Are endocasts good proxies for brain size and shape in archosaurs throughout ontogeny?

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Abstract

Cranial endocasts, or the internal molds of the braincase, are a crucial correlate for investigating the neuroanatomy of extinct vertebrates and tracking brain evolution through deep time. Nevertheless, the validity of such studies pivots on the reliability of endocasts as a proxy for brain morphology. Here, we employ micro-computed tomography imaging, including diffusible iodine-based contrast-enhanced CT, and a three-dimensional geometric morphometric framework to examine both size and shape differences between brains and endocasts of two exemplar archosaur taxa – the American alligator (Alligator mississippiensis) and the domestic chicken (Gallus gallus). With ontogenetic sampling, we quantitatively evaluate how endocasts differ from brains and whether this deviation changes during development. We find strong size and shape correlations between brains and endocasts, divergent ontogenetic trends in the brain-to-endocast correspondence between alligators and chickens, and a comparable magnitude between brain–endocast shape differences and intraspecific neuroanatomical variation. The results have important implications for paleoneurological studies in archosaurs. Notably, we demonstrate that the pattern of endocranial shape variation closely reflects brain shape variation. Therefore, analyses of endocranial morphology are unlikely to generate spurious conclusions about large-scale trends in brain size and shape. To mitigate any artifacts, however, paleoneurological studies should consider the lower brain–endocast correspondence in the hindbrain relative to the forebrain; higher size and shape correspondences in chickens than alligators throughout postnatal ontogeny; artificially ‘pedomorphic’ shape of endocasts relative to their corresponding brains; and potential biases in both size and shape data due to the lack of control for ontogenetic stages in endocranial sampling.

Key words: Alligator; diffusible iodine-based contrast-enhanced computed tomography; Gallus; geometric morphometrics; micro-computed tomography; neuroanatomy.

Introduction

Fossils are indispensable resources for elucidating ancient biotas and evolutionary dynamics through deep time (Gauthier et al. 1988a; Donoghue et al. 1989; Raff, 2007; Lee & Palci, 2015; Rabosky, 2015). Nevertheless, fossils are inherently limited in the biological information that they can provide. For example, the rarity of soft-tissue preservation requires paleontologists to frequently utilize anatomical correlates of preserved hard tissues to infer unpreserved soft-tissue characteristics of extinct taxa (e.g. Witmer, 1995; Wedel & Sanders, 2002; Watanabe et al. 2015). In the field of paleoneurology (Edinger, 1929; Kochetkova, 1978) internal molds of the cranial cavity, called cranial endocasts (hereafter ‘endocasts’), have provided crucial information on the brain morphology of extinct vertebrates (e.g. Jerison, 1963, 1969; Edinger, 1975; Hopson, 1979; Balanoff & Bever, 2017). More recently, the advent of micro-computed tomography (μCT) imaging has propelled the use of high-resolution virtual endocasts to reconstruct the neuroanatomy of fossil and extant taxa (Balanoff et al. 2016;
and references therein). Modern comparative studies have harnessed this technology to infer large-scale trends in brain evolution, including the origins of highly encephalized brains in mammals and birds (e.g. Rowe et al. 2011; Balanoff et al. 2013; Neubauer, 2014).

Despite its capacity to provide valuable information, the validity of any anatomical correlate relies on the degree to which it accurately reflects the soft-tissue structures of interest. Naturally, endocasts overestimate brain sizes due to the intermediary space between the brain and the internal surface of the skull that contains meninges, arteries, venous sinuses, cerebrospinal fluid, and the roots of cranial nerves. Seminal work based on volumetric measurements of brains and endocasts has shown that size differences between them are negligible in extant mammals (Haight & Nelson, 1987; de Miguel & Henneberg, 1999) and birds (Jerison, 1973; Zusi, 1993; Iwanuki & Nelson, 2002) because their enlarged brains occupy nearly the entire endocranial space. The neuroanatomical literature has historically referenced these volumetric analyses to justify the use of endocasts in studies of mammals, birds, and closely related extinct clades (e.g. Rowe et al. 2011; Balanoff et al. 2013). In contrast, the endocasts of most other vertebrate clades have been considered to be poor representations of brain size because the brains do not closely occupy the endocranial space (Jerison, 1969; Hopson, 1977, 1979; Kochetkova, 1978).

Although endocasts may be a good proxy for brain volume in certain taxa, quantitative assessments of brain–endocast correspondence in shape have been limited. The development of three-dimensional (3D) semi-landmark techniques (Gunz et al. 2005; Gunz & Mitteroecker, 2013) has facilitated the use of landmark-based geometric morphometric (GM) methods to characterize the relatively featureless surfaces of endocasts (Gómez-Robles et al. 2018; Pareira-Pedro & Bruner, 2018). This growing use of GM approaches in paleoneurological research prompts an examination into the degree to which endocranial shape reflects true brain morphology. Such investigations will demonstrate which brain regions are accurately or poorly approximated by the corresponding areas on endocasts.

Furthermore, both size and shape correspondences between brains and endocasts likely change throughout ontogeny. For instance, the volume of cerebrospinal fluid in the endocranial cavity increases with age in humans (Wan-uchi et al. 2002; Sherwood et al. 2011), and differential, non-linear development of the brain and the skull occurs across hominins (Neubauer et al. 2010; Bruner et al. 2015). In crocodylians, Jirak & Janacek (2017) reported changes to the brain–endocast volumetric correspondence throughout ontogeny, with closer correspondence in younger individuals. These observations imply that the inferential power of endocasts differs across not only taxa but also ontogenetic stages. However, the ontogenetic data analyzed by Jirak & Janacek (2017) covered a mixture of multiple species and sexes, preventing a systematic analysis of brain–endocast differences.

Here, we examine brain–endocast correspondence in size and shape during ontogeny in two exemplar extant archosaurs – the American alligator (Alligator mississippiensis) and the domestic chicken (Gallus gallus). Archosauria sensu Gauthier et al. (1988b) (e.g. crocodylians, birds, non-avian dinosaurs) is of interest to neurobiologists because birds possess highly encephalized brains that evolved independently from those of mammalian groups, including primates (Jerison, 1973; Northcutt & Kaas, 1995; Nieuwenhuys et al. 1998). We employ a suite of modern techniques, including standard µCT imaging, as well as diffusible iodine-based contrast-enhanced CT (diceCT) imaging (Metscher, 2009a,b; Gignac & Kley, 2014; Gignac et al. 2016), to create high-resolution endocasts and brain reconstructions, respectively. For the first time in archosaurs, we employ a high-dimensional 3D GM approach to characterize the shape of brains and endocasts and their major functional subdivisions. This dataset was then subjected to statistical methods to assess whether (1) the brain and endocranial shapes are distinct in alligators and chickens, (2) the brain–endocast deviation changes throughout ontogeny, and (3) the magnitude of this deviation could overcome real signals of intra- and interspecific variation in archosaurian neuroanatomical shape. In light of the results, we formulate important considerations for future comparative neuroanatomical studies on archosaurs.

**Materials and methods**

**Specimens**

We obtained postnatal specimens of *A. mississippiensis* from the Rockefeller Wildlife Refuge (Grand Chenier, LA, USA): four ‘perinatal’ (1 year old) individuals, five ‘yearlings’ (1-2 years old), and two ‘juvenile’ specimens (2-3 years old) (*n* = 11; Table 1). Specimens were euthanized with an overdose of 150 mg kg⁻¹ body mass solution of sodium pentobarbital (Fatal-Plus™, Vortech Pharmaceuticals, Dearborn, MI, USA) injected into the intraperitoneal space. This protocol was approved by the Stony Brook University Institutional Animal Care and Use Committee (IACUC, Protocol #236370-1) and the Oklahoma State University Center for Health Sciences (OSUCHS) IACUC (Protocol #2015-1). Specimens were then decapitated between the third and fourth cervical vertebrae and immediately fixed in 10% neutral-buffered formalin to prevent postmortem decomposition of the brain. To minimize shape distortion from tissue fixation (Weisbecker, 2012), we fixed the specimens in formalin for over 8 weeks before imaging.

The Charles River Laboratory (North Franklin, CT, USA) supplied male *G. gallus* specimens at 1 day, 1 week, 3 weeks, 6 weeks, and over 8 weeks of age (Table 1). Two individuals were sampled for each age group, with the exception of four individuals at one-day and over 8 weeks of age (*n* = 14). These specimens were euthanized at the Charles River Laboratory via cervical dislocation and decapitation, followed immediately by submersion into 10% neutral-buffered formalin solution. After 2 weeks, the fixed specimens
were transported to the American Museum of Natural History (AMNH; New York, NY, USA) wrapped in formalin-saturated gauze. Upon arrival, the specimens were again submerged in formalin for over 8 weeks before imaging to minimize distortion in brain morphology (Weisbecker, 2012). Ontogenetic sampling of alligators and chickens did not encompass equivalent developmental stages except for the trigeminal ganglion in frontal view, which allowed the clearest and most consistent delimitation of the endocast and the ganglion. Segmented endocasts were exported in Polygon file format (PLY) using ‘Precise with simplification’ setting.

DiceCT imaging utilizes Lugol’s iodine (iodine potassium-iodide, I₂KI) as a contrast agent, rendering soft tissues more radio-opaque (Metscher, 2009a,b; Gignac & Kley, 2014; Gignac et al. 2016). We used diceCT to create high-resolution in situ reconstructions of the brain from the same set of specimens used to create endocasts. The size of the specimens informed both the concentration and duration of iodine stains for optimizing the contrast among soft-tissue types (Table 1; also see Gignac et al. 2016). Although iodine staining has been associated with soft-tissue shrinkage (Vickerton et al. 2013; Cox & Faulkes, 2014), our CT images show that the brains are in close proximity to the skulls, suggesting that neither formalin fixation nor iodine staining resulted in substantial soft-tissue distortions (Fig. 1). This result is consistent with a recent assessment of potential shrinkage artifacts in bat brains, where specimens stained shortly after their collection in the field incurred minimal shrinkage effects compared to museum specimens that had been fixed in ethanol prior to staining (Hedrick et al. 2018).

The specimens were submerged in aqueous solutions of Lugol’s iodine for certain periods (Table 1) and regularly agitated on a Vertex-Genie 2 machine (Scientific Industries, Inc., Bohemia, NY, USA) for 60 s every 2–3 days to facilitate the incorporation of iodine into deeper tissue layers. During the staining process, the containers with the specimen and Lugol’s iodine solution were stored in the dark to limit loss of stain potency (Gignac et al. 2016). Processing of CT image stacks followed those for unstained specimens, with two exceptions. First, the image stacks were subjected to the enhanced local contrast (CLAHE) script (Saalfeld, 2010) in ImageJ (FIJI) with the default concentration (% I₂KI, w/v) and processing and volume-rendering CT images using the Phoenix Imaging Facility. We varied the scan parameter values in an effort to optimize the contrast and resolution of the X-ray images (Supporting Information Table S1). The creation of virtual endocasts consisted of scanning formalin-fixed heads of Alligator and Gallus, then processing and volume-rendering CT images using the Phoenix DATOS 2 reconstruction software v2.3.2 (GE Sensing & Inspection Technologies, Hürth, Germany). For larger specimens requiring multiple scans, separate image stacks were fused using the ‘3D Stitching’ function in Imaris (FIJI) v1.49u (Schindelin et al. 2012). In VGStudio MAX v2.2 (Volume Graphics, Heidelberg, Germany), we imported full X-ray image stacks of each specimen and digitally segmented the endocranial cavity following the protocol outlined by Balanoff et al. (2016). Any impressions of the cranial nerves were removed digitally from segmented regions in transverse view, except for the trigeminal ganglion in frontal view, which allowed the clearest and most consistent delimitation of the endocast and the ganglion.

Table 1 List of sampled specimens with information on age, sex, staining protocol, and neuroanatomical measurements.

| Taxon | Age       | Sex | Lugol’s iodine concentration (% I₂KI, w/v) | Stain duration (days) | Brain volume (mm³) | Endocranial volume (mm³) |
|-------|-----------|-----|-------------------------------------------|-----------------------|-------------------|-------------------------|
| Alligator | 0–1 year  | ♀   | 11.25                                     | 28                    | 520.37            | 742.42                  |
|        | 0–1 year  | ♂   | 11.25                                     | 14                    | 657.90            | 760.89                  |
|        | 0–1 year  | ♂   | 7.50                                      | 14                    | 783.90            | 814.49                  |
|        | 0–1 year  | ♂   | 7.50                                      | 14                    | 793.82            | 964.30                  |
|        | 1–2 years | ♀   | 11.25                                     | 21                    | 1157.20           | 1393.67                 |
|        | 1–2 years | ♂   | 11.25                                     | 21                    | 1331.76           | 2507.89                 |
|        | 1–2 years | ♂   | 11.25                                     | 14                    | 1544.82           | 1767.80                 |
|        | 2–3 years | ♂   | 11.25                                     | 14                    | 1649.50           | 2147.62                 |
|        | 2–3 years | ♂   | 11.25                                     | 36                    | 2164.22           | 4154.08                 |
|        | 1 day     | ♂   | 5.00                                      | 14                    | 541.66            | 901.88                  |
|        | 1 day     | ♂   | 3.00                                      | 14                    | 748.15            | 965.36                  |
|        | 1 day     | ♂   | 3.00                                      | 14                    | 796.69            | 939.77                  |
|        | 1 day     | ♂   | 3.00                                      | 14                    | 830.41            | 1016.83                 |
|        | 1 week    | ♂   | 5.00                                      | 14                    | 1007.63           | 1206.21                 |
|        | 1 week    | ♂   | 5.00                                      | 14                    | 1034.74           | 1285.18                 |
|        | 3 weeks   | ♂   | 7.50                                      | 14                    | 1643.88           | 1958.14                 |
|        | 6 weeks   | ♂   | 10.00                                     | 14                    | 1648.04           | 1891.17                 |
|        | 6 weeks   | ♂   | 10.00                                     | 14                    | 2207.13           | 2512.06                 |
|        | > 8 weeks | ♂   | 10.00                                     | 22                    | 2852.36           | 3257.38                 |
|        | > 8 weeks | ♂   | 10.00                                     | 21                    | 2856.40           | 3152.19                 |
|        | > 8 weeks | ♂   | 10.00                                     | 21                    | 3057.25           | 3277.91                 |
| Gallus | 1 day     | ♀   | 3.00                                      | 14                    | 1034.74           | 1285.18                 |
|        | 1 day     | ♀   | 3.00                                      | 14                    | 1007.63           | 1206.21                 |
|        | 3 weeks   | ♀   | 7.50                                      | 14                    | 1643.88           | 1958.14                 |
|        | 6 weeks   | ♀   | 10.00                                     | 14                    | 1648.04           | 1891.17                 |
|        | 6 weeks   | ♀   | 10.00                                     | 14                    | 2207.13           | 2512.06                 |
|        | > 8 weeks | ♀   | 10.00                                     | 22                    | 2852.36           | 3257.38                 |
|        | > 8 weeks | ♀   | 10.00                                     | 21                    | 2856.40           | 3152.19                 |
|        | > 8 weeks | ♀   | 10.00                                     | 21                    | 3057.25           | 3277.91                 |

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Fig. 1 Selected transverse (left), frontal (middle), and sagittal (right) μCT slices through the heads of perinatal Alligator (A), 2- to 3-year-old Alligator (B), 1-day-old Gallus (C), > 8-week-old Gallus (D), illustrating the minimal shrinkage artifact from staining neural tissue with high concentrations of Lugol’s iodine. Scale: 5 mm (A,C), 10 mm (B,D).
parameters to increase local grayscale contrast. Secondly, we used the Adaptive Gauss filter in VGSTUDIO MAX with the default parameters to sharpen edges, thus improving edge recognition during digital segmentation. The ventricles were left unfilled in the brain reconstructions for more accurate calculation of brain tissue volume.

**Morphological data**

We calculated the volume (mm$^3$) of endocasts and brain reconstructions in MESHLAB v2016.12 (Cignoni et al. 2008; Table 1). To characterize neuroanatomical shape, we used a 3D landmark-based GM approach. The collection of coordinate data from endocasts is difficult due to the dearth of discrete anatomical landmarks on their surfaces (Neubauer, 2014). For example, landmark configurations in previous studies on avian brain shape included one or two landmarks within major functional subdivisions (e.g. cerebrum, optic lobes, cerebellum), preventing a robust characterization of morphological variation within these structures (e.g. Kawabe et al. 2013, 2015; Marugán-Lobón et al. 2016). An automated collection and alignment of coordinate data (Boyer et al. 2011, 2015) and the landmark-free iterative closest point algorithm (Pomidor et al. 2016) are not suitable for this study because the endocasts in both species include surfaces that are interpolations and do not directly represent a structural surface (e.g. uns ossified regions of the laterosphenoid in immature specimens).

In this study, we established a landmark configuration combining discrete landmarks with semi-landmarks on curves and surfaces using LANDMARK EDITOR v3.6 (Wiley et al. 2005). The ‘patch’ tool in LANDMARK EDITOR allows the placement of discrete, consistently identifiable landmarks that define the boundaries of major functional brain divisions (i.e. left and right cerebra, left and right optic lobes, cerebellum, medulla) with a specified density of semi-landmarks sampled within these subdivisions (Table 2). The bilateral landmark data comprised 24 discrete landmarks, 87 curve semi-landmarks that define the boundaries of major functional divisions (i.e. cerebrum, optic lobes, cerebellum, and medulla), and 114 surface semi-landmarks that characterize the shape within each division. When placing surface semi-landmarks on the reconstructions of the cerebella from diceCT data, we visually confirmed that the landmarks were placed on gyri and not within the sulci.

We used the GEOMORPH R package v3.0.1 (Adams & O’tariola-Castillo, 2013) to perform a generalized Procrustes alignment on the combined Alligator and Gallus data (Gower, 1975; Rohlf & Slice, 1990), with sliding semi-landmarks minimizing total bending energy (Gunz et al. 2005; Gunz & Mitteroecker, 2013). Procrustes distances, or the sums of squared differences between corresponding landmarks and semi-landmarks, were calculated to measure shape differences among specimens. In addition to shape, we recorded the centroid sizes of the endocasts and brains from the coordinate data. After alignment, the landmarks and semi-landmarks on the left side were removed to exclude redundancy in morphological information while avoiding artifacts from aligning one-sided data of bilaterally symmetric structures (Cardini, 2016a,b). The resulting unilateral shape data comprised 16 landmarks, 49 curve semi-landmarks, and 59 surface semi-landmarks (Fig. 2, Table 2, Supporting Information Data S1). We also generated form data (combined shape and size data) by multiplying the shape data with the corresponding centroid size for each specimen. The form difference between brains and endocasts is a metric concomitant with physical distances between each corresponding landmark and semi-landmark. We computed digitization error by repeatedly collecting landmark data from a 1-day-old chicken (10 replications), which accounted for 4.14% of the total shape variation and was considered to be negligible.

**Analysis**

All statistical analyses were performed in R v3.2.4 (R Core Development Team, 2018). First, the volumetric correspondence between

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**Table 2** Landmark scheme used in this study. Each major neuroanatomical region was characterized by discrete landmarks, curve semi-landmarks that define the regional boundaries, and surface semi-landmarks to characterize the shape within each region. The numbers in parentheses denote number of median and right semi-landmarks analyzed in the study after performing generalized Procrustes alignment on bilateral data.

| Region          | No. curve semi-landmarks | No. surface semi-landmarks | Discrete landmarks defining each region                                                                 |
|-----------------|--------------------------|----------------------------|----------------------------------------------------------------------------------------------------------|
| Cerebrum        | 24 (12)                  | 78 (39)                    | Anteriormost median point of the cerebrum on dorsal side.                                                |
|                 |                          |                            | Posteromedial point of the cerebrum on dorsal side.                                                      |
|                 |                          |                            | Dorsalmost junction point of cerebrum and optic side.                                                   |
|                 |                          |                            | Ventralmost junction point of cerebrum and optic side.                                                  |
| Optic lobe      | 28 (14)                  | 24 (12)                    | Anteriormost median point of the cerebrum on dorsal side.                                               |
|                 |                          |                            | Posteromedial point of the cerebrum and optic side.                                                     |
|                 |                          |                            | Dorsalmost junction point of cerebrum and optic lobe.                                                   |
|                 |                          |                            | Ventralmost junction point of cerebrum and optic lobe.                                                  |
|                 |                          |                            | Junction point of optic lobe, cerebellum, and medulla.                                                  |
| Cerebellum      | 18 (12)                  | 8 (4)                      | Anteriormost median point of the cerebellum on dorsal side.                                             |
|                 |                          |                            | Anterolateral point of the cerebellum on the dorsal side.                                               |
|                 |                          |                            | Posteriormost median point of the cerebellum on dorsal side.                                            |
| Medulla         | 17 (11)                  | 8 (4)                      | Anteriormost median point adjacent to midbrain on ventral side.                                         |
|                 |                          |                            | Junction point of optic lobe and medulla.                                                               |
|                 |                          |                            | Posteriolateral point of medulla.                                                                      |
|                 |                          |                            | Posteriormost median point of medulla.                                                                |

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brains and endocasts was evaluated with a least-squares regression analysis on brain and endocranial sizes. We then calculated the ratio of brain to endocranial volumes. These ratios were regressed onto log-transformed brain centroid size (logCS) to identify ontogenetic trends in brain–endocast size differences in *Alligator* and *Gallus*. We employed a series of statistical analyses to investigate brain–endocast shape differences. To visualize the pattern of neuroanatomical variation, we created a morphospace of brains and endocasts using the scores along the first two principal components of shape variation. To test whether endocasts and brains differ significantly in shape, multivariate analysis of variance (MANOVA) was performed on endocranial and brain shapes of chickens and alligators separately, as well as on the combined data, using the procD.lm function in the GEOMORPH package. Localized form and shape differences were visualized by observing the direction and magnitude of changes in landmark positions. The Procrustes distances between corresponding endocasts and brains were plotted against logCS of the brain to examine whether shape differences between brains and endocasts change predictably throughout ontogeny. Least-squares regression analysis was used to detect the presence of a trend. Next, we used analysis of variance (ANOVA) to evaluate whether the magnitude of brain–endocast shape distances is different from (1) intraspecific variation within *Alligator* and *Gallus* separately; (2) interspecific variation between these two taxa; and (3) interspecific differences in endocranial shape among extant birds after conducting a generalized Procrustes alignment on a pooled coordinate dataset. For the latter comparison, we collected shape data from endocasts sampled in previously published studies (Balanoff et al. 2013). In association with these analyses, we constructed box-and-whisker plots illustrating the extent of overlap between brain–endocast differences across these hierarchical levels of neuroanatomical shape variation.

**Results**

**Brain–endocast volume**

The brain occupies 52–99% and 60–93% of endocranial volume in *Alligator* and *Gallus*, respectively. Despite these volumetric differences, endocranial volume correlates strongly with brain volume in both *Alligator* and *Gallus* (Fig. 3A; $R^2 > 0.92$, $P < 0.001$). As expected, the regression line for *Gallus* shows a steeper slope than that for *Alligator*, indicating that chickens have proportionately larger brains within the endocranial space. Regressing brain-to-endocast volume onto logCS of the brain indicates divergent trends in the two taxa (Fig. 3B). *Alligator* shows greater brain–endocast size deviation in larger specimens ($R^2 = 0.395$; $P = 0.038$), implying that the brain occupies a smaller proportion of the endocranial cavity in more mature individuals. Conversely, we find that the proportional brain size within the endocranial cavity increases throughout ontogeny in chickens ($R^2 = 0.658$; $P < 0.001$).

**Brain–endocast form**

Differences in form at each landmark illustrate localized morphological differences between brains and endocasts that reflect true physical distances (Fig. 4). Across taxa and ontogenetic stages, the endocasts generally exhibit less dorsoventral convexity in the cerebrum, greater anterior and ventral extent of the optic lobe region, less posteroventrally flexed cerebellum, and less dorsoventral convexity in the medulla. In *Alligator* the magnitude of these form differences is generally smaller in larger, more mature individuals, especially in the cerebrum (Fig. 4A). A regression analysis supports the observation that larger *Alligator* specimens exhibit smaller form differences (Fig. 3C; $R^2 = 0.478$; $P = 0.019$). In *Gallus*, form differences are concentrated in the dorsoventral extent of anterior and posterior margins of the cerebrum, the lateral extent of the cerebrum and optic lobe, and the dorsoventral extent of both the cerebellum and medulla (Fig. 4B). In contrast to *Alligator*, the magnitude of form differences in chickens does not seem to change throughout postnatal ontogeny (Fig. 3C; $R^2 < 0.001$; $P = 0.943$).

**Brain–endocast shape**

A morphospace constructed from the first two principal components (PC) illustrates the taxonomic and ontogenetic trends in neuroanatomical shape (Fig. 5A). PC1 accounts for 59.5% of the total shape variation and separates the neuroanatomical shape between *Alligator* and *Gallus*. It is primarily associated with the (1) lateral expansion of the cerebrum; (2) sphericity, relative position, and proportional size of the optic lobe; (3) dorsoventral flexion of the cerebellum; and (4) relative anteroposterior length of the medulla. PC2 accounts for 10.8% of the total shape variation and corresponds to the (1) degree of dorsoventral flexion in the entire brain and endocast; (2) relative size of the cerebrum; (3) position of the optic lobes; and (4) dorsoventral convexity of the cerebellum and medulla along the longitudinal axis. PC2 aligns with ontogenetic changes in brains and endocasts, where the ontogenetic trajectories of *Alligator* and *Gallus* are parallel to each other. Brain shape is collectively shifted to occupy areas further along the ontogenetic trajectory than the corresponding endocasts. Endocasts, therefore, exhibit an artifically ‘pedomorphic’ shape relative to brains of the same age.

In both *Alligator* and *Gallus*, the endocasts generally exhibit an anteroposteriorly restricted medulla and less dorsoventral flexion in the hindbrain (Fig. 5B,C). In *Alligator* the impression of the optic lobe on the mean endocranial shape extends further ventrally than the position of the optic lobe in the mean brain shape (Fig. 5B). Although similar shape divergences remain in the cerebellum and medulla, the divergence in the cerebrum decreases in more mature *Alligator* specimens (Fig. 5D). In *Gallus*, the endocasts exhibit greater lateral extent in the cerebrum and optic lobes that are more limited in posterolateral breadth (Fig. 5C). The overall shape divergence seems to decrease in more mature *Gallus* specimens (Fig. 5E). The MANOVA
corroborates that brains and endocasts are significantly different in their mean shapes in *Alligator* ($R^2 = 0.19; P < 0.001$) and *Gallus* ($R^2 = 0.15; P < 0.001$). The plot of Procrustes distances against logCS of brains and the regression lines (Fig. 3D) suggest that brain–endocast shape differences may gradually decrease throughout ontogeny, but MANOVA fails to reject the absence of a trend in *Alligator* ($R^2 = 0.335; P = 0.261$) and *Gallus* ($R^2 = 0.169; P = 0.144$).

Comparisons with intra- and interspecific variation

We calculated the pairwise Procrustes distances within *Alligator* and *Gallus* to represent intraspecific neuroanatomical variation for these taxa, as well as between the brains of *Alligator* and *Gallus* specimens to measure the interspecific variation between these two taxa (Table 3). Even with more restricted ontogenetic sampling, the magnitude of brain–endocast shape differences is greater in *Alligator* than in *Gallus* (Fig. 6), as confirmed by ANOVA ($R^2 = 0.303; P = 0.004$). When compared to the intraspecific variation in brain shape within these taxa, the brain–endocast shape differences are comparable (Fig. 6; *Alligator*: $R^2 = 0.026; P = 0.103$; *Gallus*: $R^2 = 0.029; P = 0.083$). The brain–endocast shape differences are smaller than the interspecific differences between *Alligator* and *Gallus* ($R^2 = 0.786; P < 0.001$) and are generally less than the magnitude of interspecific endocranial variation in crown-group birds ($R^2 = 0.117$; Fig. 2).

![3D landmark configuration used in this study on brains (left) and endocasts (right) on (A) a juvenile Alligator and (B) adult Gallus. Red, yellow, and blue points denote discrete, curve, and surface landmarks and semi-landmarks, respectively. Images not in scale.](image)

**Table 3** Mean and range of brain–endocast shape differences (Procrustes distance) at multiple levels of variation. To permit comparison of Procrustes distances, the values are based on alignment of pooled data comprising *Alligator*, *Gallus*, and interspecific sampling of extant birds. These values correspond to Fig. 6.

| Variation type                               | Mean   | Range          |
|----------------------------------------------|--------|----------------|
| Brain–endocast (*Alligator*)                 | 0.142  | 0.097–0.179    |
| Brain–endocast (*Gallus*)                    | 0.098  | 0.009–0.137    |
| Brain–endocast (*Alligator and Gallus*)     | 0.118  | 0.009–0.179    |
| Intraspecific variation (*Alligator*)        | 0.122  | 0.071–0.193    |
| Intraspecific variation (*Gallus*)           | 0.113  | 0.047–0.160    |
| Interspecific variation (*Alligator, Gallus*)| 0.245  | 0.200–0.304    |
| Interspecific variation (Neornithes)         | 0.165  | 0.081–0.309    |
Although they partially overlap in the value of Procrustes distances (Fig. 6), $P < 0.001$, our study corroborates and extends previous reports on brain–endocast congruence in archosaurs utilizing modern techniques on postnatal ontogenetic series of Alligator and Gallus. In Alligator the brain displays negative allometry relative to endocranial size throughout ontogeny (Fig. 3B), corroborating previous studies (Hopson, 1979; Rogers, 1999; Hurlburt & Waldorf, 2002; Hurlburt et al. 2013). We also find that perinatal alligators exhibit high brain–endocast correspondences in volume (> 90%). These values are consistent with Crocodylus acutus at Stage 28 embryonic stage showing 97.5% brain-to-endocast correspondence (Jirak & Janacek, 2017), and are much greater than previously reported for A. mississippiensis (i.e. 67%; Hurlburt & Waldorf, 2002). The brain occupies nearly half of the endocranial volume in the largest Alligator specimens sampled in this study (Fig. 3B), but volumetric correspondence is expected to decrease further in more mature individuals.
because previous studies have shown that the brain occupies 32 and 29% of the endocranial space in much larger specimens of *A. mississippiensis* (Hurlburt & Waldorf, 2002; Hurlburt et al. 2013) and *Crocodylus niloticus* (Jirak & Janacek, 2017), respectively. Although we did not sample somatically mature specimens in *Alligator*, our study fills a crucial gap in ontogenetic sampling that provides an evidence of consistent reduction in brain–endocast size correspondence during the first few years of life.

In contrast to *Alligator*, brain size converges towards endocranial size as growth proceeds in *Gallus* (Fig. 3B). The high proportional brain size in older individuals supports previous studies showing that the brain occupies nearly the entire endocranial space (> 90%) in somatically mature birds (Jerison, 1973; Zusi, 1993; Iwaniuk & Nelson, 2002). We also find that the brain occupies less than 80% of the endocranial cavity in neonatal chickens. Taken together, *Alligator* and *Gallus* exhibit divergent allometric trends in

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**Fig. 4** Form differences between brains and endocasts of (A) perinatal and juvenile *Alligator* and (B) 1-day-old and > 8-week-old *Gallus*. Diagrams indicate direction and magnitude of changes in landmark positions from brains to endocasts. The magnitude of vectors reflects true distance.

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brain size relative to endocranial cavity. The precise mechanism for these ontogenetic trends is yet unclear. In the Nile crocodile (C. niloticus), the central nervous system continues to grow indeterminately with body size, where the brain grows more slowly than the spinal cord relative to body size (Ngwenya et al. 2013). This combined effect of indeterminate growth and the brain-to-body allometric relationship in crocodylians is consistent with the observed reduction in proportional brain size through ontogeny. Like other birds, Gallus undergoes an abbreviated period of somatic growth that is characterized by a derived allometric relationship between brain and body size (e.g. Jerison, 1973; Tsuboi et al. 2018). Therefore, these contrasting developmental strategies likely underlie these opposing trends in brain-endocast correspondence. Fabbri et al. (2017), for instance, showed that the skull roof closely tracks the shape of adjacent regions on the forebrain and midbrain during early ontogeny that become decoupled later in development. Given this observation, Gallus, and more broadly birds, may maintain high brain-endocast correspondence due to their truncated period of somatic growth. Clarifying the molecular and functional mechanisms that explain the varying

Fig. 5 Shape variation in endocranial and brain shape data of Alligator (green) and Gallus (red). (A) Morphospace constructed from PC1 and PC2 of shape. Circles and triangles denote brains and endocasts, respectively. The size of data points corresponds to non-log-transformed centroid size. Direction and magnitude of differences in landmark positions from mean brain to mean endocranial shape in Alligator (B) and Gallus (C). Brain-to-endocast shape differences in perinatal and more mature specimens in (D) Alligator and (E) Gallus. The magnitude of vectors reflects true shape distance.
degrees of association between skull and brain development will be an important line of research in archosaur neurobiology (e.g. Marcucio et al. 2005, 2011; Young et al. 2010; Hu et al. 2015).

Form and shape correspondence

In both Alligator and Gallus, brain–endocast shape differences are concentrated in the dorsoventral convexity of the cerebellum and medulla, whereas the cerebrum (and optic lobes in Gallus) shows higher correspondences. These non-uniform deviations in shape across brain regions mirror that in form, indicating that areas with greater physical distances between the brain and endocranial surfaces also represent the regions with greater deviations in shape. The areas of relatively high deviations and close correspondences are consistent with previous studies. In crocodylians, Hopson (1979) noted that the dorsal longitudinal venous sinus and its divisions occupy a substantial portion of the area around the cerebellum and medulla. Conversely, the venous sinus is relatively thin around the cerebrum, allowing a closer correspondence between the cerebrum and adjacent areas of the braincase (Evans, 2005). This anatomical feature may extend to Gallus, where a thicker dorsal longitudinal venous sinus surrounding the hindbrain contributes to the greater morphological deviations in the cerebellum and medulla (Balanoff & Bever, 2017). In many non-avian dinosaurs, the endocast exhibits a ‘dural peak’ in the cerebellar region (Balanoff & Bever, 2017), which may represent even more size and shape deviations from the actual brain morphology.

Implications for paleoneurology

Our study shows that endocasts are clearly distinct from brains in size, form, and shape, and that the magnitude of this shape difference is comparable to intraspecific variation in brain shape. Although these results seem to suggest that endocasts are poor correlates for brain morphology, the pattern of morphological variation is congruent between brains and endocasts. For instance, endocranial volume is

Fig. 6 Box-and-whisker plots of Procrustes shape distances between corresponding brains and endocasts within Alligator and Gallus (brain-endocast difference), for intraspecific brain shape variation within Alligator and Gallus (brain-shape difference), interspecific brain shape variation between Alligator and Gallus (Alligator-Gallus brain shape difference), and interspecific endocranial shape variation in extant birds (endocranial shape variation in birds). The pooled coordinate data were re-aligned with data from extant birds to place all specimens within a single shape space to allow comparison of variation. The brackets indicate significant differences in Procrustes distances between two levels (*P < 0.05, **P < 0.01, ***P < 0.001).
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endocranial size has been known to reflect brain size in somatically mature birds accurately, the use of raw endocranial volume could overestimate brain size in perinatal specimens by 20% or more.

• Merging endocranial with brain data should be avoided. Although we are not aware of any study that combines morphometric data of endocasts and brains, we discourage such a ‘total evidence’ approach to comparative anatomy. Besides exaggerating form and shape variation in the hindbrain, these shape data would artificially incur a developmental signal, where endocasts exhibit ‘pedomorphic’ shapes relative to their corresponding brains.

• Ontogenetic stage should be controlled for purely interspecific studies whenever possible. For volumetric data, our results caution against mixing multiple species from various ontogenetic stages because the proportional brain volume varies considerably throughout ontogeny. Merging multiple ontogenetic stages would unintentionally include intraspecific variation that would be spuriously interpreted as interspecific variation. The same issue extends to form and shape data, unless an ontogenetic correction factor could be established through equivalent studies on additional archosaur taxa.

• Merging neuroanatomical data from crocodylians and birds should be avoided. Such data will tend to overestimate brain volumes in crocodylians relative to birds, assuming that the results from Alligator and Gallus are applicable to their respective crown groups. As with size, birds have a greater shape correspondence than crocodylians at similar brain sizes. Therefore, endocranial size or shape data from these two groups will distort real variation in brain morphology. To resolve this issue, identifying when particular ontogenetic trends evolved in the archosaur phylogeny is crucial for formulating a clad-specific correction factor for both size and shape (e.g. Kochiyama et al. 2018; for hominins). Endocasts of stem archosaurian clades seem to show closer associations with external brain structures, suggesting that the patterns observed here in Alligator and previously in Crocodylus (Ngwenya et al. 2013) may be a derived feature of crocodylians (Pierce et al. 2017; and references therein). Until we identify the polarity and evolutionary timing of derived ontogenetic trends in neuroanatomical size and shape, we discourage inferring the ontogenetic stage of extinct archosaur lineages based on an estimated proportional size of the brain (e.g. Hurlburt et al. 2013).

Conclusions

Through the use of µCT imaging and diceCT, we introduce a new neuroanatomical dataset comprising size and shape data from endocasts and brains of the same Alligator and
Gallus specimens. By employing a suite of computational methods on 3D GM data, we demonstrate that (1) Alligator and Gallus show discordant ontogenetic trends in volumetric correspondence between brains and endocasts; (2) the brain–endocast shape deviation is greater in Alligator than in Gallus for the ontogenetic stages sampled; (3) brains and endocasts differ significantly in shape, particularly with respect to the dorsoventral flexion of the cerebellum and medulla; and (4) the magnitude of brain–endocast shape difference is comparable to intraspecific variation within these taxa but generally lower than interspecific variation between Alligator and Gallus, as well as among extant birds. While we show that endocasts retain the overall pattern of brain shape variation, we provide several suggestions to mitigate artifacts in neuroanatomical data (see ‘Implications for paleoneurology’). Moving forward, equivalent studies on additional archosaur taxa are necessary to establish clade-specific ontogenetic trends in brain–endocast correspondence, akin to what has been achieved with research on interspecific allometric trends in brain and body size. Such endeavors will be critical for accurate and precise inferences of brain morphology in paleoneurological studies.

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Author contributions

A.W. conceived the study, created the figures, and wrote the manuscript with assistance from all co-authors. A.W., P.M.G., T.L.G., and N.J.K. collected and prepared specimens. A.W. and P.M.G. scanned the specimens and created endocasts and brain reconstructions. A.M.B. provided endocranial reconstructions of modern birds.

Conflict of interest

The authors have no conflict of interest to declare.

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Supporting Information

Additional supporting information may be found online in the Supporting Information section at the end of the article.

Table S1. Parameters used for CT imaging of specimens.

Data S1. Procrustes aligned shape data of brains and endocasts of Alligator and Gallus specimens (plain text file).