ethanol (Clemente-Carvalho et al., 2015). Tissue samples were digested with proteinase K followed by a standard three-step phenol-chloroform extraction procedure (Green & Sambrook, 2013). Amplification via polymerase chain reaction (PCR) of the rRNA 12S, COI, and Cyt b genes was based on primers developed by Goebel, Donnelly, and ATZ (1999). The rRNA 16S gene was amplified after Darst and Cannatella (2004). PCR amplification products were visualized on 1.0% agarose gels and purified using a QIAquick PCR Purification Kit (QIAGEN Inc., Venlo, The Netherlands). Purified PCR products were outsourced to Macrogen Inc. (Seoul, South Korea) for sequencing using the BigDye Terminator Kit and run on an ABI 3730xl DNA analyzer [Applied Biosystems, Inc., Grand Island, NY, USA]. Sequences were obtained in both directions with the same primers used for polymerase chain reaction amplification and subjected to BLAST searches (Altschul et al., 1997) in GenBank to determine that the target sequences had been amplified. All sequences were deposited in GenBank. Accession numbers are given in Supplementary Information S2. Sequence traces were analyzed using the phred program (Ewing, Hillier, Wendl, & Green, 1998). We obtained a total of 4,826 base pairs (bp), of which 939, 1,533, 1,369, 985 bp were from the 12S rRNA, 16S rRNA, COI, and Cyt b genes, respectively. The authenticity of the COI and Cyt b genes was confirmed by amino acid translation. Sequences were aligned using MAFFT (Katoh & Standley, 2013). We estimated a species tree for the species of *Brachycephalus* sampled here using the multi-species coalescent model implemented in BEAST 2.3 (Bouckaert et al., 2014). Sequence data containing the four mitochondrial genes were imported into BEAUti 2. Clock models and trees were linked across genes to ensure that mitochondrial genes shared the same evolutionary history. Site models were unlinked to allow for each gene to have its model of nucleotide substitution. Best-fitting DNA substitution models were selected according to the Akaike Information Criterion (AIC) in jModeltest 2.1.4 (Darriba, Taboada, Doallo, & Posada, 2012). Model GRT + G was selected for rRNA 12S and rRNA 16S, TN93 + G for COI, and HKY for Cyt b. The gamma distribution was estimated where needed from the data using four rate categories. Constant population size and strict clock (Drummond, Ho, Phillips, & Rambaut, 2006) were defined as priors. The calibrated Yule model, which is recommended for analyses using sequences
from different species (Drummond & Bouckaert, 2015), was selected as prior. Two independent MCMC chains were run for 200 million generations each, sampling values every 100,000 steps. Tracer files were examined using Tracer v1.7 (Rambaut, Drummond, Xie, Baele, & Suchard, 2018). Convergence and stationarity were verified for model parameters by visual inspecting of marginal densities and effective sample sizes ($226 < \text{ESS} > 1801$) in Tracer v1.7 (Rambaut et al., 2018). The posterior trees were summarized in TreeAnnotator using common ancestor heights to obtain the maximum clade credibility tree. Trees were visualized and edited using FigTree v1.4.3 (Rambaut, 2016).

3 | RESULTS AND DISCUSSION

3.1 | Three-dimensional geometric morphometric analysis of skull diversity in shape space

The three skeletal states are distributed in the species of *Brachycephalus* as follows (Figure 3). Nonhyperossified: *B. brunneus*, *B. didactylus*, *B. ferruginus*, *B. hermogenesi*, *B. izecksohni*, *B. pernix*, *B. pombali*, and *B. pulex*. Hyperossified bufoniform without dorsal shield: *B. alipioi*, *B. crispus*, *B. guarani*, *B. nodoterga*, *B. pitanga*, *B. toby*,

![Figure 3](image-url)
and *B. vertebralis*. Hyperossified bufoniform with dorsal shield: *B. ephippium*, *B. garbeanus*, and *B. margaritatus*.

Skull diversity in the species of *Brachycephalus* in ordinated shape space was analyzed with three procedures. In the first procedure, which was fully automated, we superimposed the virtual surfaces of the skulls of *Brachycephalus* reconstructed from 3D microcomputed tomography using GPSA. In GPSA, the optimal superposition of 3D surfaces is achieved by aligning the first and second longest axes of the surfaces, which are computed as the first two principal components of the array of 3D points making up the surfaces (Pomidor et al., 2016). This method failed, however, to correctly align the multiple 3D surfaces of *Brachycephalus*. We observed a similar outcome when we superimposed mandibles from several species of bats (Perez et al., personal observation). This suggests that the alignment algorithm used by GPSA is sensitive to objects whose major axes of variation are not well defined. Therefore, alternative algorithms will have to be devised to handle shapes that lack well defined major axes of variation. Here, we solved this problem by using instead the reconstructed images of the skull plus the vertebral column, which were then successfully superimposed using GPSA. The two other procedures we used were based on user-determined and manually registered reference points, although they differed in the superposition algorithm, which was global in one case and local in the other.

Diversity of skull shape in *Brachycephalus* is visualized in the ordinated space of the first two nmMDS axes (Figure 4). Color polygons were drawn by hand and bound the four conditions of skeletal states-Baupläne, namely, nonhyperossified leptodactyliform, nonhyperossified bufoniform, hyperossified bufoniform without dorsal shield, and hyperossified bufoniform with dorsal shield. Species of *Brachycephalus* in each of the four conditions of skeletal states-Baupläne differ markedly in skull shape, as demonstrated by nonoverlapping polygons computed from PSM, g-Procrustes, and 1-Procrustes distance matrices (Figure 4a–c). The marked dispersion in skull shape of the nonhyperossified leptodactyliforms, *B. didactylus*, *B. hermogenesi*, and *B. pulex*, relative to species of *Brachycephalus* in the other three conditions of skeletal states, is noteworthy. This pattern emerges in the space of the first two nmMDS axes computed with landmarks using the g-Procrustes and 1-Procrustes distance matrices (Figure 4b,c). Differences in skull shape between the species of *Brachycephalus* relative to the prototype are visualized using GPSA as a three dimensional heat map (Figure 5). In this map, the colors blue and red indicate low and high variance values, respectively. Differences in skull shape between the species of *Brachycephalus* involve primarily the frontoparietal and otic bones (Figure 5).

### 3.2 Three-dimensional geometric morphometric analysis of skull diversity in size-shape space

Skull diversity in the species of *Brachycephalus* is visualized in the ordinated size-shape space defined by RSC-1 and CAC (Figure 6). Color polygons were drawn by hand and bound the four conditions of skeletal states-Baupläne, namely, nonhyperossified leptodactyliform, nonhyperossified bufoniform, hyperossified bufoniform without dorsal shield, and hyperossified bufoniform with dorsal shield. The pattern of skull diversity in *Brachycephalus* that emerges from the ordinated size-shape space is similar to that in the ordinated shape space (Figures 4a–c and 6). That is, diversity in skull morphology amongst species of *Brachycephalus* is partitioned according to each of the four conditions of skeletal states-Baupläne. Again, the outstanding outcome is the marked dispersion in skull morphology of the nonhyperossified leptodactyliform species, *B. didactylus*, *B. hermogenesi*, and *B. pulex* in the space defined by the common allometric component and the first residual component (Figure 6). Another noticeable outcome is the partial overlap between nonhyperossified leptodactyliforms and nonhyperossified bufoniforms, which is caused by a single species, *B. didactylus* (Figure 6). Because this result emerged in size-shape space rather than in shape space, we conjecture that interactions between size and shape unique to *B. didactylus* determine that the skull of this nonhyperossified leptodactyliform species is more similar to those of nonhyperossified bufoniform species (Figure 6). We cannot however at this point offer any mechanistic explanation for this phenomenon.

The common allometric component scales linearly with size measured by the log of centroid size (Figure 7). Consequently, the skull diversity of *Brachycephalus* that emerged in the ordinated size-shape space does not depart from allometry. That is, diversity in skull shape in *Brachycephalus* can be explained by allometry. Paluh et al. (2020) sampled all hyperossified frog genera and their sister lineages totaling 158 species and 145 genera of 54 anuran families, and detected no departure from allometry. Thus, this macroevolutionary trend for anurans appears to hold for the case of the relatively smaller *Brachycephalus* radiation. Shape variation in the skull of *Brachycephalus* as a function of size, that is, allometry, is modelled as deformation in the size-shape formalism (Mitteroecker et al., 2004). Deformation shows that in the direction of increasing size, the common allometric component consists of a marked broadening of the frontoparietal bones in the skull vault and a posterolateral displacement of the otic bones in the otic region (Figure 7). Paluh, Stanley, and Blackburn (2020)
identified the relative length and width of the frontoparietal bones as a factor driving shape differences in the skull of all hyperossified frog genera. At a larger evolutionary scale, Bardua et al. (2020) also identified the frontoparietal and otic bones as part of 13 modules of 19 cranial regions in their analysis of the frog skull.

Whereas there is an approximately linear relationship between skull shape and size, the same is not true of the relationship between hyperossification and size (Figures 7 and 8). The degree of hyperossification in

**FIGURE 4** Ordinated shape space for the species of *Brachycephalus* derived from nonmetric multidimensional scaling. (a) Global superimposition of automated virtual surfaces. (b) Global superimposition of manually registered reference points. (c) Local superimposition of manually registered reference points. Orange, red, green, and blue polygons bound nonhyperossified leptodactyliform species, nonhyperossified bufoniform species, hyperossified bufoniform species without dorsal shield, and hyperossified bufoniform species with dorsal shield, respectively.

**FIGURE 5** Variation in skull shape between species of *Brachycephalus* relative to the prototype visualized as a three-dimensional heat map derived from the virtual surfaces using GPSA (Pomidor et al., 2016). Blue and red colors indicate low and high values of shape variation, respectively, from the designated, mean prototype surface, *Brachycephalus ferruginus*. 
Brachycephalus does not increase monotonically with increasing body size, as nonhyperossified bufoniform species and hyperossified bufoniform species without dorsal shield do share identical mean body sizes (Figure 8).

3.3 Robustness of pattern of skull diversity as a function of data acquisition and statistical formalism

High-resolution 3D microcomputed tomography imaging in connection with 3DGM revealed that the species of Brachycephalus differ in skull morphology. Furthermore, the differences are associated with the four conditions of skeletal states-Baupläne (Figures 4, 6, and 7). This pattern is to a large extent independent of the algorithm of superposition, whether global or local, and of data acquisition that is, automated, landmark-free or manually registered, landmark-based. Automated landmark-free methods sample phenotypes intensively and generate high-dimensional data (Gao et al., 2018). Conversely, such intensive sampling is not achieved by user-defined methods that rely on manually registered reference points. Nevertheless, methods such as the GPSA that use automated virtual surfaces generate maps between surfaces that lack the pointwise homology of user-defined landmarks (Pomidor et al., 2016). That is, they lack transitivity, which yields global consistency of pairwise mappings between surfaces (Gao et al., 2018). This problem is addressed in GPSA by optimally associating points in different surfaces with their nearest neighbors (Pomidor et al., 2016). Consequently, a relevant question is whether

**FIGURE 6** Common allometric component scores plotted against the first residual shape scores for Brachycephalus. Orange, red, green, and blue polygons bound nonhyperossified leptodactyliform species, nonhyperossified bufoniform species, hyperossified bufoniform species without dorsal shield, and hyperossified bufoniform species with dorsal shield, respectively.

**FIGURE 7** Log centroid size plotted against the common allometric component scores for Brachycephalus. Three-dimensional images of the skull of Brachycephalus show changes in shape as a function of the increase in the common allometric component. Orange, red, green, and blue polygons bound nonhyperossified leptodactyliform species, nonhyperossified bufoniform species, hyperossified bufoniform species without dorsal shield, and hyperossified bufoniform species with dorsal shield, respectively.
3DGM analyses predicated on user-based or automated methods of data gathering yield ordinated shape spaces that are mutually consistent (Gao et al., 2018). Furthermore, as pointed out by Gao et al. (2018), GPSA may produce a Y pattern of interspecific ordination. Here, however, the ordinated shape space generated by GPSA did not produce a Y pattern of interspecific ordination (Figure 4a). The landmark-free automated method thus behaved as well as the manually registered, landmark-based method.

However, we must point out that the larger diversity in skull morphology of the nonhyperossified lepadactyliform species, *B. didactylus*, *B. hermogenesi*, and *B. pulex*, relative to that of species of *Brachycephalus* in the other three conditions of skeletal states-Baupläne, emerged only from the landmarks-based dataset. This finding is relevant because landmarks defined by experienced morphometricians are regarded as providing a “ground truth” with respect to the derived ordinated size and/or size-shape space (Gao et al., 2018; Gao, Kovalsky, Boyer, & Daubechies, 2019). Therefore, the landmark-based method of data acquisition will likely remain relevant as a tool to reveal morphological variability and diversity.

### 3.4 | Species tree estimation

Our mitochondrial species tree revealed that *B. pulex* is sister to all species of *Brachycephalus* (Figure 9). Next
in the branching order, *Brachycephalus brunneus*, *B. ferruginus*, *B. izecksohni*, *B. permix*, and *B. pombali* are monophyletic. This lineage is the sister to all other species and *B. hermogenesi* is the sister to the remaining species. *Brachycephalus didactylus* is sister to two monophyletic lineages. One lineage includes *B. ephippium*, *B. garbeanus*, and *B. margaritatus*. The other monophyletic lineage includes *B. alipioi*, *B. crispus*, *B. guarani*, *B. nodoterga*, *B. pitanga*, *B. toby*, and *B. vertebralis*. Earlier, Clemente-Carvalho et al. (2011) estimated a species tree for *Brachycephalus* based on mitochondrial genes that sampled a single specimen per species, which yielded a trichotomy with many branches lacking adequate support, as measured by posterior probabilities. Our current species tree, which sampled multiple individuals per species, resolved the phylogenetic relationships between the species of *Brachycephalus* sampled here reasonably well. Our mitochondrial species tree implies the following about the evolution of the conditions of skeletal states—Baupläne in *Brachycephalus*. First, the nonhyperossified skeletal state is the ancestral condition for *Brachycephalus*. Second, the hyperossified skeletal state evolved once in *Brachycephalus*, with the appearance of a dorsal shield in the lineage leading to *B. ephippium*, *B. garbeanus*, and *B. margaritatus*. Third, the leptocephaliform Bauplan is the ancestral condition for *Brachycephalus*, whereas the bufoniform Bauplan evolved twice independently in *Brachycephalus*. Recently, Condez et al. (2020) demonstrated quantitatively that the nonhyperossified skeletal state is indeed the ancestral skeletal state in *Brachycephalus*.

The most striking result, however, is the lack of monophyly of the nonhyperossified leptocephaliform species of *Brachycephalus*. Condez et al.’s (2020) also demonstrated the same result with their mitochondrial species tree. The branching order of the nonhyperossified leptocephaliform species differs though between Condez et al.’s (2020) and our mitochondrial species trees (present work). In Condez et al.’s (2020) mitochondrial species tree *B. hermogenesi* is the sister taxon to all other *Brachycephalus*, whereas *B. pulex* is the sister taxon in our species tree. Remarkably, the phylogenetic diversity of the nonhyperossified leptocephaliform revealed by the mitochondrial species tree is mirrored by our analysis of the diversity in skull morphology in shape and size-shape space (Figures 4, 6, and 9). Skull diversity in the nonhyperossified leptocephaliform species of *Brachycephalus* is markedly larger when compared to skull diversity in species in the other three conditions of skeletal states—Baupläne. The diversity of nonhyperossified leptocephaliforms is not limited, however, to skull morphology or molecular phylogenetic branching patterns. Recently, Condez et al. (2020) described a fourth species of nonhyperossified leptocephaliform, *B. sulfuratus* and, in addition, discovered three putative new species of nonhyperossified leptocephaliforms based on their magnitude of molecular variation (Condez et al., 2020). *Brachycephalus sulfuratus* extended the distribution of the nonhyperossified leptocephaliform species from the State of Bahia in Northeastern Brazil (Napoli et al., 2011) into the State of Santa Catarina in Southern Brazil (Condez et al., 2016). The discovery of *B. sulfuratus* revealed that the geographic distribution of the nonhyperossified leptocephaliform species is larger than the distribution of any of the three other conditions of skeletal states—Baupläne.

The nonhyperossified leptocephaliform condition thus emerges as evolutionarily complex in terms of morphological, molecular diversity, and geographic distribution. Therefore, additional morphological, molecular, and distributional data for the nonhyperossified leptocephaliform skeletal state—Baupläne condition will be needed to further understand the evolutionary history of *Brachycephalus*.

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Caio dos Santos: Formal analysis; methodology; writing-review and editing. S. Perez: Conceptualization; formal analysis; methodology; writing-original draft; writing-review and editing. Prianda Laborda: Formal analysis; methodology; writing-review and editing. Ricardo Lopes: Formal analysis; funding acquisition; methodology; writing-review and editing. Rute Clemente-Carvalho: Formal analysis; methodology; writing-review and editing. Sergio dos Reis: Conceptualization; formal analysis; methodology; writing-original draft; writing-review and editing. Fernando Von Zuben: Formal analysis; methodology; software; writing-original draft; writing-review and editing.

CONFLICT OF INTEREST
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