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MORPHOLOGICAL DISPARITY IN THEROPOD JAWS: COMPARING DISCRETE CHARACTERS AND GEOMETRIC MORPHOMETRICS

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Abstract: Disparity, the diversity of form and function of organisms, can be assessed from cladistic or phenetic characters, and from discrete characters or continuous characters such as landmarks, outlines, or ratios. But do these different methods of assessing disparity provide comparable results? Here we provide evidence that all metrics correlate significantly with each other and capture similar patterns of morphological variation. We compare three methods of capturing morphological disparity (discrete characters, geometric morphometric outlines and geometric morphometric landmarks) in coelurosaurian dinosaurs. We standardize our study by focusing all our metrics on the mandible, so avoiding the risk of confounding disparity methods with anatomical coverage of the taxa. The correlation is strongest between the two geometric morphometric methods, and weaker between the morphometric methods and the discrete characters. By using phylogenetic simulations of discrete character and geometric morphometric data sets, we show that the strength of these correlations is significantly greater than expected from the evolution of random data under Brownian motion. All disparity metrics confirm that Maniraptoriformes had the highest disparity of all coelurosaurians, and omnivores and herbivores had higher disparity than carnivores.

Key words: dinosaur, coelurosaur, maniraptoriform, disparity, morphometrics.

Studies of the amount of morphological variation, commonly referred to as ’disparity’, have become common in palaeontology. It might be argued that disparity (form) and diversity (species richness) should track each other in a model of homogenous evolution, but they are frequently decoupled, with some clades showing high species richness but limited diversity of form, and smaller clades showing high disparity (Wills et al. 1994; Fortey et al. 1996; Foote 1997; Ruta et al. 2013). Further, disparity is often high early in the evolution of a clade, suggesting some kind of ‘early burst’ model of evolution (Foote 1997; Erwin 2007). Disparity studies have provided insights into the evolution of novel body plans and ecological innovations (Goswami & Polly 2010; Brusatte et al. 2014; Deline et al. 2018), the impact and selectivity of mass extinction events (Brusatte et al. 2008; Friedman 2009; Bapst et al. 2012), and morphological expansion during evolutionary radiations (Foote 1997; Erwin 2007; Hughes et al. 2013; Stubbs et al. 2013; Close et al. 2015; Cooney et al. 2017).

Disparity should be considered in a comparative framework, and there are several analytical approaches. The most common methods use discrete descriptive characters or geometric morphometrics. Describing morphological variation using discrete characters has usually focused on cladistic data sets as a ready source of rich data on trait variation (Wills et al. 1994; Lloyd 2016; Gerber 2019). This approach involves analysing character–taxon matrices where morphologies are scored using character states, including the presence and absence of features, the numbers of certain elements (e.g. teeth or limbs), the relationships between, or orientation of, elements and even general features relating to size and shape. Geometric morphometric methods, such as landmark coordinates and outlines, measure the shape of a structure, with outlines measuring the outer margin of a morphology and landmarks measuring the location of homologous features in a Cartesian coordinate system (MacLeod 1999; Zelditch et al. 2012).

These methods can be used in different circumstances. For some studies, geometric morphometric analyses are not possible due to a lack of homologous points, the complexity of the morphology, or a lack of completely preserved specimens, in which case discrete characters...
may represent a suitable alternative. In other studies, observing shape changes, and linking these to evolutionary hypotheses, is pivotal, and discrete characters are not appropriate. Because these methods are often used interchangeably in the literature to describe disparity, it is important to consider whether all methods give similar results or not, when discussing large-scale evolutionary patterns. If not, then the methods must be applied and interpreted with extra care. Most comparisons of different disparity methods (Villier & Eble 2004; Anderson et al. 2011; Anderson & Friedman 2012; Foth et al. 2012; Hetherington et al. 2015; Hopkins 2017; Maclaren et al. 2017; Romano 2017) have shown similar results, but more or less strongly. On the other hand, Mongiardino Koch et al. (2017) found disagreements when comparing traditional morphometric data and discrete characters, and they strongly advocated incorporating a phylogenetic framework.

Here we compare different methods of assessing disparity, including the two main methods in geometric morphometrics (landmarks and outlines), as well as discrete cladistic characters. We use a case study focusing on a single anatomical region, to ensure that we compare like with like. Our case study looks at coelurosaurian dinosaur mandibles, and we use these for several reasons: they have a good fossil record, all clades and time intervals are sampled by multiple specimens, often with complete mandibles preserved. Further, they have a wide range of morphologies, ranging from the elongated jaws in Mesozoic birds to the robust jaws in tyrannosaurs, and the bizarre oviraptorids (Weishampel et al. 2004), often associated with different diets. Many coelurosaurians were carnivores, while some clades, such as therizinosaurs, oviraptorids and birds show specializations for herbivory or omnivory (Zanno & Makovicky 2011). Mandibular disparity can also be effectively measured using all analytical approaches. Finally, disparity in vertebrate jaws has been the subject of previous studies, and it is accepted that characters of the mandible and it is accepted that characters of the mandible and ecomorphological and morphometric analyses, the jaw samples had to be complete and undistorted, but they need not have been statistically. The 40 coelurosaurian taxa used in these analyses had mandibles that were complete, without taphonomic distortion and had also been coded for jaw and dental characters in the discrete character matrix of Brusatte et al. (2014) (see Schaeffer et al. 2019, tables S1, S3). Most taxa are known from a single mandible fossil, but for species where multiple specimens exist, a single representative was chosen for the geometric shape analyses, but the discrete character analyses sampled all material to maximize coding. In the second series of extended analyses, we used the maximum possible sample size available based on the restrictions of the methods. For inclusion within the discrete character analyses, the taxa had to be coded for mandibular characters in the character matrix from Brusatte et al. (2014), even if the specimens were fragmentary or partially incomplete. We used the function TrimMorphDistMatrix from Claditis (Lloyd 2016) to remove highly incomplete taxa that generated non-applicable distances due to a lack of shared characters, leaving a sample size of 89 taxa (see Schaeffer et al. 2019, tables S2, S4). For the extended geometric morphometric analyses, the jaw samples had to be complete and undistorted, but they need not have been included within the character matrix of Brusatte et al. (2014), giving a sample size of 60 (see Schaeffer et al. 2019, table S5).

**Comparative groupings**

We examined morphospaces and calculated disparity statistics for comparative groupings, aiming to replicate the types of analyses common in the literature. We quantified disparity in clades, firstly comparing the two major coelurosaurian groups, Maniraptoriformes and Tyrannosauroidea, and then the following subgroups within Maniraptoriformes: Dromaeosauridae, Aves, Oviraptorosauria and Ornithomimosauria. Too few taxa were sampled from Troodontidae, Scansoriopterygidae and Therizinosauria, so these maniraptoriform subgroups were excluded from the disparity calculations. We also compared disparity in three broad ecological groupings, encompassing the dietary and size diversity within coelurosaurians: small carnivores (<4 m total body length), large carnivores (>4 m total body length) and herbivores plus omnivores. Dietary categories were assigned based on consideration of a wide range of features of anatomy and associated fossils, from the literature (Zanno & Makovicky 2011; Brusatte et al. 2012; Foth & Rauhut 2013). Herbivores and omnivores were grouped together because it is difficult to distinguish between

**MATERIAL AND METHOD**

**Taxon sampling**

We followed two sampling approaches. In the first, a consistent sample of 40 coelurosaurian taxa was used across all three analytical approaches, so they had identical composition in terms of phylogenetic and temporal coverage, and inter-taxon distances could be directly compared statistically. The 40 coelurosaurian taxa used in these analyses had mandibles that were complete, without taphonomic distortion and had also been coded for jaw and dental characters in the discrete character matrix of Brusatte et al. (2014) (see Schaeffer et al. 2019, tables S1, S3). Most taxa are known from a single mandible fossil, but for species where multiple specimens exist, a single representative was chosen for the geometric shape analyses, but the discrete character analyses sampled all material to maximize coding. In the second series of extended analyses, we used the maximum possible sample size available based on the restrictions of the methods. For inclusion within the discrete character analyses, the taxa had to be coded for mandibular characters in the character matrix from Brusatte et al. (2014), even if the specimens were fragmentary or partially incomplete. We used the function TrimMorphDistMatrix from Claditis (Lloyd 2016) to remove highly incomplete taxa that generated non-applicable distances due to a lack of shared characters, leaving a sample size of 89 taxa (see Schaeffer et al. 2019, tables S2, S4). For the extended geometric morphometric analyses, the jaw samples had to be complete and undistorted, but they need not have been included within the character matrix of Brusatte et al. (2014), giving a sample size of 60 (see Schaeffer et al. 2019, table S5).

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them using dietary proxies. All taxa were classified into groups using the literature (Schaeffer et al. 2019, tables S3–S6).

**Discrete character analyses**

Character-based disparity was assessed using the coelurosaurian discrete character matrix of Brusatte et al. (2014). First the data set was reduced to contain only mandibular and dentary tooth characters (78 characters, see Schaeffer et al. 2019, tables S1, S2). As previously noted, we first ran analyses with a reduced sample to match the geometric morphometric analyses (40 taxa with complete mandibles), and then additional analyses were run using the full sample. The inclusion of dentary teeth characters within the analyses was assessed by running supplementary analyses using a mandible-only data set (66 characters).

The character matrices were converted into pairwise distance matrices based on maximum observable rescaled distances (MORD; Lloyd 2016). The distance matrices were then subjected to principal coordinates analysis (PCOA) with Cailliez negative eigenvalue correction (Cailliez 1983), and the latter had the effect of reducing the proportion of variance expressed by each PCO-axis compared to uncorrected PCOA (i.e. PCO1 10.7% vs 25.1% and PCO2 9.6% vs 21.3%) (Schaeffer et al. 2019, figs S1, S2). However, the distribution of taxa in morphospace is almost identical in biplots generated from the major axes of variation (PCO1–PCO5), as previously noted by Hopkins (2017) and Nordén et al. (2018). Therefore, although the percentage of variance reported is comparatively low, the biplots of PCO1–PCO2 morphospace still represent a decent proportion of overall variance and illustrate the major axes of variation. The disparity metric sum of variances (SOV) was calculated from all coordinate axes scores for each of the comparative groupings. The SOV metric is generally robust to sample size differences in comparative groupings (Ciampaglio et al. 2001), but to test this we also calculated SOV with rarefaction to equalize sample sizes based on the group with smallest sample size. All analyses were performed using the packages phytools (Revell 2012), Cladistics (Lloyd 2016), dispRity (Guillerme 2018) and vegan (Oksanen et al. 2017), all in the R coding environment (R Core Team 2017).

**Geometric morphometric analyses**

The geometric shape analyses were based on lateral profiles of the mandibles in two dimensions. We first assembled a collection of lateral mandible images from museum specimens and the primary literature (Schaeffer et al. 2019, table S8). For the outline analyses, the jaw samples were converted to silhouettes, excluding the dentition, and imported into R. We used elliptical Fourier analysis (EFA) to compare outline shapes with the R package Momocs (Bonhomme et al. 2014; Navarro et al. 2018). The jaws were converted to digitized outline closed curves (Fig. 1C), and then aligned, centred and scaled, before a set of 500 x and y-coordinate points was assigned to each outline profile. The number of harmonics required to account for the shape variation was calibrated. Each harmonic yielded four Fourier coefficients describing the shapes. Fourier coefficient data accounting for 99% of the shape variation in the original jaws (here coefficients from 11 EFA harmonics) was subjected to principal component analyses (PCA) to ordinate the data, explore major aspects of the geometric variation and generate morphospaces using the first two principal components axes. The same jaw samples were used in the landmark geometric analyses. Landmarks were digitally added to the specimens using tpsDig2 and tpsUtil (Rohlf 2017a, b) and consisted of 6 fixed landmarks and 50 semi-landmark points along 6 curves (Bookstein 1991) (Fig. 1B, Schaeffer et al. 2019, files S1–S4). To remove noise effects such as size and orientation, Procrustes aligned landmarks were calculated using a generalized Procrustes analysis in tpsRelw, that incorporated a sliding procedure for minimizing bending energy (Rohlf 2017c). The Procrustes-aligned landmark data were then subjected to PCA in the R package geomorph (Adams & Otárola-Castillo 2013), and morphospaces were created using the scores from the first two axes. For both geometric morphometric analyses, the function PCcontrib from the R package Momocs (Bonhomme et al. 2014) was used to visualize and plot shape changes along principal component axes 1, 2 and 3, and the sum of variances disparity statistic was calculated for the comparative groupings using all morphospace axes scores with the R package dispRity, both with and without rarefaction (Guillerme 2018).

**Correlation tests**

Morphological disparity can be expressed as pairwise distances between taxa. To assess whether the three measures of disparity, based on characters, landmarks and outlines, provide similar insights into morphological disparity, we used Mantel tests (Anderson & Friedman 2012; Hetherington et al. 2015) to examine the correlation between pairwise distances derived from each sample-standardized analysis. Pairwise distances for the three data sets are based on Euclidean distances in the multidimensional morphospace coordinates from all axes, generated from the protocols described above. Ancillary tests were
performed based on pairwise distances from character data with dentary teeth characters excluded, to ensure that the inclusion or exclusion of teeth in the character data was not significantly impacting the measured disparity.

**Phylogenetic simulation tests**

To provide a null model with which to compare our empirical correlation results we used a simulation approach. We tested whether the empirical correlation results between different data sets describing the same anatomical unit are significantly different to correlations expected from the evolution of unrelated traits or morphological data sets evolving under Brownian motion on a phylogeny.

For the phylogenetic data we sampled 10 coelurosaurian trees from Brusatte *et al.* (2014). To calculate phylogenetic branch lengths, each topology was time-calibrated 50 times using both the equal (Brusatte *et al.* 2008) and minimum branch length (Laurin 2004) dating approaches with the `timePaleoPhy` function of the `paleotree` R package (Bapst 2012). For each dating replicate, ages were randomly sampled from between each taxon’s first and last appearance dates (age data from Brusatte *et al.* 2014). The resulting 1000 coelurosaurian trees therefore incorporated phylogenetic uncertainty (10 topologies), two dating methods, and temporal occurrence uncertainty for the tips (through the replicates). All trees were cropped to contain only the 40 taxa that were present in the empirical correlation tests. We then simulated two types of morphological data, with similar properties to those in our study, for each of the 1000 dated phylogenies.

First, we simulated discrete character data sets. The function `sim.morpho` from the R package `dispRity` (Guillerme 2018) was used to simulate data sets with 78 characters (77% of which had two states and 23% had three states) using the equal-rates (ER = Mk) model (Paradis & Schliep 2018). For the model parameters of each simulated data set, we randomly sampled rates of 5, 10 or 20 (higher rates increased phylogenetic signal) and distribution shapes of 0.5, 1 or 2. Missing data, ranging from 21.5% to 37.8%, was introduced to each simulated discrete character data set to reflect the nature of the empirical jaw character data (and fossil-based discrete character data in general). The simulated discrete character data were converted into MORD pairwise distance matrices and subjected to PCOA with Cailliez negative eigenvalue correction (Cailliez 1983), giving 1000 sets of PCOA scores for 40 taxa.

Next, we simulated geometric shape data evolving under Brownian motion for the same 1000 dated coelurosaurian trees. We aimed to generate 40 ‘jaw-like’ structures defined by 56 landmark coordinates for each tree. As a starting

![Fig. 1.](image)
point, we used the consensus shape from our sample of 40 theropod jaws as the ancestral morphology. The method required co-variance data to guide how the shapes would transform during the simulations, so we calculated Procrustes residuals from our original landmark configurations as input. With these three inputs, a phylogeny, ancestral morphology and residuals data, we simulated landmark configurations at all nodes and tips evolving under phylogenetic Brownian motion for all 1000 topologies, using custom code built around the SimEvo function from Evomorph (Cabrera & Giri 2016). This model simulates shapes by calculating the product of a phenotypic covariance matrix (which is the Procrustes residuals not the phylogenetic variance co-variance matrix) with the vector $\beta H$. The $H$ component transforms the $P$ matrix into the $G$ matrix which describes the heritable proportion of $P$ (Polly 2004), with $\beta$ summarizing the selection coefficients (i.e. morphological change over time $= \beta H^*P$). This is identical to the formulation in SimEvo, except that instead of running the simulation for $n$ generations, our method runs a single generation in which variation of $\beta H$ is sampled according to a mean zero normal distribution with variance equal to its branch length. As a Brownian process evolution occurs independently on each branch, with each descendant branch inheriting the shape from its parent. After running the simulations, the 1000 sets of 40 landmark configurations were converted to TPS format and subjected to PCA in geconnect (Adams & Otárola-Castillo 2013), giving 1000 sets of PC scores (code is available in Schaeffer et al. 2019).

Mantel correlation tests were performed for each pair of simulated data sets (characters and landmarks simulated from the same tree) based on pairwise Euclidean distances from the multidimensional morphospace coordinates from all axes. The distribution of correlation statistics from these 1000 paired simulated data sets was then statistically compared to the observed correlation results from the jaw discrete character and geometric shape data sets, to test the null hypothesis of no difference between the simulated data sets and real data.

## RESULTS

### Morphospace occupation

The morphospace biplots (Figs 2, 3) illustrate the morphological dissimilarity between taxa based on major features of morphological variation and they show divisions according to taxonomic grouping (Fig. 2) and dietary category (Fig. 3). In both figures, the morphospace axes represent the same major features of variation, so this is described first, before exploring how the taxonomic and dietary groupings are distributed in morphospace.

In the outline morphospaces, using the standardized sample of 40 jaws (Figs 2A, 3A), PC1 describes 61.4% of variation, representing the thickness of the dentary region and jaw mid-length, while PC2 (23.2%) encapsulates variation in the dorsoventral curvature of the mandible and the relative position of the cranium–mandible articulation (Fig. 4). The remaining outline morphospace axes (PC3–PC40) describe 15.4% of variation (Schaeffer et al. 2019, fig. S3). In the standardized landmark analyses (Figs 2B, 3B, 4), PC1 (40.7%) reflects changes in jaw depth, particularly at the mid-length and anteriorly, like the outline analysis. However, PC2 (22.8%) represents the overall thickness of the jaw, unlike the outline analysis. Variation in the dorsoventral curvature of the jaw is represented by PC3 (13.6%) in the landmark analysis (Schaeffer et al. 2019, fig. S4). The remaining axes in the landmark analysis (PC4–PC40) describe a further 22.9% of variation (Schaeffer et al. 2019, fig. S4). In the standardized discrete characters analyses (Figs 2C, 3C), PCO1 accounts for 10.7% of variation while PCO2 represents a further 9.6% (Schaeffer et al. 2019, fig. S2). As previously noted, these relatively low percentages of variation are the combined result of having many characters, missing data and using negative eigenvalue correction during PCOA (Hopkins 2017; Nordén et al. 2018).

In the morphospaces highlighting taxonomic groupings we compare the distribution of maniraptoriforms and tyrannosauroids (Fig. 2). In all analyses, the former occupies a much larger region of morphospace than the latter. The jaw outlines morphospace (Fig. 2A) shows a near-complete separation between maniraptoriforms and tyrannosauroids. Three basal tyrannosauroids, Proceratosaurus, Guanlong and Dilong, overlap with maniraptoriforms in central morphospace. The maniraptoriforms occupy a large range along PC1, mainly due to the divergent oviraptoroids, and a smaller range along PC2, whereas the tyrannosauroids have a large range on PC2 but limited expanse on PC1. The landmarks morphospace (Fig. 2B) shows complete overlap of the two groups. The maniraptoriforms occupy a broad area that is extended along both PC1 and PC2, whereas tyrannosauroids have limited expanse on both PC1 and PC2. The discrepancy between the two morphometric morphospaces results from different aspects of shape variation being represented by PC2 (Fig. 4). If landmark morphospace is plotted based on PC1 and PC3, the overall pattern is similar to the outline morphospace, with tyrannosauroids diverging on PC3 (Schaeffer et al. 2019, fig. S7). The discrete characters morphospace (Fig. 2C) shows similar patterns to the outlines morphospace (Fig. 2A), with two basal tyrannosauroids, Proceratosaurus and Guanlong, overlapping with maniraptoriforms, while tyrannosauroids generally occupy a distinct area of morphospace and a restricted range of PCO1, but expand on PCO2. The
FIG. 2. Patterns of morphospace occupation and disparity (sum of variances, SOV) of Coelurosauria grouped according to taxonomy, measured using three metrics. The bivariate morphospaces and disparity plots were created using: A, D, geometric outlines; B, E, geometric landmarks; C, F, discrete jaw and dental characters. Legend: triangles, Tyrannosauridae; circles, Maniraptoriformes; stripes (D–F), disparity for the Ornithomimidae, Oviraptoridae, Dromaeosauridae and Avialae. In the morphospace plots (A–C) open circles denote coelurosaurians that are not classified as Tyrannosauridae or Maniraptoriformes, namely Sinosauropteryx prima, Compsognathus longipes and Ornitholestes hermanni. Dinosaur silhouettes from http://phylopic.org: Scott Hartman, Nobu Tamura, Brad McFeeters, Emily Willoughby and Matt Martyniuk (CC BY 3.0). Colour online.
FIG. 3. Patterns of morphospace occupation and disparity (sum of variances, SOV) of Coelurosauria grouped according to diet, measured using three metrics. The bivariate morphospaces and disparity plots were created using: A, D, geometric outlines; B, E, geometric landmarks; C, F, discrete jaw and dental characters. Legend: circles, small carnivores; dark circles, large carnivores; triangles, omnivores and herbivores. Dinosaur silhouettes from http://phylopic.org: Scott Hartman, Brad McFeeters and Emily Willoughby (CC BY 3.0). Colour online.
Maniraptoriforms occupy a wide zone, particularly extending along PCO1 due to the divergent oviraptorids. Morphospace plots with all taxa individually labelled are provided in Schaeffer et al. (2019, figs S5, S6, S8, S9). When using a reduced discrete character data set of 66 characters, based only on the mandibular bones (excluding teeth), the overall distribution of taxa is similar, as is the relative expanse of both clades on the major PCO axes (Schaeffer et al. 2019, fig. S10A).

In the morphospace plots showing the taxa distinguished by diet there are many commonalities in the distribution of groups and some nuanced differences (Fig. 3). No single morphotype or discrete character combination characterizes omnivores and herbivores, but instead the grouping is widely distributed in all morphospaces, occupying extreme positions on the major axes of variation and more central areas. The wide distribution in morphospace probably reflects the fact that this is an ill-defined dietary category. Some of the sampled small carnivores overlap with omnivore–herbivores in central morphospace, but the grouping does contain some divergent forms, particularly in the landmark morphospace on PC2 (Fig. 3B). The sampled large carnivorous taxa occupy a reduced area of morphospace in all analyses. Their morphospace is distinct from the small carnivores and omnivore–herbivores in the outline and character morphospaces, but not in the landmark morphospace (Fig. 3B). Large carnivorous taxa generally have jaws that are dorsoventrally deep in the

**FIG. 4.** Mandible shape changes along the three main principal components axes (PC1, PC2, PC3). A, shape changes based on outline and landmark approaches for the standardized sample (n = 40). B, shape changes based on outline and landmark approaches for the full sample (n = 60). The y-axes are the principal component axes. The x-axes illustrate the mean shape and the shapes at standard deviations for each of the PC-axes. Darker outlines are realized morphologies whereas lighter outlines are theoretical morphologies not shown by the sampled taxa.
postdentary region. Once again, when using a reduced discrete character data set with only mandibular bones the relative distribution and expanse of dietary groups in morphospace is consistent (Schaeffer et al. 2019, fig. S10B).

**Disparity patterns**

As with the morphospaces, the sum of variances (SOV) disparity patterns converge for the different metrics but show some subtle differences between the data types (Figs 2, 3). In the taxonomic results (Fig. 2), the tyrannosaurids always have lower mandibular disparity than the maniraptoriforms. These results are statistically significant, based on non-overlap of the 95% confidence intervals generated through bootstrapping, for the landmarks and characters (Fig. 2E, F, Schaeffer et al. 2019, fig. S10C) but not the outlines (Fig. 2D). The confidence intervals around the outline-based SOV statistic for maniraptoriforms are large, probably owing to the inclusion/exclusion of certain divergent oviraptorid taxa during the bootstrapping procedure.

Within Maniraptoriformes, the greatest disparity in the outline analysis (Fig. 2D) is seen in oviraptorids, and a smaller portion from the Dromaeosauridae, Avialae and Ornithomimosauria. The landmarks results (Fig. 2E) show a similar pattern, with the oviraptorids showing greatest disparity, but this time closely followed by the Avialae.

The character-based results (Fig. 2F, Schaeffer et al. 2019, fig. S10C) show something completely different. Within the Maniraptoriformes, the Avialae show the highest subgroup disparity, followed by the ornithomimosaurids, oviraptorids and dromaeosaurids, but all are within the error bar ranges of each other. The dietary groupings also show some differences between the three methods (Fig. 3). In all three cases, the omnivore–herbivore grouping shows highest disparity, but this is notably higher in the shape analyses (Fig. 3D, E) and only slightly so according to discrete characters (Fig. 3F) reflecting the fact that the category may include several feeding types that we cannot further subdivide. Further, the large carnivores always show lowest total disparity, but significantly so only for landmarks (Fig. 3E) and discrete characters (Fig. 3F, Schaeffer et al. 2019, fig. S10D). All disparity results are consistent when using rarefaction to standardize sample sizes, but the confidence envelopes are wider due to reduced sample sizes (Schaeffer et al. 2019, table S9).

**Correlation between morphological distances**

Across all correlation tests based on morphological distances, both Pearson and Spearman Mantel tests show evidence of statistically significant correlation at a 0.001 threshold level. The strength of correlations varies between tests. As expected, from the two geometric morphometric approaches, the outline and landmark-based distances, have a strong and significant correlation (Pearson $r = 0.791$, $r^2 = 0.626$, $p = 0.001$; Spearman $\rho = 0.579$, $p = 0.001$). The strength of correlations between morphological distances in the discrete character morphospace and the shape morphospaces are weaker. The discrete character data does show a relatively strong correlation with the outline distances (Pearson $r = 0.611$, $r^2 = 0.373$, $p = 0.001$; Spearman $\rho = 0.7147$, $p = 0.001$), while the weakest correlation between distances is recovered from tests of the discrete character distances and the landmark distances (Pearson $r = 0.502$, $r^2 = 0.252$, $p = 0.001$; Spearman $\rho = 0.491$, $p = 0.001$). When dental characters are excluded from the discrete character analyses, the correlation tests again show statistically significant correlations with the outline data (Pearson $r = 0.578$, $r^2 = 0.334$, $p = 0.001$; Spearman $\rho = 0.7278$, $p = 0.001$) and weaker correlation with the landmark data (Pearson $r = 0.452$, $r^2 = 0.205$, $p = 0.001$; Spearman $\rho = 0.4591$, $p = 0.001$). The correlation between morphological distances in discrete character morphospaces with and without dental characters is very strong and significant (Pearson $r = 0.964$, $r^2 = 0.929$, $p = 0.001$; Spearman $\rho = 0.970$, $p = 0.001$), suggesting the addition of 13 dantary tooth characters does not alter the character-based disparity patterns.

**Correlations and phylogenetic simulations**

Correlation tests performed on the phylogenetically simulated discrete character and geometric shape data also return statistically significant results, but the correlation coefficients and coefficients of determination are significantly lower than in the empirical jaw data sets. Mantel tests performed on the 1000 pairs of phylogenetically simulated discrete character data and shape data show that 81.1% (Pearson) and 90% (Spearman) of iterations give statistically significant correlations at a 0.05 threshold level, while 67.3% (Pearson) and 79.7% (Spearman) return statistically significant correlations at a 0.001 threshold. However, when comparing the distribution of correlation coefficients and coefficients of determination from the simulated results to the empirical results, it is clear that the Pearson’s $r$ values (outlines vs characters $p < 0.001$; landmarks vs characters $p = 0.001$), $r^2$ values and Spearman’s $\rho$ values (outlines vs characters $p < 0.001$; landmarks vs characters $p = 0.016$) are significantly higher in the real mandibular data than in the correlations from 1000 phylogenetically simulated character and shape data sets (Fig. 5).
Full sample size

For the full sample (Figs 6, 7) the same disparity patterns are present as in the standardized sample (Figs 2, 3). The relative distributions of tyrannosauroids and maniraptoriforms remain similar for all analyses (Fig. 6A–C), and the relative proportions of morphospace occupied by all clades remain comparable. In the two shape analyses, the same geometric changes are recovered on PC1–PC3 in the full sample size analyses (Fig. 4; although note that PC2 and PC3 are reversed), suggesting the inclusion of additional samples does not alter our understanding of the major shape innovations in coelurosaurian jaws. Expanding the sample for the geometric analyses introduced two outlying forms; the bird Longipteryx notably diverges along PC2 in the outline morphospace and the oviraptorid Gigantoraptor expands the bounds of PC2 in the landmark morphospace. When using the full sample, the maniraptoriforms again show significantly greater SOV disparity than the tyrannosauroids, but now in all three analyses (Fig. 6D–F), while the Avialae show relatively higher disparity in the outline analysis when compared to the standardized sample (Fig. 6D). In the discrete character analysis, the various maniraptoriform subclades show much more uniform (total disparities for the full data set (Fig. 6F) than for the standardized data set (Fig. 2F).

In terms of dietary categories, the morphospace and disparity patterns are also similar (Figs 3, 7). Omnivores–
herbivores again show the greatest disparity in all analyses. The disparity of large carnivores is greater when using the full sample, and the group is represented by 11 taxa rather than 6 taxa. This is particularly noticeable in the extended discrete characters study (Fig. 7C, F), where large carnivores now occupy a larger area of morphospace and overlap the small carnivores when more taxa are included. The relative proportions of the total morphospaces are similar between the reduced taxon study (Fig. 3D–F) and the study with all taxa (Fig. 7D–F), except for the relative levelling of total SOV for all three categories when sampling is increased (cf. Figs 3F, 7F).

All disparity results are consistent when using rarefaction to standardize sample sizes (Schaeffer et al. 2019, table S9).

**DISCUSSION**

Morphological disparity and a comparison of methods

The three methods of measuring morphological disparity share many commonalities and show moderate correlations between morphological distances. By incorporating a simulation approach, we tested whether the observed correlation results should be expected from random evolution under phylogenetic Brownian motion, and we used a novel modelling approach to simulate geometric data. It is widely accepted that shared phylogenetic history leads to phenotypic similarity between related species (Felsenstein 1985; Harvey & Pagel 1991; Freckleton et al. 2002). It is therefore unsurprising that our tests recover significant correlations between the majority of phylogenetically simulated shape and discrete character data sets, highlighting a common phylogenetic structure. However, the strength of the correlations between the three coelurosaur jaw data sets could not be replicated by simulations and the contrasting measures of morphological disparity are significantly more similar than expected from the evolution of random data under Brownian motion. Given that the shape analyses and discrete characters record morphological differences in very different ways, even if measured from the same anatomy, the moderate to strong correlations suggest that the three methods capture the same major patterns of morphological disparity, and this is not solely the outcome of random evolution along the branches of a phylogeny.

In light of our results, it is important to consider the extent to which we should expect the geometric shape and discrete character data to capture the same disparity patterns. Of the 78 jaw and dental characters used, 66 specifically relate to the mandible. However, only 28.8% of these characters (19 characters: numbers 1, 2, 3, 6, 8, 9, 14, 26, 28, 37, 42, 43, 44, 45, 47, 51, 61, 72, 78) describe overall shape variation, and just 18.2% of the 66 characters (12 characters: numbers 2, 3, 6, 14, 26, 28, 43, 45, 47, 51, 61, 78) could be captured by 2-D lateral jaw profiles in our analyses (other variation is in the dorsoventral shape or fenestrae shape/position). The remaining discrete mandibular characters encompass more nuanced morphological features, such as sutural contacts between constituent bones, presence, size and location of foramina and fossae, bone textures, pneumaticity, and features of muscle attachment sites. Therefore, it is perhaps surprising that we recover moderate correlations between the shape and character data. This may result from concerted evolution between geometry and other characters of the mandible, potentially highlighting morphological integration, an ecological constraint, or an underlying phylogenetic signal (Hetherington et al. 2015). Furthermore, it is surprising that correlations between the geometric data and discrete character data are stronger when teeth are included (all 78 characters), compared to comparisons with only the mandibular bones (66 characters). This implies that taxa with divergent jaw shapes share dental character scoring, and therefore the inclusion of dental characters does not erode the major dissimilarity patterns. Again, this could represent an ecological characteristic or an underlying phylogenetic signal.

Our study can be compared with previous endeavours, which have so far yielded differing results. Hetherington et al. (2015), for example, working on caecilian amphibians, showed significant and moderate to good correlations (Spearman’s $\rho = 0.36–0.66$ and Pearson $r = 0.38–0.65$, for comparisons of Euclidean distance matrices from the different methods) between landmark-based morphometrics and discrete character methods, confirming earlier results on echinoderms by Villier & Eble (2004). On the other hand, Mongiardino Koch et al. (2017), in their study of scorpion disparity, found significant differences between discrete character and traditional morphometric methods, and they showed that these differences were greater than expected from the evolution of random data on a phylogeny. They also simulated discrete character data and compared this to simulated traditional morphometric data (not landmark coordinates).

As expected, the two shape-based methods (outlines and landmarks) show the greatest similarity, both in terms of the relative areas of morphospace occupation, the overlaps in morphospace, the total disparity measured by SOV, and in correlation tests. The similarities between the outline and landmark measures are readily understood when shape transformations are compared using both methods and both sample sizes for the first three principal components (Fig. 4). In both cases, the PCA is capturing similar aspects of variation: jaw height, relative attenuation along the jaw length and jaw curvature. The
FIG. 6. Patterns of morphospace occupation and disparity (sum of variances, SOV) of Coelurosauria, for the full sample, grouped according to taxonomy, measured using three metrics. The bivariate morphospaces and disparity plots were created using: A, D, geometric outlines; B, E, geometric landmarks; C, F, discrete jaw and dental characters. Legend: triangles, Tyrannosauroidea; circles, Maniraptoriformes; stripes (D–F), disparity for the Ornithomimosauria, Oviraptoridae, Dromaeosauridae and Avialae. In the morphospace plots (A–C) open circles denote coelurosaurians that are not classified as Tyrannosauroidea or Maniraptoriformes, these taxa are listed in Schaeffer et al. (2019, tables S4, S5). Dinosaur silhouettes from http://phylopic.org: Scott Hartman, Nobu Tamura, Brad McFeeters, Emily Willoughby and Matt Martyniuk (CC BY 3.0). Colour online.
FIG. 7. Patterns of morphospace occupation and disparity (sum of variances, SOV) of Coelurosauria, for the full sample, grouped according to diet, measured using three metrics. The bivariate morphospaces and disparity plots were created using: A, D, geometric outlines; B, E, geometric landmarks; C, F, discrete jaw and dental characters. Legend: circles, small carnivores; dark circles, large carnivores; triangles, omnivores and herbivores; open circles (C), coelurosaurians that could not be confidently assigned a dietary grouping, these taxa are listed in Schaeffer et al. (2019, tables S4, S5). Dinosaur silhouettes from http://phylop.org: Scott Hartman, Brad McFeeters and Emily Willoughby (CC BY 3.0). Colour online.
differences between the two shape analyses are because PC2 and PC3 variously capture the concavity of the jaw outline.

The discrepancy between PC2 in the shape analyses may be driven by our sampling of landmarks. In the landmark geometric approach, two fixed landmarks were positioned on the anterior part of jaw, at the dorsal and ventral angles of the symphysis, and a landmark curve with five points was located between these fixed points. Morphological changes in the relative locations of these anterior landmarks were dominant factors loading on PC2 in the landmark analyses and resulted in the dorsoventral curvature being expressed on PC3, instead of PC2 as in the outline analyses (Fig. 4). It could therefore be argued that this region of the jaw was oversampled, while the outlines more effectively sampled the jaws edges with densely sampled, but evenly distributed, points. Landmarking procedures are an important part of shape analyses (Mitteroecker & Gunz 2009; Cardini 2016; Watanabe 2018) and our results highlight how the selection of methods and landmarks could impact the interpretations of shape evolution based on the major axes of variation. Although, it is important to note that the SOV disparity measured from all PC axes (total shape variation) gave consistent patterns in the outline and landmark methods and the correlation tests based on all axes show strong and significant correlations.

Perhaps our application of landmarks tended to increase the similarity of the outlines and landmark results. By adopting multiple richly sampled semi-landmark curves, we effectively outlined each mandible with 50 points, which gives nearly as much detail as the outline approach (Figs 1, 4). Nonetheless, we felt it was inappropriate here to use a reduced number of fixed homologous landmarks, because there were only a limited number of confidently identifiable homologous points and introducing semi-landmark curves more accurately captures additional information on the overall shape of the mandible. Ideally, it would have been interesting to incorporate landmarks measuring the size and location of the functionally significant external mandibular fenestra, but this feature is absent in some Mesozoic birds and therefore could not be measured on all samples.

The SOV results all suggest that Maniraptoriformes are more disparate than Tyrannosauroidea (Figs 2, 6). The relatively low disparity of Tyrannosauroidea could reflect the lower diversity of this clade in comparison to the Maniraptoriformes, but we argue this is not a problem of experimental design. First, we performed additional analyses with rarefaction to standardize sample sizes for SOV disparity comparisons, and the results were consistent with un rarefied analyses (Scha effe r et al. 2019, table S9). Second, we included all possible specimens (Fig. 6), and so the differences reflect reality, namely the fact it is a smaller clade and with more uniform anatomy and adaptation than Maniraptoriformes. Finally, we used Brusatte et al. (2014) as the data source for discrete cladistic characters, and those authors included the maximum possible sample of tyrannosaurid taxa compared to the other groups (such as Avialae); this does have the effect of diminishing the difference between Maniraptoriformes and Tyrannosauroidea in morphospace occupation (Figs 2C, 6C).

Some authors have advocated caution in using discrete character cladistic data matrices as a source of data for disparity studies (Anderson & Friedman 2012; Benson 2018). These authors note several advantages of such data sets: they are readily available, they typically document broad anatomical coverage, they enable comparisons between taxa of very different form, ancestral taxa can be reconstructed, and they can be scored for fragmentary material. On the other hand, the use of cladistic data matrices entails some problems: it is unclear how to relate anatomical and functional variation both for individual characters or for overall morphologies; such data matrices may reflect phylogenetic signal rather than any aspect of ecomorphology; they may concentrate on obscure anatomical details of, say, the braincase or maxilla orientation, which may have limited functional or evolutionary importance; other significant characters are ignored because they are either autapomorphies or record shape but not phylogenetic signal. Benson (2018) gives examples of published data matrices that may give apparently high rates for certain clades, such as birds, but it cannot be said whether the evolutionary rates are truly high or simply reflect excessive research interest and coding of many small-scale anatomical characters. Gerber (2019) highlights problems of excessive missing data and the meaning of the axes in cases where cladistic characters are used, but they and others (Lloyd 2016; Hopkins & Gerber 2017) support the use of discrete characters to document disparity when appropriate methods are used to minimize bias and understand the data. Equally, of course, outline and landmark studies may be capturing aspects of shape that have no evolutionary or ecological significance at all or might be simply size-related (Gould 1966).

Anderson & Friedman (2012) recommend that analysts devise metrics that have functional significance, such as ratios of relative lengths of portions of the jaw or limbs. Another productive approach could be to use subsets of discrete character data sets that have hypothesized ecomorphological or evolutionary significance (Gerber 2019; Stubbs et al. 2019), such as the appendicular skeleton in the fish–tetrapod transition (Ruta & Wills 2016), the display crests and feeding apparatus in hadrosaurid dinosaurs (Stubbs et al. 2019) and this contribution. Alternatively, researchers could construct or combine new matrices from multiple sources for disparity studies that...
focus on specific parts of anatomy, such as the teeth (Strickson et al. 2016) or teeth and jaw (Nordén et al. 2018).

Ecomorphology

Several ecomorphological features can be seen in our results. First, it is very noticeable that oviraptorids dominate all morphospace analyses (Figs 2, 6). Their bizarre mandible shape dominates PC1 and separates them from the general cluster of taxa. Oviraptorids were also identified as an aberrant group in the studies of skull shape by Brusatte et al. (2012) and Foth & Rauhut (2013). Second, the maniraptoriforms have an overall higher disparity than the tyrannosaurids. This reflects the high disparity of the oviraptorids, but also the fact that the clade includes birds and their ancestors, which were mainly carnivores and insectivores, as well as omnivores and herbivores, some without teeth. These maniraptoriforms encompassed a substantially greater amount of total phylogenetic branch duration than tyrannosaurids, giving them more opportunity to accumulate ecological diversity.

It is interesting that the omnivore–herbivore category, even though sparsely sampled, occupies a great area of morphospace according to all methods (Figs 3, 7) and this, as noted earlier, may simply reflect the fact that this is an ill-defined or waste-basket dietary category. The relative amount of morphospace is especially pronounced with the shape methods (cf. Figs 3D–E, 7D–E). This too reflects the high disparity of oviraptorids, and the generally astonishing variety of shapes of mandibles in that clade, the ornithomimids and others. The differences are less pronounced in the discrete character analyses (Figs 3F, 7F) perhaps because relatively few cladistic-style characters are used to describe ecomorphological differences in the mandible.

CONCLUSION

Our results suggest that all methods of measuring morphological disparity give comparable results, and this is not solely the result of phylogeny. In our case, the two shape-based methods (outlines, landmarks) gave very similar results as they are both recording the same features and rendering them on the dominant multivariate axes in the same ways. We did note some subtle differences in recording of jaw attenuation and bending between the methods, and we highlight the importance of exploring shape changes along axes beyond those shown in biplots of PC1 and PC2 morphospace. By comparing different sampling regimes our comparisons show that a reduced sample of taxa recover the same major axes of shape variation and a full sample with more taxa largely saturates morphospace without considerably expanding it. Our key findings, that the disparity of the major clades and dietary categories were comparable among all three methods, matches earlier findings from other such comparisons of disparity data sets for different taxa. But, as we noted, each method gives subtly different degrees of separation and overlap in disparity between different clades, and so we recommend that future studies should use multiple approaches when assessing disparity as each has its advantages, and each data type reflects different aspects of morphology in relation to function and evolution.

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DATA ARCHIVING STATEMENT

Data for this study are available in the Dryad Digital Repository: https://doi.org/10.5061/dryad.70rk2xf

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