A histological survey of avian post-natal skeletal ontogeny

Jessie Atterholt¹,² and Holly N. Woodward³

¹ Graduate College of Biomedical Sciences, Western University of Health Sciences, Pomona, California, United States
² Department of Integrative Biology, University of California, Berkeley, California, United States
³ Department of Anatomy and Cell Biology, Oklahoma State University Center for Health Sciences, Tulsa, Oklahoma, United States

ABSTRACT

Bone histology of crown-group birds is a research topic of great interest, permitting insight into the evolution of remarkably high growth rates in this clade and variation across the altricial-precocial spectrum. In this study, we describe microanatomical characteristics of the humerus and femur in partial growth series from 14 crown group birds representing ten major clades (Struthioniformes, Galliformes, Apodiformes, Columbiformes, Charadriiformes, Accipitriformes, Strigiformes, Psittaciformes, Falconiformes, and Passeriformes). Our goals were to: (1) describe the microanatomy of each individual; (2) make inter-and intra-taxonomic comparisons; (3) assess patterns that correspond with developmental mode; and (4) to further parse out phylogenetic, developmental, and functional constraints on avian osteological development. Across taxa, the femoral and humeral tissue of neonates can be broadly characterized as highly-vascularized, disorganized woven bone with great variation in cortical thickness (inter-and intrataxonomically, within an individual specimen, and within a single section). The tissue of precocial chicks is relatively more mature at hatching than in altricial, but other categories along the developmental spectrum were less easy to distinguish, thus we were unable to identify an definitive histological proxy for developmental mode. We did not find evidence to support hypotheses that precocial chicks exclusively have thicker cortices and more mature bone in the femur than the humerus at time of hatching; instead, this is a characteristic of nearly all taxa (regardless of developmental mode), suggesting deep evolutionary origins and the effects of developmental channeling. Bone tissue in adults exhibited unexpected variation, corresponding to differences in body size. Large-bodied birds have cortices of fibrolamellar bone, but organization of tissue increases and vascularity decreases with diminishing body size. The outer circumferential layer (OCL) also appears at earlier growth stages in small-bodied taxa. Thus, while the OCL is indicative of a cessation of appositional growth it is not always indicative of cortical maturity (that is, maximum organization of bony tissue for a given taxon). Small size is achieved by truncating the period of fast growth; manipulation of the timing of offset of bone growth is therefore an important factor in changing growth trajectories to alter adult body size.
INTRODUCTION

The description and evolution of osteohistological structures and characteristics of avian adult bone have been a subject of scientific investigation for decades, particularly following the discovery of the dinosaurian ancestry of birds. Modern birds are extremely fast-growing animals (de Ricqlès et al., 1991; Starck & Ricklefs, 1998; Erickson et al., 2009; Wilson & Chin, 2014), as were many non-avian dinosaurs. However, many initial studies of bone histology in Mesozoic avialans concluded that stem-group birds were either moderately more slow-growing (as in ornithuromorphs (Chinsamy, Chiappe & Dodson, 1995; Bell et al., 2010)), or drastically slower (as in the case of enantiornithines and Archaeopteryx (Chinsamy, Chiappe & Dodson, 1994; Cambra-Moo et al., 2006; Erickson et al., 2009; O’Connor et al., 2014)). Subsequent work has begun to reveal a greater diversity and complexity of life history strategies among these Mesozoic clades than previously understood. In Archaeopteryx, Voeten et al. (2018) demonstrated that immature ontogenetic stages of this taxon have vascular areas comparable to some extant taxa, and therefore likely a higher rate of bone growth than previously known (at least at certain growth stages). O’Connor et al. (2015) show evidence of a growth strategy similar to modern birds in the derived ornithuromorph Iteravis huchzermeyeri. Wang et al. (2019) conclude the same for the ornithuromorph Yanornis. In an ontogenetic study of Confuciusornis, Chinsamy et al. (2020b) report that this taxon likely experienced rates of growth comparable to some modern birds for a time in early-to mid-ontogeny, and furthermore present evidence that Confuciusornis retained a degree of developmental plasticity that allowed this taxon to phenotypically respond to environmental cues. Atterholt et al. (2021) report on fibrolamellar and incipient fibrolamellar bone in at least some skeletal elements of a Late Cretaceous skeletally-mature enantiornithine. A growing body of evidence supports the idea that fast, yet intermittent growth is the plesiomorphic condition for Avialae, and that a complex evolutionary pattern of both losses and amplifications of these features appears in Mesozoic and extant members of this clade (Prondvai et al., 2018; Wang et al., 2019; Chinsamy et al., 2020b; Wang et al., 2020).

Amid an increasing abundance of paleohistological studies on avialan and nonavialan dinosaurs, still relatively little is known about how bone develops in Aves, their closest living relatives. Bird bones are very thin-walled because resorption is a highly active process in avian bone growth, erasing most ‘history’ of development, so adult osteohistological features reveal very little about earlier growth stages. We instead must rely on ontogenetic studies of bone histology, which fortunately are increasingly common. Post-natal histology of bird bone has been examined in several taxa, however, the vast majority of studies focus on paleognathous birds or galloanseriforms (Castanet et al., 1996; Rath et al., 1999; Castanet et al., 2000; de Margerie, Cubo & Castanet, 2002; Skedros & Hunt, 2004; Turvey & Holdaway, 2005; Kuehn et al., 2019). The former are of particular interest likely because they are the early-branching clade of Aves, and thus are often
perceived as being more representative of dinosaurian growth. Galloanseriforms understandably draw the attention of researchers because of their importance in the poultry industry, and because specimens are readily available since they are domestically bred and farmed. Comparatively few studies have investigated histological characteristics outside of these early-diverging clades (de Margerie et al., 2004; Watanabe, 2018; McGuire et al., 2020).

Here, we present a histological description of post-natal skeletal development of the humerus and femur in 14 taxa representing a phylogenetically diverse sampling of ten major clades: Struthioniformes, Galliformes, Apodiformes, Columbiformes, Charadriiformes, Accipitriformes, Psittaciformes, Falconiformes, and Passeriformes (Fig. 1; Table S1). Specifically, we describe periosteal growth that occurs after a chick hatches, assessing patterns qualitatively and quantitatively, and making inter- and intrataxonomic comparisons. Bone is a highly complex tissue, and its growth and development are influenced by a variety of complicated functional and phylogenetic factors. In the case

![Phylogenetic tree used as evolutionary context in this study (topology based on Prum et al. (2015); branch lengths shown here do not reflect time calibration of the original tree). Colors represent developmental mode of each taxon: orange, precocial; yellow, semi-precocial; green, semi-altricial 1; blue, semi-altricial 2; purple, altricial. See Supplemental Table 2 for a break-down of traits used to define each category.](https://doi.org/10.7717/peerj.12160/fig-1)
of birds, this includes the challenge of adapting a skeleton simultaneously to aerial locomotion and terrestrial locomotion (or at least the demands of weight-bearing when perched). This is further complicated by shifting demands on bones throughout ontogeny, as degree of locomotion and locomotion type generally shift dramatically through development as chicks fledge and take to flight. Previous studies began identifying the specific ways in which phylogeny and function influence bone histology (Cubo et al., 2005; Montes, Castanet & Cubo, 2007; Cubo et al., 2008; Montes, Castanet & Cubo, 2010; Legendre et al., 2013; Padian & Lamm, 2013; Legendre et al., 2014; Padian & de Ricqlès, 2020). A major goal of this study is to further identify aspects of bone microstructure affected by these two channeling mechanisms, as well as by developmental constraint.

The final major objective of this investigation is to study ontogenetic histological patterns across the altricial-precocial spectrum. Birds, like mammals, are notable for the range of morphologies and behaviors they display at hatching (or birth), described as the altricial-precocial spectrum (Starck & Ricklefs, 1998). Neonatal chicks and their parents exhibit a full range of intermediate behavioral and morphological characteristics. At the precocial extreme of the spectrum, neonates are independent of their parents at hatching and even possess contour feathers and flight capabilities; here, there is no parental care after the eggs hatch. At the opposite end of the spectrum, the most altricial chicks hatch naked and with closed eyes. They remain nest-bound for the first few weeks of life, and are entirely dependent upon their parents for food, protection, etc. Various discretizations of the spectrum exist, with minor differences in the names and number of groups used to break it down into various developmental modes (e.g., Portmann, 1935; Nice, 1962; Skutch, 1976; Starck, 1993), but all rely on traits such as whether neonate eyes are open or closed, if chicks hatch naked or with feathers (and, in the latter case, what type of feathers), locomotor activity of the chick, and degree of parental care (e.g., showing to food, direct feeding, brooding, etc.) Here, we adopt the categories of developmental mode outlined by Starck & Ricklefs (1998), summarized in Table S2.

Of particular relevance to this investigation is evidence from previous studies showing that precocial chicks have growth rates much lower than those observed in altricial chicks (Ricklefs, 1968; Ricklefs, 1973; Starck, 1989; Starck, 1993; Starck & Ricklefs, 1998). Because growth rates, along with functional and physiological demands, vary so greatly across this spectrum, it is reasonable to expect substantial osteohistological variation in neonates of different developmental modes. Furthermore, identification of histological features that correlate with different developmental modes could be used as proxies for developmental mode in extinct taxa.

In studies of bone growth in the chicks of California gulls Carrier & Leon (1990) provided evidence that weak skeletal tissue (which they define to include a woven matrix) in very young individuals can be partially compensated for by increasing cortical thickness. Therefore, relative to other birds, gull chicks have very thick cortical walls in the femur because they begin locomoting terrestrially very soon after hatching, but the humerus has a thinner cortical wall until fledging, when the wing is finally used in flight. These data suggest certain predictions about cortical wall thickness of the two major limb elements in chicks at hatching, given information on developmental mode (whether or not the chicks
are active at hatching) and primary locomotor module (whether they will ultimately use primarily their wings, legs, or both to locomote), and the correlation between the two. Dial (2003) identified the primary locomotor module of adults of more altricial taxa as the pectoral limb, and of precocial taxa as the pelvic limb. Additionally, chicks on the altricial end of the spectrum do not locomote at all until weeks after hatching, while precocial chicks can generally walk independently very soon post-hatching. Therefore, altricial and semi-altricial chicks should have thin-walled bones in both the humerus and the femur or a slightly thicker-walled humerus (if investment in the elements of their primary locomotor module begins very early), and precocial and semi-precocial chicks (such as the gull) should have a thicker-walled femur.

Similarly, a difference in bone maturity between pelvic and pectoral limbs was reported by Dial & Carrier (2010) in Mallard ducks, who found that the functional maturity of the hindlimb was much greater than that of the forelimb through most of post-natal ontogeny, up until the time of fledging. Prondvai et al. (2020) also report a difference in cortical apposition rates related to developmental timing of the bone of limb elements in ducks. Evidence from such studies suggests that a difference in bone maturity is linked to the precocial and semi-precocial developmental mode of these taxa; such chicks locomote using the pelvic limb from the time of hatching, but do not require the pectoral limb for flight until much later. It follows, therefore, that these earlier functional demands on the pelvic limb have led to selection for this locomotor module maturing at an earlier time.

Therefore, we have a clear hypothesis regarding a histological signal of developmental mode: more precocial chicks should have more mature femoral bone, defined here as possessing a mosaic of various features indicating relatively more growth has occurred (a thicker cortex, more organized tissue, smaller vascular openings, and/or thicker woven bone trabeculae). In contrast, chicks closer to the altricial end of the spectrum should have more mature humeral bone than femoral (or at least of equal maturity and thickness, if delayed locomotion is not enough to select for a functional difference in neonate limb bones). The range of developmental modes covered by the taxa in this dataset present an excellent opportunity for testing this hypothesis.

**MATERIALS & METHODS**

**Acquisition of specimens**

Taxa for which complete or partial post-natal growth series were collected were included in the study. A post-natal growth series was defined as ranging from neonate chick to somatically mature adult (i.e., final size, after which little or no morphological change takes place); in some instances, individuals of intermediate ages were also available. Ultimately, 14 taxa from 10 avian families were used (Fig. 1; Table 1; Table S1), comprising 89 specimens and generating 356 histological thin sections. Because a major goal of this study was to acquire specimens representing a phylogenetically broad sampling of Aves, specimen acquisition was opportunistic and dependent upon donations from the OK Corral Ostrich Farm in Oro Grande, CA, USA (the source of the ostriches); Avian Resources in San Dimas, CA, USA (the source of the green-cheeked conures); the Lindsay Atterholt and Woodward (2021), PeerJ, DOI 10.7717/peerj.12160
Wildlife Museum in Walnut Creek, CA and the Society for Prevention of Animal Abuse in Monterey Co., CA, USA for all other taxa. Beyond the broad phylogenetic sampling that this method of specimen acquisition allows, another strong benefit is that most of the birds in the dataset (the native Californian taxa accumulated from local wildlife hospitals) were all wild birds that hatched and lived in their natural habitat and conditions, where they would have been subject to normal behavioral and biomechanical impacts on the growing skeleton; only the two taxa not endemic to California (ostriches and green-cheeked conures) were hatched and raised by the breeder/farmer.

All specimens were dissected and skeletonized at the Museum of Vertebrate Zoology (MVZ) at the University of California, Berkeley. Skeletons, histological slides (plus residual tissue blocks), and frozen soft-tissue samples all are housed in the MVZ.

### Table 1: Summary of sampling across growth stages for each taxon (number of specimens per growth stage), and timing of the first appearance of an incipient OCL (light red) and a distinctly-formed OCL (dark red).

| Taxon                | Neonate | Downy | Pin-feathered | Pre-fledgling | Fledgling | Sub-adult | Adult |
|---------------------|---------|-------|---------------|---------------|----------|-----------|-------|
| California Quail    | 8       | —     | 2             | —             | —        | —         | 2     |
| Wild Turkey         | 1       | —     | —             | 1             | —        | —         | 1     |
| Western Gull        | 2       | —     | —             | —             | —        | —         | 2     |
| American Kestrel    | 2       | —     | —             | —             | 1        | —         | 2     |
| Red-tailed Hawk     | —       | 1     | —             | 1             | —        | —         | 2     |
| White-tailed Kite   | —       | —     | 1             | 2             | 3        | —         | 1     |
| Barn Owl            | —       | 1     | 2             | 2             | 1        | —         | 2     |
| Great-horned Owl    | —       | —     | 1             | 1             | 1        | —         | 1     |
| Mourning Dove       | —       | —     | 1             | 2             | 2        | 2         | 1     |
| Anna's Hummingbird  | —       | —     | 1             | —             | —        | 1         | 1     |
| Green-cheeked conure| 1       | 1     | 1             | 1             | 1        | —         | 1     |
| Western Scrub-Jay   | 1       | —     | 1             | 3             | 2        | 1         | —     |
| House Finch         | 1       | —     | 1             | 2             | 1        | —         | 1     |

**Note:** This indicates the first appearance of this structure in each taxon over all, either in the humerus and/or the femur. Developmental mode is indicated by color number superscript: orange1, precocial; yellow2, semi-precocial; green3, semi-altricial 1; blue4, semi-altricial 2; purple5, altricial. Six ostrich chicks also comprise part of the dataset, but are not included here because (beyond neonate an adult) they do not fit into the growth stages used for other taxa.

This study examines bone as a mineralized tissue and is not an investigation of the process of ossification. Therefore, tissue samples were harvested from the youngest individuals with ossified diaphyses; in some cases, neonate bone was suitable, but in several altricial taxa neonate chicks were too small and had bones still too cartilaginous to be appropriate for the methods employed. By the time of hatching, most neonates have fully ossified bony diaphyses of the femur and humerus with cartilaginous proximal and distal ends, so tissue samples were taken at the mid-shaft portion of each element.

The humerus and femur were obtained from skeletonized individuals, and the mid-shaft regions harvested. Whenever possible, the left elements were used, though breakage or missing bones sometimes necessitated use of right elements. This investigation limited focus to the humerus and femur because these are two bones that, at their...
midshafts, undergo minimal secondary growth and remodeling, and therefore preserve a clear ontogenetic record (Padian & Lamm, 2013).

Tissue samples were fixed in 10% neutral buffered formalin for 48 h (with one change of solution after the first 24), then transferred to a solution of 70% ethyl alcohol for 48 h (with one change of solution after the first 24), subsequently placed in a solution of 85% ethyl alcohol for 48 h (with one change of solution after the first 24), and finally cleared in Histo-Clear (National Diagnostics) for 4–8 h, contingent upon the size of the tissue sample. Bone samples were then embedded in Epothin and sectioned to one mm wafers using a diamond-embedded saw. Wafers were mounted on glass slides and ground to approximately 100 µm thickness using a lap grinder and grit paper. Two sections from each bone were retained, and one of these mounted sections was stained in a 0.7% solution of toluidine blue. Finally, coverslips were applied using Permount (Fisher Chemical, Waltham, MA, USA). Slides were photographed using a Nikon digital sight camera and petrographic microscope (DS-U3 and DS-Fi2), and captured using the computer program NIS-Elements (F4.00.00). Sections were visualized under regular light and with cross-polarized light (XPL). Measurements were taken using ImageJ (1.48v).

**Experimental design & terminology**

Precise ages of most individuals at time of death are unknown due to the method of specimen collection. Therefore, in cases where exact numerical age was missing, growth series were divided into the following qualitative growth stages, modified from identifications made by the wildlife hospitals and based on body size and general external morphology (primarily the condition of the feathers): neonate, downy chick, pin-feathered chick, pre-fledgedling chick, fledgling chick, sub-adult, and adult. These categories were used to describe all taxa except for the ostrich, because these birds do not neatly fit into these categories. We note that, though these are artificial categories that fall along a spectrum, they roughly approximate the stages of feather development described by Prum & Brush (2002). Comparisons among intermediate growth stages are generally limited, and intertaxonomic comparisons are focused on adults and neonates (the two most ‘equal’ growth stages).

Due to decomposition and/or immaturity of specimens, sex was indeterminable by dissection for many specimens in the dataset. However, this is not considered a serious impediment to the study. The only major recorded histological difference in bone between male and female birds is medullary bone, variably present in adult females (Bonucci & Gherardi, 1975; Miller & Bowman, 1981; Van de Velde, Vermeiden & Bloot, 1985; Dacke et al., 1993).

This project is primarily descriptive. Each growth stage is described qualitatively in terms of general bone type, cross-sectional shape, variability of cortical thickness, density and orientation of blood vessels, density and shape of osteocyte lacunae, and presence of primary and secondary osteons. Osteocyte lacunal volume and density is known to be extremely variable intrataxonomically, among skeletal elements, and even regionally within a single bone, and is most accurately estimated when using multiple thin sections (D’Emic & Benson, 2013); therefore while a qualitative description of osteocyte lacunae is
included in this study, we do not quantify this feature or infer relative rates of growth and metabolism.

The terminology used to qualitatively classify and describe bone tissues follows the precedent of Francillon-Vieillot et al. (1990) and de Ricqlès (1975), as updated by Huttenlocker, Woodward & Hall (2013). Bone is described both in terms of matrix classification (based on fiber organization), and vascular classification (based on type and orientation of vascular channels).

The altricial-precocial spectrum has been divided into a number of categories based on morphological and behavioral features by multiple authors (see Starck & Ricklefs, 1998); here, we used the Starck & Ricklefs (1998) reorganization of Nice’s (Nice, 1962) developmental classes. Starck and Ricklefs’ categories are summarized in Table S2.

Bone histology was also analyzed quantitatively; all statistical tests were run using the program R (v4.0.3). For each section, average cortical thickness and average total cross-sectional diameter were measured in μm. The cortical thickness was an average of eight measurements evenly distributed across the cortex (midpoints of the cranial, caudal, medial, and lateral regions, and positions equidistant between these). Average total diameter of each sample was taken as an average of four measurements of the total diameter of the cortex, along the craniocaudal and mediolateral axes. and the two axes at the mid points between these. These data were used to calculate ratios for comparisons of relative cortical thickness as 2 (cortical thickness)/total diameter.

Cortical thickness is analyzed and discussed in two ways: absolute cortical thickness (ACT) and relative cortical thickness (RCT). These ratios together with average measures of cortical thickness and cortical diameter can be found in Table S1. A log10 transformation was applied to measures of absolute cortical thickness (to correct for strong allometry as associated with body size) and to relative cortical thickness (to normalize the data, which had an exponential distribution). General patterns of variation and changes in cortical width (by growth stage and intertaxonomically) are visualized and described using boxplots. Phylogenetic signal was tested for in ACT and RCT of the humerus and femur of adults and neonates, as well as in developmental mode, using Blomberg’s K test statistic (Blomberg, Garland & Ives, 2003). The phylogenetic topology used for this test is based on the tree in Prum et al. (2015), using the same calibrations as the original published phylogeny. Finally, regression tests were implemented to assess intertaxonomic relationships (with relevant metrics from all individuals pooled into a common dataset) between cortical thickness of the adult humerus and femur, cortical thickness of the neonate humerus and femur, and developmental mode with cortical thickness of each element for neonates. Specific models (and results for each) are described in Tables 2 and 3. For these tests, the developmental spectrum was treated as a pseudo-continuous ordinal predictor (Symonds & Blomberg, 2014) and numerically discretized (based on the categorization of Starck & Ricklefs (1998) as follows: 1, precocial; 2, semi-precocial; 3, semi-altricial 1; 4, semi-altricial 2; 5, altricial. A phylogenetic generalized least squares regression (PGLS) was used in cases when model residuals had phylogenetic signal; in all other cases, an ordinary least squares regression (OLS) was used.
RESULTS

Broad intertaxonomic patterns and comparisons are reported here; for a detailed description of each individual and intrataxonomic comparisons, please see Supplementary Material.

General observations of variation in cortical thickness

The thickness of the cortex, both on an absolute and relative scale, was highly variable across growth stages, taxa, and developmental mode (Figs. 2–4). Analysis of both RCT and

---

**Table 2** Results of tests for phylogenetic signal in developmental mode and cortical thickness of humeri and femora of adults and neonates, using Blomberg’s $K$ test-statistic.

| Parameter tested | $K$ Statistic | $P$-value |
|------------------|---------------|-----------|
| Adults, Humerus (ACT) | 0.804 | 0.591 |
| Adults, Femur (ACT) | 1.060 | 0.197 |
| Neonates, Humerus (ACT) | 0.804 | 0.613 |
| Neonate, Femur (ACT) | 1.061 | 0.207 |
| Adults, Humerus (RCT) | 0.860 | 0.433 |
| Adults, Femur (RCT) | 1.101 | 0.089 |
| Neonates, Humerus (RCT) | 0.813 | 0.689 |
| Neonate, Femur (RCT) | 1.235 | 0.043 * |
| Developmental Mode | 1.317 | 0.005 * |
| log10(NHACT) ~ log10(NFACT) + log10(NBM) | 1.286 | 0.028 * |

**Note:** ACT, absolute cortical thickness; RCT, relative cortical thickness; measured as the ratio of average cortical thickness to average total cross-sectional diameter. The final row shows results of a test for phylogenetic signal in the residuals from a PGLS analysis of the model described (NHACT, neonate humerus ACT; NFACT, neonate femur ACT; NBM, neonate body mass). Significant $P$-values are indicated by asterisks.

**Table 3** Results of regression analyses between adult and neonate humerus and femur cortical thickness, and developmental mode.

| Regression Model | $F$ Statistic | $P$-value ($F$ Statistic) | Adjusted $R^2$ | $P$-value (Adjusted $R^2$) |
|------------------|---------------|---------------------------|----------------|---------------------------|
| log10(NHACT) ~ log10(NFACT) + log10(NBM) | 0.81; 0.96 | 0.466; 0.392 | 0.65 | 0.054 |
| log10(AHACT) ~ log10(AFACT) + log10(ABM) | 5.53; −0.40 | 0.007 $\times 10^{-4}$; 0.692 | 0.91 | 0.002 $\times 10^{-5}$ * |
| log10(AHRCT) ~ log10(AFRCT) | — | — | 0.23 | 0.029 |
| log10(NHRCT) ~ log10(NFRCT) | — | — | 0.15 | 0.095 |
| DM ~ log10(NHACT) + log10(NBM) | 0.45; −0.75 | 0.660; 0.468 | −0.11 | 0.709 |
| DM ~ log10(NFACT) + log10(NBM) | 3.06; −3.15 | 0.051; 0.023 * | 0.39 | 0.050 |
| DM ~ log10(NHRCT) | — | — | −0.03 | 0.442 |
| DM ~ log10(NFRCT) | — | — | −0.06 | 0.657 |

**Note:** The first model was assessed using PGLS. All other models were assessed using OLS. ACT, absolute cortical thickness; RCT, relative cortical thickness; measured as the ratio of average cortical thickness to average total cross-sectional diameter. Significant $p$-values are indicated by asterisks. Abbreviations used in models: NHACT, neonate humerus ACT; NFACT, neonate femur ACT; NBM, neonate body mass; AHACT, adult humerus ACT; AFACT, adult femur ACT; ABM, adult body mass; AHRCT, adult humerus RCT; AFRCT, adult femur RCT; NHRCT, neonate humerus RCT; NFRCT, neonate femur RCT; DM, developmental mode.
Figure 2  Boxplots showing the variation of cortical thickness by growth stage, where red represents measurements of the humerus and green measurements of the femur. (A) Absolute cortical thickness (ACT). (B) Relative cortical thickness (RCT). Within each growth stage, all samples (from the humerus and femur) across all taxa are included, excepting ostriches (which were not categorized using these growth stages). Boxes = interquartile range, or values between the 25th and 75th percentiles; horizontal black bars = median values; whiskers = minimum and maximum values, not including outliers; circles = outliers.
ACT is included in the interest of conducting a thorough study, and because in some instances each metric reveals interesting patterns that the other does not. Absolute cortical thickness appears to undergo an overall trend of increase in early ontogeny to a maximum thickness around the pin-feathered stage, a subsequent decrease from the fledgling to sub-adult stages, and final moderate increase to adult cortical thickness (Fig. 2A). Interestingly, such a trend was also frequently observed at the scale of individual taxa (see Supplementary Material), though this more nuanced and detailed description also revealed

![Boxplots showing variation in cortical thickness by taxon, where red represents measurements of the humerus and green measurements of the femur.](Full-size DOI: 10.7717/peerj.12160/fig-3)
a frequent pattern of a secondary increase at the fledgling stage (rather than a uniformly high ACT from pin-feather to fledgling stages). The pin-feathered stage exhibits the highest variance in both elements, both overall and for the interquartile range. In most growth stages, variance of the humerus and femur are similar in magnitude. Most measures of ACT fall within a narrower range of values for the femur than the humerus in neonates and adults. In pre-fledgling chicks, the humerus appears to have a more restricted range of ACT. In neonates, ACT of the femur is generally greater than in the humerus; this relationship is inverted in adults, where ACT of the humerus is generally higher.

Ontogenetic changes in RCT reflect the overall patterns observed in ACT (Fig. 2B): an increase in early ontogeny peaking at the pin-feathered stage, followed by a decrease, and final moderate increase from sub-adult to adult. However, the early increase is disrupted by a reduction in RCT from the neonate to downy stages before reaching a maximum at the pin-feathered stage. At hatching, neonate RCT is more variable in the

---

**Figure 4** Boxplots showing variation in cortical thickness by developmental mode, where red represents measurements of the humerus and green measurements of the femur. (A) Absolute cortical thickness (ACT). (B) Relative cortical thickness (RCT). For each group, the humerus and femur of all neonate specimens within a particular developmental mode are represented. Additionally, “semi-altricial” here refers to “semi-altricial 1;” the elements of the semi-altricial 2 neonate were not ossified and thus not examined in this study. Boxes = interquartile range, or values between the 25th and 75th percentiles; horizontal black bars = median values; whiskers = minimum and maximum values, not including outliers; circles = outliers.
femur than the humerus. Neonate median values of RCT are higher in the humerus than femur, but the medians are equal at the adult stage. Highest RCT is reached at the pin-feathered chick stage. The greatest individual value is seen in the humerus of the pin-feathered house finch chick (RCT = 0.78). However, this stage also has the highest variance, so certainly not all pin-feathered chicks have a very thick cortex. Anna’s hummingbird pin-feathered chicks had the lowest RCT in the dataset (RCT = 0.12 in the humerus and femur). Overall variance and interquartile variance generally tend to decrease through ontogeny.

Predictably, absolute cortical thickness is highest in ostriches, and lowest in Anna’s hummingbird, reflecting the body size extremes represented in this dataset (Figure 3A). However, the wild turkey, Western gull, great-horned owl, red-tailed hawk, barn owl, and white-tailed kite all have high ACT, within range of the values observed in the ostriches (though the dataset notably is lacking samples from an adult ostrich). Other taxa appear to roughly group according to medium values of ACT (California quail, mourning dove, American Kestrel, green-cheeked conure, and house finch). These general groupings also parallel body size differences, in spite of the log transformation of the data.

In many taxa (great-horned owl, red-tailed hawks, Anna’s hummingbird, white-tailed kites, American kestrels, barn owls, scrub jays, and house finches), the femur exhibits higher variance overall in ACT than the humerus. Variance in ACT is greater in the humerus than the femur in Western gulls, green-cheeked conures, and ostriches, and roughly equal between the two elements in California quail, wild turkeys, and mourning doves. Highest levels of variance in ACT are seen in the California quail, Western gull humerus, wild turkey, green-cheeked conure, and ostrich humerus (Fig. 3A). It is striking that these taxa span a wide ranges of body sizes, primary locomotor module, and developmental mode. Variance of ACT is low in great-horned owls, barn owls, white-tailed kites, red-tailed hawks, mourning doves, and scrub jays, though the degree of difference in variation is quite possibly due to a lack of available samples representing the extremes of the growth stages (no adult for scrub jays, and no neonates for the rest).

Intertaxonomically, the highest values of RCT were observed in the humerus and femur of the house finch pin-feathered chick, where the cortex comprises 78% and 70% of total cross-sectional diameter, respectively. Taxa represented by more specimens at the pin-feathered and pre-fledging stage exhibit higher variance than others, further underscoring the major increase in cortical thickness during these early growth stages (Fig. 3). Interquartile range and total variance for each element are close and usually overlapping in nearly all taxa. However, the RCT of ostrich chick humeri is much higher than that of the femora, reflecting the sharp differences in cortical maturity observed between the two elements in this taxon. The predominant pattern across nearly all other taxa is a higher variance of RCT in the femur, either drastically (in Anna’s hummingbirds, Western gulls, American kestrels, Western scrub jays, and house finches) or moderately (as in California quail, wild turkeys, mourning doves, green-cheeked conures, white-tailed kites, and red-tailed hawks). The single exception to this pattern is the barn owl, in which the humerus exhibits higher variance. We are cautious about interpreting these observations since intrataxonomic sample metrics would be altered by the addition of
more specimens representing more growth stages in most if not all cases. However, the
consistency of this pattern across a range of samples sizes, phylogenetic differences, and
various developmental modes, is notable. Overall, however, the interspecific interquartile
range of RCT falls within a relatively narrow span of values across taxa (especially
compared to ACT); only the ostrich deviates from this trend, and only then with respect
to the humerus. In the femur, this pattern is sustained when making intraspecific
observations, but in notable contrast, intraspecific variation is generally greater in the RCT
of the humerus than in ACT.

Distributions of ACT and RCT by developmental mode are presented in Fig. 4.
The semi-precocial developmental mode appears to have the highest values of ACT for
both elements (based on our limited sampling) (Fig. 4A). The greatest variance of ACT is
observed in altricial neonates, which represent both the lowest and some of the highest
values of ACT (excepting semi-precocial ACT). ACT of semi-altricial chicks appears to fall
within the same range as that of altricial neonates. Interestingly, there is relatively little
difference in ACT between the two extremes of the spectrum. ACT of the humerus in
precocial chicks falls also within the range of measurements of humeral ACT for altricial
neonates (though with a variance of a much lesser magnitude). Femoral ACT of precocial
neonates also overlaps with the range observed in altricial chicks, though values are
generally a bit higher in the latter group. Additionally, femoral ACT is higher than humeral
ACT across the altricial precocial spectrum (in terms of variance, interquartile range, and
median values), though when considering taxa on an individual basis this pattern is not
born out in the wild turkey neonate.

In terms of RCT (Fig. 4B), altricial neonates have the highest values for both the
humerus and femur, though precocial neonates fall within a very similar range. Once
again, this developmental mode also has a variance of the greatest magnitude, for both
elements. Notably, the RCT of precocial chicks is generally quite a bit lower than all other
developmental modes. This is also the only group in which RCT is higher in the humerus
than in the femur. Furthermore, values of RCT for semi-altricial chicks are closer to
those for precocial neonates than altricial neonates.

**Phylogenetic tests**
We tested for phylogenetic signal in average cortical width of the humerus and femur in
the adult and neonate, as well as in developmental mode (Table 2); cortical width was
analyzed both as an absolute measurement (ACT), and relative (RCT). Phylogenetic
signal was not significant in ACT of either element for either growth stage. Significant
signal was recovered in the RCT of the neonate femur, but not the neonate humerus.
No significant phylogenetic signal was detected in RCT of either bone in adults.
Developmental mode has significant phylogenetic signal, a result that has been reported by
previous authors (Starck & Ricklefs, 1998; Atterholt, 2011).

Regression analyses were used to assess relationships between RCT and ACT in
adult and neonate humeri and femora, comparing element to element within the same
growth stage, and comparing each neonate metric individually to developmental mode
(see results in Tables 2 and 3). The only model with significant phylogenetic signal was that
assessing the relationship between ACT of the neonate humerus and femur \((K = 1.286; P = 0.028)\). The PGLS results for this regression, however, were not indicative of a significant effect of femoral ACT or body mass on humeral ACT in neonates, either individually or together (Table 3). In contrast, ACT of the adult humerus and femur samples is strongly correlated \((R = 0.91; P = 0.002 \times 10^{-5})\). Furthermore, results indicate that body mass is not significantly related to humerus ACT when the effects of femur ACT is accounted for \((F = −0.40; P = 0.692)\). However, RCT of the humerus is not significantly related to that of the femur in neonates or adults (Table 3).

None of the various measures of cortical thickness in neonates exhibit significant correlations with developmental mode, though we note this may be due to the relatively small sample size of neonates in this study. However, body mass has an effect even at the neonate growth stage in ACT of the femur, with larger-bodied chicks having thicker femoral cortices.

**Histology of neonates & other immature developmental stages**

The wild turkey and Western gull neonates have the thickest humeral cortices as an absolute measure \((220.28\) and \(218.98\) µm respectively), and the conure the thinnest \((39.38\) µm) (Table 4). In terms of RCT (Table 5), the Western scrub-jay has the thickest humeral cortex at 48% total cross-sectional diameter; the conure has thinnest at 18% total cortical diameter. Other taxa fall within a range of 24–34%. Qualitative characteristics of neonate humeri are summarized in Table 6.

Humeri of neonates are almost uniformly circular in cross-section (Fig. 5); the two exceptions are the turkey, which has a rounded-triangle cross-sectional shape, and the quail, in which it is elliptical with a flat lateral margin. Vascular density is very high in all individuals, with many canals perforating a disorganized cortex of woven bone (Figs. 6–10). However, the thickness of the developing cortical trabeculae of the bone and the size of the channels \((i.e.,\) porosity) differ among taxa. Precocial chicks have more ‘mature’ humeri with relatively lower vascular porosity and thicker struts of woven bone (Figs. 6A & 6B). This is particularly noticeable in the quail and turkey, which have humeral bone of the most mature appearance out of all neonates examined. Apart from porosity, neonate humeri are all remarkably similar, consisting of woven bone not only with collagen fibers arranged in a disorganized way, but also with vascular channels displaying much irregularity in size, shape, and orientation. Most humeri can be characterized no more specifically than as having many longitudinal and circumferential channels of greatly variable shapes. Cortical thickness is often asymmetrical in the neonate humerus, to greater or lesser degrees, always showing variation in porosity in correlation with cortical width; narrow cortical areas have denser bone, while vascular channels are larger and more numerous in the thicker regions. This variation did not occur consistently in any particular directional region.

The thickness around the cortex of neonate femora was less variable than in the humerus, displaying patterns of variation more similar to that observed in adults (Fig. 5); qualitative features of neonate femora are summarized in Table 7. The femur therefore appears less variable and subject to change through ontogeny in terms of these
macroscopic characteristics, possibly a reflection of the microanatomical evidence that this element is relatively further along in development than the humerus (as observed in most taxa in this dataset—see below for detailed description). In terms of ACT, the Western gull has the thickest cortex (405.71 µm) and the conure has the thinnest (113.68 µm), though the quail is nearly as thin (117.99 µm). The relatively thickest cortex was observed in the scrub jay (29%), and the thinnest in the wild turkey (9%), though the California quail has a nearly equivalent cortex at 11% total cross-sectional diameter (Table 5).

Neonate femora are all circular in cross-section (Fig. 5). The bone of the neonate femur shares some features with the humerus. It is composed of a woven bone matrix of disorganized fibers perforated by numerous vascular channels that are highly irregular in shape and orientation. The relative size of these channels, and thus the porosity of the bone, is one of the characteristics that differs most among taxa. The California quail and wild turkey have the lowest porosity, i.e., canals of the smallest relative size. Vascular channels in the femora of these two taxa are also most organized of all the neonates, and can be unambiguously classified as longitudinal. In the gull, porosity is high, but the

| Taxon                                      | Neonate humeral cortical thickness | Neonate femoral cortical thickness | Adult humeral cortical thickness | Adult femoral cortical thickness |
|--------------------------------------------|-----------------------------------|-----------------------------------|---------------------------------|---------------------------------|
| California Quail ([Callipepla californica](#)) | 90.67                             | 117.99                            | 395.36                          | 380.76                          |
| Wild Turkey ([Meleagris gallopavo](#))     | 220.28                            | 185.92                            | 1223.12                         | 1240.14                         |
| Western Gull ([Larus occidentalis](#))      | 218.98                            | 405.71                            | 1105.81                         | 632.64                          |
| American Kestrel ([Falco sparverius](#))   | 124.86                            | 153.60                            | 359.75                          | 261.49                          |
| Red-tailed Hawk ([Buteo jamaicensis](#))   | no sample                         | no sample                         | 800.49                          | 625.18                          |
| White-tailed Kite ([Elanus leucurus](#))   | no sample                         | no sample                         | 503.06                          | 382.01                          |
| Barn Owl ([Tyto alba](#))                  | no sample                         | no sample                         | 463.79                          | 545.27                          |
| Great-horned Owl ([Bubo virginianus](#))   | no sample                         | no sample                         | 752.79                          | 736.66                          |
| Mourning Dove ([Zenaida macroura](#))      | unossified                        | unossified                        | 350.61                          | 229.60                          |
| Anna’s Hummingbird ([Calypte anna](#))     | unossified                        | unossified                        | 70.55                           | 46.18                           |
| Green-cheeked conure ([Pyrrhura molinae](#)) | 39.38                             | 113.68                            | 266.38                          | 184.81                          |
| Western Scrub-Jay ([Aphelocoma californica](#)) | 208.99                           | 361.33                            | no sample                       | no sample                       |
| House Finch ([Haemorhous mexicanus](#))    | unossified                        | 166.63                            | 190.34                          | 98.44                           |

Note:
Developmental mode is indicated by color number superscript: orange, precocial; yellow, semi-precocial; green, semi-altricial 1; blue, semi-altricial 2; purple, altricial. All measurements are given in µm.

Table 4: Average absolute cortical thickness (ACT) of the humerus and femur for all adults and neonates used in this study.
Table 5  Average relative cortical thickness as compared to the average total cross-sectional diameter (RCT) of the humerus and femur for all adults and neonates used in this study.

| Taxon                            | Neonate humeral cortical thickness | Neonate femoral cortical thickness | Adult humeral cortical thickness | Adult femoral cortical thickness |
|----------------------------------|-----------------------------------|-----------------------------------|----------------------------------|----------------------------------|
| California Quail*                | 0.24                              | 0.22                              | 0.28                             | 0.26                             |
| (Callipepla californica)         |                                   |                                   |                                  |                                  |
| Wild Turkey*                    | 0.30                              | 0.18                              | 0.20                             | 0.24                             |
| (Meleagris gallopavo)           |                                   |                                   |                                  |                                  |
| Western Gull*                   | 0.34                              | 0.44                              | 0.34                             | 0.28                             |
| (Larus occidentalis)            |                                   |                                   |                                  |                                  |
| American Kestrel*               | 0.24                              | 0.28                              | 0.22                             | 0.18                             |
| (Falco sparverius)              |                                   |                                   |                                  |                                  |
| Red-tailed Hawk*                | no sample                         | no sample                         | 0.20                             | 0.18                             |
| (Buteo jamaicensis)             |                                   |                                   |                                  |                                  |
| White-tailed Kite*              | no sample                         | no sample                         | 0.20                             | 0.18                             |
| (Elanus leucurus)               |                                   |                                   |                                  |                                  |
| Barn Owl*                       | no sample                         | no sample                         | 0.20                             | 0.30                             |
| (Tyto alba)                     |                                   |                                   |                                  |                                  |
| Great-horned Owl*               | no sample                         | no sample                         | 0.18                             | 0.22                             |
| (Bubo virginianus)              |                                   |                                   |                                  |                                  |
| Mourning Dove*                  | unossified                        | unossified                        | 0.20                             | 0.20                             |
| (Zenaida macroura)              |                                   |                                   |                                  |                                  |
| Anna’s Hummingbird*             | unossified                        | unossified                        | 0.18                             | 0.18                             |
| (Calypte anna)                  |                                   |                                   |                                  |                                  |
| Green-cheeked conure*           | 0.18                              | 0.40                              | 0.22                             | 0.20                             |
| (Pyrrhura molinae)              |                                   |                                   |                                  |                                  |
| Western Scrub-Jay*              | 0.48                              | 0.58                              | no sample                        | no sample                        |
| (Aphelocoma californica)        |                                   |                                   |                                  |                                  |
| House Finch*                    | unossified                        | 0.46                              | 0.26                             | 0.18                             |
| (Haemorhous mexicanus)          |                                   |                                   |                                  |                                  |

Note: Developmental mode is indicated by color number superscript: orange 1, precocial; yellow 2, semi-precocial; green 3, semi-altricial 1; blue 4, semi-altricial 2; purple 5, altricial.

Trabeculae of woven bone are quite thick. In the conure, scrub-jay, house finch, and kestrel, porosity is highest and the trabeculae are thinnest. In all of these taxa, it is difficult to classify predominant vascular orientation because there is such variation in the shape of channels and (Figs. 11–15; Table 7). Overall, these differences appear to reflect differences in maturity of the bone at hatching. Precocial quail and turkeys have the most mature bone, followed by semi-precocial gulls, while the most immature bone is seen in the altricial and semi-altricial chicks. Also as in the humerus, neonate femora display a hugely varying asymmetry in cortical thickness, and corresponding differences in density/porosity.

Chicks in the pin-feathered through fledgling stages frequently are distinguished by the presence of a thick, prominent endosteum that was seen only as a very thin layer in adults and neonates (when visible at all). Though we did not prepare our sections for study of soft tissues, when inadvertently preserved in a section, the soft tissue endosteum was consistently thick and prominent in the pin-feathered through fledgling stages, and only observed as a thin layer in neonates and adults. Indeed, a thickened endosteum was of one
of the only distinguishing features between sub-adult and adult birds. Of particular importance to growing chicks, the endosteal tissue functions to house osteoprogenitor cells and providing an important environment for production of hematopoetic stem cells and multipotent cells (Hall, 2005; Eroschenko, 2008; Cordeiro-Spinetti, Taichman & Balduino, 2015), all of which would be particularly important in growing chicks.

In the two passerines included in this study, a unique feature was observed in early growth stages: a circlet of bone in the endosteal cavity almost completely detached from the rest of the developing cortex (Figs. 5H, 10C, 15H). In Western scrub jays, this was present in the humerus and femur of the neonate. In the house finch, this was observed in both the humerus and femur of the pin-feathered stage. We interpret this intracortical gap as evidence of a Kastschenko line (Francillon, 1980), a thin layer of osteoid tissue that occurs between growing endosteal and periosteal bone, degraded and no longer present in these thin sections but leaving behind a space where it used to be.

Chondroid bone has been described as abundantly present in the bones of developing Rouen ducks by Prondvai et al. (2020). Similarly, chondroid bone was very common in many growth stages of most taxa observed here (e.g., Figs. 6A, 14A, 15D). It is present in

Table 6 Qualitative comparisons among neonate humeri.

| Taxon                     | Cross-sectional shape         | Bone composition & porosity                        | Primary vascular orientation | Other notable features                                                                 |
|---------------------------|-------------------------------|---------------------------------------------------|------------------------------|----------------------------------------------------------------------------------------|
| California Quail¹         | elliptical, flattened lateral margin | moderate, canals relatively small; bone predominates cortex | longitudinal                 | lateral edge thinner and with fewer vascular canals                                    |
| (Callipepla californica)  |                               |                                                   |                              |                                                                                        |
| Wild Turkey³              | rounded triangle               | moderate and very uneven porosity                 | longitudinal                 | medial cortex much thicker than lateral, canals much more numerous in thicker regions   |
| (Meleagris gallopavo)     |                               |                                                   |                              |                                                                                        |
| Western Gull²             | circular                       | high porosity, but woven bone still substantial    | longitudinal/irregular        | some variation in cortical thickness, canals larger in thicker regions                  |
| (Larus occidentalis)      |                               |                                                   |                              |                                                                                        |
| American Kestrel³         | circular                       | moderate porosity, substantial bone present       | longitudinal/irregular        | very thin cortex; few vascular canals                                                  |
| (Falco sparverius)        |                               |                                                   |                              |                                                                                        |
| Green-cheeked conure⁴     | circular                       | moderate porosity, substantial bone present       | all orientations             | extremely thin cortex; few vascular canals                                              |
| (Pyrrhura molinae)        |                               |                                                   |                              |                                                                                        |
| Western Scrub-Jay⁴        | circular                       | high; bony bone very thin                         | all orientations             | solid endosteal circlet of bone present; highly asymmetrical cortical thickness         |
| (Aphelocoma californica)  |                               |                                                   |                              |                                                                                        |

Note: Note that vascular density was high almost ubiquitously among these individuals, therefore porosity is compared among neonates, nearly all have high porosity relative to adults. Results are shown for taxa for which neonate specimens were available, and in which the humerus was ossified and viable for this study (the humeri of the mourning dove, Anna’s hummingbird, and house finch were not included for the latter reason). Developmental mode is indicated by color number superscript: orange¹, precocial; yellow², semi-precocial; green³, semi-altricial 1; purple⁴, altricial.
Figure 5 Diaphyseal cross-sections of youngest individuals with ossified humeral and femoral diaphyses included in this study, showing general cross-sectional geometry and variation in cortical thickness. All are neonates, except in the case of Anna’s hummingbird and mourning dove, shown here at the pin-feathered stage. Only femur of the neonate house finch was ossified, and therefore included in this study. For each pair, the humeral section is on the left and the femoral on the right. In all images cranial is up and lateral is left. Sections are grouped by developmental mode. Precocial (orange): (A) Wild turkey (Meleagris gallopavo; MVZ190764); (B) California quail (Callipepla californica; MVZ190745). Semi-precocial (yellow): (C) Western gull (Larus occidentalis; JAA264). Semi-precocial 1 (green): (D) American kestrel (Falco sparverius; MVZ190890). Semi-precocial 2 (blue): (E) mourning dove (Zenaida macroura; MVZ190778). Altricial (purple): (F) green-cheeked conure (Pyrrhura molinae; MVZ190895); (G) Western scrub jay (Aphelocoma californica; MVZ190927); (H) house finch (Haemorhous mexicanus; MVZ190969); (I) Anna’s hummingbird (Calypte anna; MVZ190799). Scale bars for the turkey sections = 500 µm. All other scale bars = 100 µm. For cross-sections of all ages, see figures in Supplemental Material. DOI: 10.7717/peerj.12160
Figure 6 Close-up photographs of microanatomical details in the humeri of precocial birds, sections from neonates shown in the top row and from adults in the bottom row. (A) Wild turkey neonate (MVZ190764); (B) California quail neonate (MVZ190751); (C) wild turkey adult (MVZ190763), showing fibrolamellar bone of the middle cortical layer where it meets the ICL; (D) California quail adult (MVZ190762). In all images the periosteal surface is oriented to the upper left and the endosteal to the lower right. Scale bar = 100 µm.

Figure 7 Close-up photographs of microanatomical details in the humeri of semi-precocial birds, sections from neonates shown in the top row and from adults in the bottom row. (A) Western gull neonate (JAA64); (B) Western gull neonate (MVZ190822); (C) Western gull adult (MVZ190829); (D) Western gull adult (MVZ190831). In all images the periosteal surface is oriented to the upper left and the endosteal to the lower right. Scale bar = 100 µm.
large areas of bone in younger chicks, and disappears approximately by the fledgling stage in most taxa, though was observed up to the subadult stage in Anna’s hummingbird. This discovery of wide-spread chondroid bone across avian taxa supports the results and predictions of Prondvai et al. (2020), and will be described and addressed in greater detail in a future publication.

**Adult histology**

Across somatically mature adults, the humerus varied in cross-sectional shape from roughly triangular (turkey) to ovate (quail, dove, hummingbird) to circular (all other taxa); many humeri of adult individuals are moderately flattened along one margin (Fig. 16); qualitative features of adult humeri, including cross-sectional shape, are summarized in Table 8. Some of this variation may reflect which portion of the shaft the tissue sample is from (the midshaft region was sampled generally, but some sections are marginally more proximal or distal), though it is notable that the femur, sampled in the same way, does not show such variation. This developmental stage displayed the least variance in cortical thickness throughout the circumference of the bone, with all cortices of uniform or nearly uniform breadth. In many taxa ACT reaches a maximum at adult body size, but RCT of the humerus was generally thinnest in adults (though sometimes very early growth stages were equally thin, see Table S1). Western gulls have the highest RCT (0.34). In all other birds, cortical width ranged from 18–28% of cortical diameter, a consistency
remarkable considering the phylogenetic breadth and locomotor diversity of the sample size.

High levels of variation were also observed in the density of vascular canals relative to anatomical location within the cortex (particularly traversing from inner to outer cortex), orientation of these channels, and relative thinness of the ICL and OCL. The size of vascular channels was small in all individuals, but qualitative data suggest porosity may be correlated with body size: canals occurred in very low densities in hummingbirds, and moderate numbers in doves, conures, finches, and kestrels; in all other taxa, vascular density was high. All humeri showed almost the full range of vascular canal orientations, but in the majority a reticular or longitudinal pattern was dominant; only the wild turkey and barn owl displayed a laminar arrangement. At the microstructural level, adult bone was very similar across taxa in having small, elliptical or compressed osteocyte lacunae.

Avian adult femora display even less variation than the humeri; qualitative characteristics of these elements are summarized in Table 9. Almost all are perfectly circular in cross-section (Fig. 16), with the exception of one of the adult quail. In nearly all taxa, the ACT of the femur is less than that of the humerus (Table 4). The thickest femoral
**Figure 10** Close-up photographs of microanatomical details in the humeri of altricial birds. Sections from youngest individuals with ossified diaphyses shown in the top row, and sections from adults in the bottom row. (A) Anna’s hummingbird pin-feathered chick (MVZ190799); (B) green-cheeked conure neonate (MVZ190895); (C) Western scrub jay neonate (MVZ190927), arrow indicates endosteal bone that separated from the periosteal bone at a Kastschenko line; (D) Anna’s hummingbird adult (MVZ190807); (E) green-cheeked conure adult (MVZ190917); (F) house finch adult (MVZ190993). In all images the periosteal surface is oriented to the upper left and the endosteal to the lower right. Scale bar = 100 µm.

**Table 7** Qualitative comparisons among neonate femora.

| Taxon                     | Cross-sectional shape | Bone composition & porosity                                                                 | Primary porosity orientation | Other notable features                                                                 |
|--------------------------|-----------------------|---------------------------------------------------------------------------------------------|------------------------------|----------------------------------------------------------------------------------------|
| California Quail         | circular              | moderate porosity                                                                          | longitudinal                 | uniform thickness with evenly-distributed vascular canals                               |
| (Callipepla californica) |                       |                                               |                              |                                                                                        |
| Wild Turkey              | circular              | moderate-low porosity, smaller canals than in humerus                                        | longitudinal                 | some incipient primary osteons already present; slightly uneven cortical thickness w/ lower porosity in thin regions |
| (Meleagris gallopavo)    |                       |                                               |                              |                                                                                        |
| Western Gull             | circular              | high porosity, but woven trabeculae thick and well-developed                                | longitudinal/irregular       | some variation in cortical thickness, canals larger in thicker regions                   |
| (Larus occidentalis)     |                       |                                               |                              |                                                                                        |
| American Kestrel         | circular              | high porosity; woven trabeculae very thin                                                  | longitudinal/irregular       | highly asymmetrical cortical thickness with larger canals in thicker regions             |
| (Falco sparverius)       |                       |                                               |                              |                                                                                        |
| Green-cheeked conure     | circular              | high porosity, but trabeculae thick and well-developed                                      | all orientations             | cortex not of uniform thickness; canals larger in thicker regions                        |
| (Pyrrhura molinae)       |                       |                                               |                              |                                                                                        |
| Western Scrub-Jay        | circular              | high porosity; very thin trabeculae                                                         | all orientations             | solid endosteal circlet of bone present; highly asymmetrical cortical thickness         |
| (Aphelocoma californica) |                       |                                               |                              |                                                                                        |
| House Finch              | circular              | high porosity; very thin trabeculae                                                          | longitudinal/irregular       | solid circlet of endosteal bone present                                                  |
| (Haemorhous mexicanus)   |                       |                                               |                              |                                                                                        |

**Note:**

Note that vascular density was high almost ubiquitously among these individuals, therefore porosity is compared among neonates. Results are shown for taxa for which neonate specimens were available, and in which the femur was ossified and viable for this study (the femora of the mourning dove and Anna’s hummingbird were not included for the latter reason). Developmental mode is indicated by color number superscript: orange¹, precocial; yellow², semi-precocial; green³, semi-altricial; purple⁴, altricial.

---

Atterholt and Woodward (2021), *PeerJ*, DOI 10.7717/peerj.12160
Figure 11 Close-up photographs of microanatomical details in the femora of precocial birds, sections from neonates shown in the top row and from adults in the bottom row. (A) Wild turkey neonate (MVZ190764); (B) California quail neonate (MVZ190751); (C) wild turkey adult (MVZ190763); (D) California quail adult (MVZ190762), showing a region of woven bone with incipient primary osteons (white arrows) and simple vascular canals (black arrows). In all images the periosteal surface is oriented to the upper left and the endosteal to the lower right. Scale bar = 100 µm.

Figure 12 Close-up photographs of microanatomical details in the femora of semi-precocial birds, sections from neonates shown in the top row and from adults in the bottom row. (A) Western gull neonate (JAA64); (B) Western gull neonate (MVZ190822); (C) Western gull adult (MVZ190829); (D) Western gull adult (MVZ190831). In all images the periosteal surface is oriented to the upper left and the endosteal to the lower right. Scale bar = 100 µm.
cortices are found in the turkey (1240.14 µm) and the great-horned owl (736.66 µm). The thinnest is in the hummingbird (70.55 µm). In terms of RCT, the barn owl has the greatest cortical thickness in the femur (0.30). The RCT of all other taxa falls in a range from 0.18–28; notably, five of these taxa (representing a wide range of body sizes and flight styles) all have an average adult femoral RCT of 0.18 (Table 5).

Vascular channels were very small in size but present in large numbers in most taxa, as in the humerus. A low vascular density was only observed in the house finch, and the hummingbird femur is avascular. Notably, these are the two smallest adults included in this study. These observations are consistent with results reported by Cubo et al. (2014), who discovered a significant correlation between vascular density and body size in a number of adult birds and lepidosaurs. This is a reminder that, in mature bone, metabolic requirements and method of oxygen and nutrient receipt are related to size of an element, which should be considered when making paleohistological interpretations. A predominantly laminar orientation of vascular channels was observed only in the wild turkey and one barn owl specimen. In all other taxa, canals were arranged longitudinally, in an anastomosing network, or some variation between these two.

Overall, there is a fair amount of variance in the humeral and femoral histology of adults, with cortices ranging from predominantly fibrolamellar to avascular parallel-fibered bone (Figs. 6, 11C & 11D; Figs. 7,12C, 12D; Figs. 8, 13B–13F; Figs. 9 & 14B; Figs. 10D–10F; Figs. 15E–15G). The wild turkey, Western gull, red-tailed hawk, barn owl, and great-horned owl have humeral cortices composed of a thick layer of fibrolamellar...
bone between a clear OCL and ICL. In California quail, American kestrels, white-tailed kites, and mourning doves, the cortex is mainly composed of a woven or parallel-fibered matrix with regional fibrolamellar bone or incipient primary osteons. In the house finch and green-cheeked conure the humerus is composed of a thin, middle layer of woven matrix with simple vascular canals, between a strongly parallel-fibered OCL and ICL.

Finally, in Anna’s hummingbird the cortex of the humerus is nearly avascular and composed entirely of parallel-fibered bone.

All individuals classified here as “adults” did indeed have an OCL in the humerus and femur. However, in some taxa an OCL appeared in some individuals assigned to other growth stages (summarized in Table 1). In mourning doves, an incipient OCL is present at the subadult stage in both the humerus and femur (Figs. S18F & S18G, S20G). In green-cheeked conures, an incipient OCL is visible by the fledgling stage in the humerus, and the pre-fledging stage in the femur (Fig. S32D). The femur of the fledgling has a thick, prominent OCL, and has achieved a cross-sectional size nearly equivalent to that of the adult. In Western scrub jays, the humerus of the subadult individual shows signs of early OCL formation (Fig. S34G), and the femur of the fledgling and subadult
stages both exhibit incipient OCLs (Figs. S36F & S36G). In house finches, the OCL first starts to develop in the fledgling stage in the humerus (Fig. S38D), and the pre-fledgling stage in the femur (Figs. S40C & S40D). By the fledgling stage, the OCL is thick and clearly developed, and cross-sectional size of this element is within the range of the adult (in fact, slightly larger in the two individuals compared here). In the great-horned owl, OCL formation begins in the fledgling chick in the humerus and femur (though it is in very early stages of development; Figs. S42D and S44C). In barn owls, an incipient OCL is also present in the femur as early as the fledgling stage (Fig. S48F), and by the subadult stage appears well-developed and has achieved the same thickness as that of the adult stage. The difference, instead, is lack of an ICL in the subadult individual. The humerus of the barn owl also has an incipient OCL by the subadult stage (Fig. S46G).

Anna’s hummingbird is the most outstanding exception to an OCL correlating with “adulthood” based on plumage. An OCL was present in the humerus as early as the fledgling growth stage (Fig. S14B), and persists through the subadult stage. In the adult, the OCL is no longer visible as a distinct layer (Fig. 10D). Rather, it appears to have widened so much that, in combination with endosteal resorption, it now comprises nearly all the bone of the cortex. Furthermore, the size of the fledgling humerus is comparable to that of the adult.

An ICL was also present in the femur and humerus of nearly all adult birds in this dataset. This feature often appeared as a very distinctive layer of highly organized tissue, parallel-fibered bone bordering on lamellar in some instances. It appeared to be only in early stages of formation in the adult hummingbird (Figs. 10D and 15E), and was entirely absent in femur of the adult house finch, which instead had a scalloped endosteal

---

**Figure 15** Close-up photographs of microanatomical details in the femora of altricial birds. (A) Anna’s hummingbird pin-feathered chick (MVZ190799); (B) green-cheeked conure neonate (MVZ190895); (C) house finch neonate (MVZ190969); (D) Western scrub jay neonate (MVZ190927); (E) Anna’s hummingbird adult (MVZ190807); (F) green-cheeked conure adult (MVZ190917); (G) house finch adult (MVZ190993). (H) house finch pin-feathered chick showing inner circle of “floating” bone interpreted as evidence of a Kastschenko line. In all images the periosteal surface is oriented to the upper left and the endosteal to the lower right. Scale bar = 100 µm. Full-size [DOI: 10.7717/peerj.12160/fig-15]
Figure 16 Diaphyseal cross-sections of all adults included in this study, showing general cross-sectional geometry and variation in cortical thickness. For each pair, the humeral section is on the left and the femoral on the right. In all images cranial is up and lateral is left. Sections are grouped by developmental mode. Precocial (orange): (A) Wild turkey (*Meleagris gallopavo*; MVZ190763), scale bar = 1,000 µm. (B) California quail (*Callipepla californica*; MVZ190762), scale bar = 500 µm. Semi-precocial (yellow): (C), Western gull (*Larus occidentalis*; MVZ190831), scale bar = 1,000 µm. Semi-altricial 1 (green): (D) great-horned owl (*Bubo virginianus*; MVZ190883), scale bar = 1,000 µm; (E) white-tailed kite (*Elanus leucurus*; MVZ190861), scale bar = 500 µm; (F) barn owl (*Tyto alba*; MVZ190877), scale bar = 500 µm; (G) American kestrel (*Falco sparverius*; MVZ190892), scale bar = 500 µm; (H) red-tailed hawk (*Buteo jamaicensis*; MVZ190855), scale bar = 1,000 µm. Semi-altricial 2 (blue): (I) mourning dove (*Zenaida macroura*; MVZ190775), scale bar = 500 µm. Altricial (purple): (J) green-cheeked conure (*Pyrrhura molinae*; MVZ190917), scale bar = 500 µm; (K) house finch (*Haemorhous mexicana*; MVZ190993), scale bar = 250 µm; (L) Anna’s hummingbird (*Calypte anna*; MVZ190807), scale bar = 100 µm. For cross-sections of all ages, see figures in Supplemental Material.
margin indicative of active resorption (Fig. 15G). Such variation may be linked to differing stages of the life-cycle in which endosteal resorption and deposition of bone are activated. Additionally, of the two adult barn owls sectioned in this study, one lacked an ICL in the humerus, indicating that tissue growth and/or resorption along the endosteal margin was still active, and suggesting this individual had either not yet achieved the same level of cortical maturity as the other, or had resorbed the endosteal surface for mineral mobilization (however, both individuals have an ICL and OCL in the femora).

Relatively little remodeling in humeri and femora has been described in previous studies of the limb bones of adult birds (Enlow & Brown, 1957; Currey, 2003; Simons & O’connor, Table 8 Qualitative comparisons among adult humeri.

| Taxon               | Cross-sectional shape | Presence of ICL and OCL          | Composition of middle cortical layer                                      | Relative vascular density | Primary vascular orientation |
|---------------------|-----------------------|----------------------------------|--------------------------------------------------------------------------|---------------------------|-----------------------------|
| California Quail1   | circular to ovate     | yes; OCL thicker (~1/3 of cortex) but ICL more distinct | woven/incipient fibrolamellar                                             | moderate                  | longitudinal                 |
| (Callipepla californica) |                     |                                  |                                                                           |                           |                             |
| Wild Turkey1        | rounded triangle      | yes; ICL and OCL more distinct   | fibrolamellar                                                             | high                      | circumferential/ plexiform   |
| (Meleagris gallopavo) |                       |                                  |                                                                           |                           |                             |
| Western Gull2       | circular              | yes; ICL and OCL more distinct   | fibrolamellar                                                             | high                      | reticular                   |
| (Larus occidentalis) |                       |                                  |                                                                           |                           |                             |
| American Kestrel3   | circular              | yes; equal thickness, ICL more distinct | woven w/simple canals and regional fibrolamellar                         | moderate                  | longitudinal/ reticular      |
| (Falco sparverius)  |                       |                                  |                                                                           |                           |                             |
| Red-tailed Hawk3    | circular              | yes; ICL and OCL more distinct   | fibrolamellar                                                             | high                      | longitudinal                 |
| (Buteo jamaicensis) |                       |                                  |                                                                           |                           |                             |
| White-tailed Kite3  | circular              | yes; OCL, ICL more distinct      | fibrolamellar bone with regional woven bone w/ simple canals             | moderate                  | longitudinal                 |
| (Elanus leucurus)   |                       |                                  |                                                                           |                           |                             |
| Barn Owl3           | circular              | yes; equal thickness, ICL more distinct | fibrolamellar                                                             | high                      | reticular/ circumferential   |
| (Tyto alba)         |                       |                                  |                                                                           |                           |                             |
| Great-horned Owl3   | circular              | yes; ICL and OCL more distinct   | fibrolamellar                                                             | high                      | longitudinal                 |
| (Bubo virginianus)  |                       |                                  |                                                                           |                           |                             |
| Mourning Dove4      | ovate (main medullary cavity) | yes; ICL incipient, OCL 1/3-1/2 cortex | weakly woven with primary osteons (middle layer is of equal thickness to OCL in some areas) | moderate                  | longitudinal                 |
| (Zenaida macroura)  |                       |                                  |                                                                           |                           |                             |
| Anna’s Hummingbird5 | ovate (main medullary cavity) | yes; OCL dominates cortex | none                                                                      | avascular                 | —                            |
| (Calypte anna)      |                       |                                  |                                                                           |                           |                             |
| Green-cheeked conure5 | circular              | yes (but ICL thin/ irregular); OCL thicker | woven bone with primary vascular canals; patches of fibrolamellar          | moderate-low              | longitudinal                 |
| (Pyrrhura molinae)  |                       |                                  |                                                                           |                           |                             |
| House Finch5        | circular              | yes; OCL is thicker, ~1/3-1/2 of cortex | woven bone with primary vascular canals; patches of fibrolamellar          | moderate-low              | longitudinal                 |
| (Haemorhous mexicanus) |                      |                                  |                                                                           |                           |                             |

Note: Vascular density in adults was ubiquitously lower than in younger growth stages, thus relative vascular density among adults only is reported here. “Middle cortical layer” refers to the portion of the cortex located between the OCL and the ICL. Descriptions of the ICL as “more distinct” refer to instances in which the ICL is more advanced in development than the OCL, and to the very high level of organization of tissue in this layer (which appears strongly lamellated and has a distinct, high birefringence). Developmental mode is indicated by color number superscript: orange1, precocial; yellow2, semi-precocial; green3, semi-altricial 1; blue4, semi-altricial 2; purple5, altricial.
with the exception of extremely dense Haversian systems present in penguin bones (related to the aquatic lifestyle of these animals (Meister, 1962) and paleognaths (Bourdon et al., 2009; de Ricqlès et al., 2016; Chinsamy et al., 2020a)). With respect to the mid-shaft of the humerus and femur, the results of this investigation concur with these studies, and only rare, small, localized areas of osteonal remodeling were observed (e.g., in the wild turkey humerus).

## DISCUSSION

Several osteohistological attributes characterize Aves as a whole, both in terms of general ontogenetic trends and similarities among specific growth stages. In all neonates, the humerus and femur are composed of a disorganized matrix of woven bone with many large, irregular vascular channels, and a high osteocyte density (with round, large osteocyte

---

### Table 9 Qualitative comparisons among adult femora.

| Taxon                                        | Cross-sectional shape | ICL and OCL present | Composition of Middle Cortical Layer | Relative vascular density | Primary vascular orientation |
|----------------------------------------------|-----------------------|---------------------|-------------------------------------|--------------------------|-----------------------------|
| California Quail (Callipepla californica)     | circular to ovate     | yes; OCL thicker (~1/4-1/3 cortex) but ICL more distinct | woven with simple canals/regional fibrolamellar | moderate                 | longitudinal                |
| Wild Turkey (Meleagris gallopavo)            | circular to ovate     | yes; ICL thicker and more distinct | fibrolamellar                       | high                     | circumferential/plexiform    |
| Western Gull (Larus occidentalis)             | circular              | yes; equal thickness, ICL more distinct | fibrolamellar                       | high                     | reticular                   |
| American Kestrel (Falco sparverius)          | circular              | yes; equal thickness, ICL more distinct | parallel-fibered with regional fibrolamellar | moderate                 | longitudinal                |
| Red-tailed Hawk (Buteo jamaicensis)          | circular              | yes; ICL thicker and more distinct | fibrolamellar                       | high                     | reticular-longitudinal      |
| White-tailed Kite (Elanus leucurus)           | circular              | yes; equal, ICL more distinct | fibrolamellar bone with regional woven bone w/simple canals | moderate                 | longitudinal                |
| Barn Owl (Tyto alba)                         | circular              | yes; equal thickness, ICL more distinct | fibrolamellar                       | high                     | reticular-longitudinal      |
| Great-horned Owl (Bubo virginianus)          | circular              | yes; ICL thicker and more distinct | fibrolamellar                       | high                     | longitudinal                |
| Mourning Dove (Zenaida macroura)              | circular              | yes; OCL thicker (comprises almost entirety of cortex) | small patches of weakly woven with sparse simple canals | low                      | longitudinal                |
| Anna’s Hummingbird (Calypte anna)            | circular              | yes; ICL incipient(?), OCL dominates cortex | none                               | avascular                | —                           |
| Green-cheeked conure (Pyrrhura molinae)       | circular              | yes; OCL thicker, about 1/3 of cortex | woven bone with simple vascular canals; patches of fibrolamellar bone | low                     | longitudinal                |
| House Finch (Haemorhous mexicanus)           | circular              | ICL absent; OCL ~1/2 of cortex | parallel-fibered with regional woven bone | low                     | longitudinal                |

**Note:**
Vascular density in adults was ubiquitously lower than in younger growth stages, thus relative density among adults only is reported here. "Middle cortical layer" refers to the portion of the cortex located between the OCL and the ICL. Descriptions of the ICL as “more distinct” refer to the very high level of organization of tissue in this layer, which appears strongly lamellated and has a high birefringence. Developmental mode is indicated by color number superscript: orange 1, precocial; yellow 2, semi-precocial; green 3, semi-altricial 1; blue 4, semi-altricial 2; purple 5, altricial.
lacunae) (Figs. 6–15). In many hatchling chicks, it was difficult to identify a primary orientation of vascular canals because their arrangement appeared so haphazard. This is unsurprising, and corroborates with both previous reports on bone histology of avian chicks (Castanet et al., 2000; de Margerie, Cubo & Castanet, 2002; de Margerie et al., 2004), and data available on high growth rates in birds (Starck, 1989, Starck, 1993, Starck, 1994; Starck & Ricklefs, 1998) as bone with such a lack of organization is known to be very fast-growing. Additionally, nearly all taxa hatch with a femoral cortex that is thicker than that of the humerus (as an absolute measure), but this relationship becomes inverted by the adult stage, when the bone of the humerus is almost invariably thicker than the femur. Absolute cortical thickness generally increases through ontogeny, reaching highest values in adults. Conversely, relative cortical thickness often decreases between the neonate and adult stages. Below we discuss further patterns related to phylogeny, biomechanics, and ontogeny.

**Phylogenetic patterns**

In this study, only RCT of the neonate femur has significant phylogenetic signal ($K = 1.235; P = 0.043$). Phylogenetic signal has been reported in other histological characteristics of adult birds by Houde & Olson (1986) and Houde (1987), in ostriches and the extinct bird Hesperornis. Other examples include signal in the density of Haversian bone (Ponton et al., 2007); relative cortical thickness of sauropsid femora, including a diverse sampling of extant birds (Cubo et al., 2005); and a wide range of features (e.g., vascular density and vascular orientation) across paleognathous birds (Legendre et al., 2014). It is perhaps surprising, then, that significant phylogenetic signal was not detected in more traits in the present investigation. However, we note that this may be because low sample sizes, such as in this study, predispose these tests to type II error, and others urge the use of large datasets with thorough phylogenetic coverage to rigorously support a hypothesis of phylogenetic signal (Legendre et al., 2014). Future studies incorporating more taxa may yield different results.

Some features observed appear to be clade-specific. Very young chicks of both passerine taxa included in this study (the house finch and scrub-jay) have a unique characteristic in the humerus and femur: a thin, solid ring of bone that lines the endosteal edge of the woven bone comprising the cortex, making the medullary cavity very distinct (Figs. 5H, 10C, 15H). This circlet of bone appears to detach and ‘float’ into the medullary cavity later in development, where it is presumably resorbed as it appears in no later stages of ontogeny. We interpret this as evidence of a Kastschenko line, a thin layer of tissue that is a remnant of the cartilaginous precursor that forms earlier in bone development, located between the growing periosteal and endosteal bone (Kastschenko, 1880; Francillon, 1980; Castanet & Smirina, 1990). In this case, the space observed between the inner circlet of bone and the rest of the cortex is all that remains of the osteoid layer. During the specimen preparation process, periosteal and endosteal layers have separated along this region of thin, delicate tissue, forming a space between the two layers of bone. This is a very common osteohistological feature in frogs and salamanders (e.g., Rozenblut & Ogielska, 2005; Song et al., 2017), but is relatively rare in other groups. To date, the only amniotes in
which a Kastschenko line has been identified are elasmosaurid and polycotylid plesiosaurs (O’gorman, Talevi & Fernández, 2016; O’Keefe et al., 2019), early-diverging sauropodomorphs (Cerda, Pol & Chinsamy, 2014), a titanosaur sauropod (González et al., 2020), and theropods (de Ricqlès et al., 2001). This study therefore presents the first evidence of this structure in Avialae. Furthermore, both taxa in which it was identified belong to the clade Passeriformes, suggesting that the Kastschenko line is indicative of a growth strategy specific to members of this clade, however, sampling of additional passerine and non-passerine birds will be necessary to confirm the biological reality of this pattern.

**Histological variation**

RCT is remarkably similar across adults of all taxa. This is particularly the case for the humerus, likely because the bones of flighted taxa thin out to a relatively restricted range of RCT values to simultaneously support the strains of active flight and reduce body weight. Both RCT and ACT are some of the most highly variable traits in growing chicks. Across taxa, birds of the same growth stage often exhibited very different cortical thicknesses, and within a given element great asymmetry in width of the cortical bone was also frequently observed, indicating that growth stage based on external, gross anatomical features (such as plumage) are unsurprisingly not “equivalent” between taxa in terms of bone development. Alternatively, it may reflect varying mechanical forces on the bone through growth, though the lack of pattern for a given element among individuals of the same growth stage and taxon suggests that this characteristic is non-adaptive, or that a greater sample size is needed for the adaptive pattern to become apparent. These differences among birds of the same growth stage later become highly reduced as elements converge to similar thicknesses among adults of the same taxon. This was observed across clades and implies that selective forces affecting cortical thickness may be less strong at earlier stages of development. It also suggests that environmental factors, differences in parental care, and quality/amount of food available from clutch to clutch may have a strong effect on the thickness of bone in chicks.

Overall, regional differences consistently related to anatomical direction within a section (that is, differences among cranial, caudal, medial, and lateral quadrants) were minimal across growth stages. Other studies have also reported a lack of regional histovariability. For instance, Simons & O’connor (2012) examined laminarity in adult birds of a range of flight styles, and found no significant variation among the four cortical quadrants. Here, the only major regional variation observed within a given bone cross-section was in the seemingly arbitrary thickness of cortical bone discussed above. Most of this variation was seen in the humerus, and primarily at younger developmental stages where it was very common.

**Mechanical signal**

Almost invariably, a trend of decreasing RCT is observed through post-natal ontogeny, with adults having, on average, lower RCT than chicks (Table 5). However, it must be noted that this trend is not always a constant, unidirectional decrease, and that there are some exceptions to this pattern. In the turkey and quail, adult RCT of the femur is greater
than in the neonate. Both taxa are precocial, and both are members of Galliformes, an early-diverging clade, so this characteristic may be related to life history or inherited from early common ancestors. Alternatively, this could reflect functional demands; both birds rely primarily on the pelvic limb for locomoting and are not strong fliers.

When considering more complete growth series (that is, not only the end points), other trends in cortical growth become apparent. In many taxa, there is an increase in RCT from the neonate to pin-feathered stage followed by a decrease, and an increase from pre-fledgling to fledgling stage followed by a decrease. Similarly, some taxa also undergo an increase in ACT at the pin-feathered or fledgling stage, followed by a decrease in later ontogenetic stages. However, cortical thickness within and between individual growth stages exhibited such great variability that it is impossible to comment on the biological reality of such trends without a greater sample size. Nonetheless, this does suggest a general trend of highly dynamic osteoblast and osteoclast activity, with rates of endosteal resorption and periosteal deposition fluctuating through ontogeny. Patterns of periosteal growth in birds are therefore complex, and the relationship between rates of periosteal deposition and endosteal resorption is ever-shifting.

Some trends in cortical thickness are clearly explicable. In adults, ACT is typically at a peak because individuals have reached maximum body size and thus have larger bones than earlier stages. Also, adults tend to have measures of lowest RCT because thin-walled bones are an adaptation to flight. However, proximate functional causes of other ontogenetic changes in the cortical thickness of the humerus and femur are more ambiguous.

A previous study suggests (Carrier & Leon, 1990) that the structural weaknesses of the immature bone of young birds may be compensated for by a greater cortical thickness. Time of fledging, when a bird begins to fly, places especially high demand on the wing bones, potentially explaining the secondary increase in relative cortical thickness often observed in fledgling chicks. Carrier & Auriemma (1992) note that time of fledging is limited by relative length of certain limb elements (birds with proportionally longer humeri and ulnae take longer to reach the fledgling stage). These authors infer that longitudinal growth may be a developmental constraint imposing a minimum duration to the fledgling stage. This critical stage of avian ontogeny is strongly affected by the dynamics of bone growth, and diametric thickening of the bone may be a way to offset the limitations of longitudinal growth and/or weaknesses of the immature tissue.

Other histological features have also been suggested as adaptations to locomotion, and to resisting the various strains that it imposes. de Margerie, Cubo & Castanet (2002) and de Margerie (2006) proposed that circular cross-sectional shape, a thin cortical wall, collagen fibers arranged spirally around the bone shaft at a 45° angle, and bone laminarity (i.e., many circumferential vascular canals) are all functional adaptations for torsion resistance. They further report that degree of laminarity varies across skeletal elements, and is highest in the humerus, radius, ulna, and femur.

If true, we expect laminar vascularity will be found in the major pectoral elements across volant avian taxa. The range of taxa explored in this study allows a test of this hypothesis, which does not appear to hold true across the avian phylogeny. Laminar vascularity was in
fact quite rare in the samples studied here. In a few adult birds, laminar bone did dominate the humerus (the wild turkey, one of the barn owls, and weak lamination in some regions of the red-tailed hawk cortex), but it was absent in the vast majority of specimens. Certainly, a larger sample size and additional sectioning of each element would provide a more rigorous test, but preliminary results indicate that laminar vascularity does not dominate the humerus in a majority, or even many, birds. This corroborates recent findings in homing pigeons (McGuire et al., 2020) and bats (Lee & Simons, 2015).

Of particular note, McGuire et al. (2020) observed a higher degree in laminarity in younger individuals, with vascular orientation becoming increasingly longitudinal with maturity. Though additional quantitative studies of data presented here are necessary, current qualitative observations suggest this ontogenetic pattern is likely common across Aves. Interestingly, the femur of the wild turkey was also predominantly laminar, as were the femora of the older ostrich chicks. These results are in keeping with the idea that this orientation of vascular canals is an adaptation to torsion, because the avian femur, which is obliquely oriented in the cranio-caudal plane, is subject to high torsion, particularly in large terrestrial birds such as the turkey, ostrich, and emu (de Margerie, Cubo & Castanet, 2002; de Margerie et al., 2005; Kuehn et al., 2019).

Developmental mode & onset of bone growth

At the start of this investigation, we hypothesized that a histological signal of developmental mode would be present in the form of differing cortical thickness in pectoral versus pelvic limb elements. Precocial chicks locomote earlier, and with the pelvic limb, and should have an absolutely thicker and more mature femoral cortex than humeral; in contrast, altricial chicks will ultimately locomote primarily with the pectoral limb, and should have an absolutely thicker and more mature humeral cortex relative to that of the femur (or at least of equal maturity and thickness, if delayed locomotion is not enough to select for a functional difference in neonate limb bones). However, the data here do not bear out these predictions, instead indicating that the functional relationship between cortical thickness and maturity of neonates is more ambiguous.

With respect to thickness of the cortical bone, neonates of some taxa do fit the model described above. The femur of the semi-precocial Western gull chick is thicker than the humerus (by absolute and relative measures), matching Carrier & Leon’s (1990) observations in the California gull. The precocial California quail neonate also has a thicker femoral than humeral cortex by absolute measure. However, in the wild turkey chick, the humerus is absolutely thicker than the femur, and in the altricial Western scrub-jay and green-cheeked conure, the femur is absolutely and relatively thicker than the humerus. Additionally, the highest values of relative cortical thickness of both elements are predominantly seen in the altricial birds. Instead, precocial birds appear to compensate for greater functional demands on the pelvic limb simply by having more mature bone in the femur than the humerus (though this is not unique to precocial chicks, as discussed in greater detail below).

The idea of cortical thickness compensating for a lack of structural strength is born out by three of the four precocial taxa represented in this dataset, but this does not fully explain
why the pattern of the femoral neonate cortex being thicker than the humeral does not appear in the precocial wild turkey, or why it does appear in altricial taxa. These differences in cortical thickness may not be a functional adaptation at all, instead representing a phylogenetic constraint as implied by how common this trait is in the diverse array of birds sampled in this study. On the other hand, Carril, Tambussi & Rasskin-Gutman, 2021 purport that the early ossification of the pelvic limb relative to the pectoral limb in altricial monk parakeet chicks is adaptive because chicks engage in moderate walking movements while still in the nest and before fledging (Aramburú, 1997). If such behavior is also present in other altricial chicks, this would potentially account for the patterns of bone maturity observed here. Future studies of locomotory behavior in chicks may help reveal whether or not it is adaptive to have a thicker femoral cortex than humeral at hatching.

Ultimately, it is striking that no distinct pattern of differences in either measure of cortical thickness was observed across the altricial-precocial spectrum. RCT of precocial and semi-altricial chicks falls within a similar range, as does RCT of semi-precocial and altricial chicks (Fig. 5B). Results for ACT across developmental modes are similar, though the differences not as great (Fig. 5A). Whatever contributes to variation in cortical thickness, it does not appear related to place along the altricial-precocial spectrum.

In contrast to cortical thickness, relative maturity of the bone at hatching does appear related to developmental mode. In precocial taxa, the bone of the humerus and femur was more mature (as inferred from the relatively smaller vascular channels and more advanced state of fibrolamellar development), while the most immature bone was observed in more altricial neonates (with very thin cortices, large vascular openings, thin trabeculae of woven bone, and in some cases totally cartilaginous elements at hatching). Again, this very likely reflects differences in functional demands and growth rates. Starck & Ricklefs (1998) have reported higher overall rates of growth at the altricial end of the developmental spectrum and lower rates at the precocial end. Ricklefs (1979) specifically identified an inverse relationship between growth rate and maturity of a tissue. Kirkwood et al. (1989) demonstrate that an increase in tibiotarsal length occurs at an average rate three times greater in altricial hatchlings than in precocial neonates. The more immature bone of altricial chicks indicates a relatively higher growth rate of the humerus and the femur than in precocial chicks. The more mature bone of precocial chicks is evidence of the higher functional demands placed on neonates that must locomote, find food (in some cases), and maintain at least some degree of independence very soon after hatching.

However, when comparing relative maturity of the femur and humerus of the same individual, another pattern becomes apparent: in all of the birds in this dataset for which a neonate specimen was available, the tissue of femur at hatching is comparatively more mature than the humerus, having relatively smaller vascular channels and a more organized bone matrix. In the case of the house finch, this difference is manifest as only the femur being ossified enough to be included in this study, while the humerus was still entirely cartilaginous. At a proximate level, this difference in maturity is indicative of varying growth rates between these two major limb bones. Ricklefs (1979) reported that degree of functional maturity of a tissue is inversely related to rate of growth, a hypothesis that has been substantiated by subsequent studies (e.g., Montes et al., 2005). Others
describe an inverse, negative relationship between high growth rates and strength of ossified tissues (Carrier & Leon, 1990; de Margerie et al., 2004), indicating a functional constraint on the relationship between growth rate and maturity, an idea further supported by differences in bone maturity of altricial and precocial neonates observed here. Therefore, the more immature bone of the humerus suggests that this element grows at a faster rate than the femur. It is likely that the difference in bone maturity between the humerus and femur is partly due to early functional demands placed on the femora of precocial taxa (both bones are more mature in precocial than altricial hatchlings); however, the presence of this characteristic in all birds across the altricial-precocial spectrum indicates it may also be the result of ontogenetic (wherein possible phenotypes are constrained by the developmental program) or phylogenetic channeling.

Precocial and semi-precocial birds may have, through selection, exapted and exaggerated this pre-existing characteristic to facilitate the chicks’ lifestyle, but the data presented here suggest this difference in maturity may initially (and in other developmental modes) represent a non-adaptive trait. Alternatively, as suggested by Carril, Tambussi & Rasskin-Gutman (2021), this may in fact be adaptive in even altricial chicks contingent upon the degree of in-nest locomotion that they exhibit. In either case, it is notable that a growing body of histological studies of Mesozoic birds and non-avialan theropods hints that a greater functional maturity of the hindlimb relative to the forelimb may have evolutionary origins even deeper than crown group birds (e.g., Prondvai et al., 2018).

Ultimately, the only consistent histological patterns related to developmental mode is the maturity of femoral and humeral bone together at the time of hatching (precocial chicks have more mature bone than altricial). However, because this characteristic is continuous and not discrete, it is difficult to define a precise way of predicting developmental mode from qualitative assessment of neonate histology alone, perhaps outside of the two most extreme ends of the altricial-precocial spectrum. The bone of precocial taxa appears more mature at hatching than that of altricial chicks, however, altricial chicks pass through stages later in their long bone ontogeny where they resemble the bone of precocial chicks, suggesting that this difference results from changes to pre-natal onset of bone growth. More altricial chicks have accomplished less growth in the egg, but their post-natal growth trajectory is still very similar to that of precocial chicks, only with a different post-hatching ‘starting point.’ That is, altricial chicks will undergo ontogenetic stages where their bone resembles that of precocial chicks, but have to first experience early stages of post-natal growth that have already been achieved by more precocial chicks at the time of hatching.

**Body size and offset of bone growth**

The presence of the OCL is conventionally interpreted as a signal of reduction or cessation of growth, an indicator that skeletal maturity has been achieved (Chinsamy & Elzanowski, 2001; de Ricqlès et al., 2003; Chinsamy et al., 2013). In this study, where the “adult” growth stage was based on plumage and body size, we were able to test the relationship between “adult” external morphology and skeletal maturity.
An OCL was observed in both the humerus and femur of all taxa in this dataset for which an adult sample was available. However, in almost all instances, the OCL was also observed at earlier growth stages in both elements. In eight of the nine taxa for which mid-to late-growth stages were available, the OCL first appears at the pre-fledgling (house finch), fledgling (white-tailed kite, barn owl, great-horned owl, Anna’s hummingbird, green-cheeked conure, Western scrub jay) or subadult (mourning dove) stage (Table 1). In nearly all first appearances, the OCL is incipient, comprising a very thin portion of the periosteal cortex that is parallel-fibered and avascular. However, in some instances a fully-developed, ‘mature’ OCL was present before the adult stage. Anna’s hummingbird represents the most drastic example of this, with a very thick OCL in the cortex of the fledgling humerus. In all such cases, the diametric size of the humerus or femur is comparable to that of the adult. Therefore, these examples suggest that appositional growth ceases at a relatively early developmental stage, while the bone continues to change and mature with a parallel-fibered matrix becoming more prevalent than a woven matrix or chondroid tissue, and endosteal resorption still active. Not only is an OCL decisively not a sign of senescence (as also argued by Woodward et al., 2020), but the presence of an OCL does not always even imply that an individual has achieved cortical maturity and does not necessarily correlate with the presence of “adult” gross morphology, further underscoring the complexity of defining “adulthood.” In short, skeletal maturity cannot be equated with the cessation of appositional growth.

Additionally, the microanatomical features of the middle cortical layer (adjacent to the OCL) of “adults” in this study was distinct relative to all earlier growth stages. Prior studies describe the cortical bone of modern birds as generally composed of a layer of highly-vascularized fibrolamellar bone (de Ricqlès, 1975; de Ricqlès et al., 1991; Castanet et al., 1996), though de Margerie, Cubo & Castanet (2002) do describe avascular to poorly-vascularized parallel-fibered bone in the humerus of a mallard. A cortex mainly composed of fibrolamellar bone was observed in many skeletally mature individuals in this dataset, but not all. Specifically, this best describes the larger-bodied birds (the wild turkey, Western gull, red-tailed hawk, barn owl, and great-horned owl). The adult histology of these taxa corroborates the known description of crown-group birds growing very quickly until adult size is achieved, then rapidly slowing down, resulting in the familiar pattern of a fibrolamellar cortex (reflecting early fast growth) and a parallel-fibered OCL and ICL (reflecting the later short stage of slower growth).

However, this pattern becomes less clear as body size decreases. In birds of medium body-size (quail, American kestrels, white-tailed kites, and mourning doves), the middle cortical layer is composed of a woven or parallel-fibered matrix with regional fibrolamellar bone or incipient primary osteons. The fast growth of earlier developmental stages is not as clearly reflected in these adults. A true fibrolamellar complex never quite fully forms in these birds and seems to only ever achieve early stages of development with, at most, regional and/or incipient fibrolamellar bone present.

In small birds (house finch and green-cheeked conure), the long bones are composed of a thin, middle layer of a woven matrix with simple vascular canals, between a strongly parallel-fibered OCL and ICL. Vascularity is also reduced, with relatively sparse
longitudinal canals present. Finally, in the smallest bird in this dataset (Anna's hummingbird), the cortices of the humerus and femur are avascular and composed entirely of parallel-fibered bone. In short, the spectrum of body sizes parallels the spectrum of structural organization of bone, ranging from fibrolamellar to parallel-fibered with a variety of intermediates between the two. In smaller birds, the OCL appears to grow so thick that, combined with endosteal resorption, it ultimately comprises the majority or the entirety of the cortex.

This trend of apparently slower growth in small-bodied taxa that belong to clades including large, fast-growing taxa (e.g., dinosaurs, pterosaurs, and mammals) has also been reported in other groups and is not unique to crown-group birds (Case, 1978; Padian, de Ricqlès & Horner, 2001; Padian, Horner & de Ricqlès, 2004; Lehman & Woodward, 2008). These patterns provide evidence that most birds in this dataset follow an ontogenetic trajectory of very fast growth in early stages (neonate to pre-fledgling), which abruptly slows or even stops at later stages (pre-fledgling to subadult), followed by a period dominated by bone maturation with little new appositional growth. This supports the idea suggested by Padian, Horner & de Ricqlès (2004) that avialans evolved a relatively small body size by truncating the early period of fast growth. Furthermore, this appears to be a growth strategy that has been manipulated and exaggerated repeatedly in the evolution of increasingly smaller birds. Histological traits of small-bodied taxa studied here indicate that the period of fast growth is much shorter than in large-bodied taxa. This pattern is most extreme in Anna’s hummingbird, in which a fully-formed OCL is present by the fledgling stage in the humerus, and the adult cortex is completely parallel-fibered. However, these small birds still ultimately complete their growth fast enough not to lay down any LAGs or other growth marks, as none were observed in the adults of this dataset.

What is notable is that this ontogenetic trend seems entirely driven by body size, and independent of developmental mode; birds of medium to small body size across the altricial-precocial spectrum follow this model. This is not to say that these taxa share similar growth rates, however. It has been well-documented that birds on the precocial end of the spectrum grow at lower rates than those at the altricial end (Ricklefs, 1968; Ricklefs, 1973; Starck, 1989; Starck, 1993; Starck & Ricklefs, 1998). Instead, this seems to result from a manipulation in the timing of off-set of bone growth (a truncated period of fast growth followed by a relatively longer period of slowed growth and bone maturation) in small-bodied birds across the altricial-precocial spectrum. While absolute growth rates of more altricial taxa are undoubtedly higher during the fast growth phase, any taxon that only has to achieve small body size still employs the strategy of truncating this part of the growth trajectory. In parallel to this and as described above, data presented here also suggest that change in the timing of onset of bone growth is related to place along the altricial-precocial spectrum.

**CONCLUSIONS**

We tested hypotheses that cortical thickness is used to compensate for biomechanical weakness of woven bone in the hindlimbs of precocial chicks, and that precocial chicks exclusively have more mature bone in the femur than the humerus at the time of hatching.
No evidence was found that unequivocally supports either prediction. While most precocial taxa in this dataset have thicker femoral cortices than humeral at hatching, this relationship was also observed in most other taxa, and irrespective of place along the altricial-precocial spectrum, the femoral bone of most neonates was comparatively more mature than that of the humerus. Once again, this indicates a difference in relative growth rates, and is a somewhat surprising result given the different functional demands on altricial and precocial chicks based on behavior and primary locomotor module. This may instead represent developmental channeling in avian growth, a feature with such deep evolutionary history that it is a nearly indelible part of their ontogeny. While maturity of these two limb bones later becomes coincident, as reported here and by others (e.g., Starck & Ricklefs, 1998), the initial difference appears to be common to all or most avians.

Furthermore, differences in cortical thicknesses across growth stages and within single elements suggest that width of the cortex may represent a non-adaptive feature of immature bird bones. Ultimately, there are in fact fewer histological differences along the altricial-precocial spectrum than anticipated, and it is difficult based on qualitative descriptions to identify a clear histological signal of developmental mode beyond relative maturity of both skeletal elements at the extreme ends of the spectrum. However, it is very possible that quantitative analyses of histological attributes would be more successful in parsing out finer categories along the altricial-precocial spectrum.

If relative growth rates can be inferred from tissue maturity as many previous studies suggest (e.g., Ricklefs, 1979; Kirkwood et al., 1989; Carrier & Leon, 1990; Montes et al., 2005) this research does support the conclusion of lower growth rates in precocial neonates, and higher growth rates in altricial neonates. The humerus and femur of chicks at the precocial end of the developmental spectrum were observed to be more mature than that of the humerus and femur respectively in chicks along the altricial end of the spectrum. This not only reflects the faster growth of altricial chicks, but also the higher functional demands on precocial chicks, which need to locomote and be relatively independent at an earlier age.

Bone growth appears to be a balance between growth rate, structural weakness/strength, and functional demands. This aspect of life history is a compromise between functional and developmental channeling, and it is the interplay of these two constraints that limits viable growth trajectories. However, within these constraints, different taxa alter their developmental trajectories by changing the timing of the onset and offset of bone growth. The bone of precocial chicks is more functionally mature than that of altricial neonates because of the biomechanical demands that come with precociality. The onset of bone growth is affected by developmental mode, with precocial chicks accomplishing more growth in the egg than altricial. Altricial chicks ultimately follow a similar trajectory of bone development, but in a shorter span of time (hatching to adult). On the other hand, timing of the offset of bone growth is independent of developmental mode, and instead changes with adult body size. Smaller birds achieve smaller body size by truncating their period of fast growth, parallel to the strategy used by avialans to evolve a small body size relative to their dinosaurian ancestors. As a result, small taxa achieve appositional “adult” size by the pre-fledgedling to subadult stage when the OCL appears. However, this does not correlate with complete skeletal maturity. Instead, the tissue continues to change
and mature through a combination of endosteal resorption, increased organization of bone texture (including a thickening of the OCL), and a reduction in vascularity.

Avian osteological development is clearly influenced by the canonical triad of constraints: functional, phylogenetic, and developmental (Seilacher, 1970; Gould, 2002). Functionally, the demands of locomotion (to varying degrees across the developmental spectrum) and strain on bone are high but are necessarily compromised by the need for fast growth which results in weaker bone. Phylogenetically, several histological characteristics observed are correlated with particular clades, including cortical thickness and unique features (e.g., intercortical gap left by the Kastschenko line in passeriforms). And developmentally, all birds are constrained by high growth rates and consequently weak bones, as well as ontogenetically channeled attributes, such as relatively more mature bone in the femur compared to the humerus at hatching.

This research lays the groundwork for future studies, such as longitudinal sections of the samples described here, sections from other skeletal elements, a detailed description of the presence of chondroid bone, quantification of additional microstructural features (e.g., vascular porosity and size and shape of osteocyte lacunae), inclusion of more specimens and taxa, and further investigation into the functional and phylogenetic differences between developmental modes and avian clades to gain greater insight into the influence of these factors on bone growth and evolution.

ACKNOWLEDGEMENTS

We are grateful to Doug Osbourne Jr. and Sr. of the OK Corral Ostrich Farm, Steve Duncan and Avian Resources, the SPCA for Monterey County, and the Lindsay Wildlife Museum for donating specimens to this research. The team running the prep lab at the Museum of Vertebrate Zoology was an incredible help in skeletonizing specimens, led by Theresa Barclay. Andrew Lee was a great source of support regarding methodology of preparing thin-sections of undecalciﬁed modern bone. Thanks also to Marvalee Wake, David Lindberg, Rauri Bowie, and Sabrina Agarwal who provided feedback on earlier versions of this manuscript. Lucy Chang and Jeremiah Scott provided guidance on the statistical analyses and interpretations. Finally, we are grateful to Lucas Legendre and Edina Prondvai for their constructive criticism of this paper. As reviewers, both went above and beyond to provide extensive helpful feedback. We are especially grateful to Dr. Legendre for sharing his expertise regarding the statistical analyses, and for providing additional support and clarification. We are likewise grateful to Dr. Prondvai for having such an eye for detail, and for catching many errors that would have been easy to overlook. We also express our gratitude to her for providing feedback and assistance identifying chondroid bone in earlier stages of this work.

ADDITIONAL INFORMATION AND DECLARATIONS

Funding

This work was supported by the Joseph Mallard Graduate Fellowship from the Museum of Vertebrate Zoology at the University of California, Berkeley; and the Doris O. and Samuel
P. Welles Research Fund from the University of California Museum of Paleontology. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Grant Disclosures
The following grant information was disclosed by the authors:
Museum of Vertebrate Zoology at the University of California, Berkeley.
University of California Museum of Paleontology.

Competing Interests
The authors declare that they have no competing interests.

Author Contributions
• Jessie Atterholt conceived and designed the experiments, performed the experiments, analyzed the data, prepared figures and/or tables, authored or reviewed drafts of the paper, and approved the final draft.
• Holly N. Woodward analyzed the data, authored or reviewed drafts of the paper, and approved the final draft.

Data Availability
The following information was supplied regarding data availability:
Raw data (measurements of histological structures) are available in the Supplemental Tables.

Supplemental Information
Supplemental information for this article can be found online at http://dx.doi.org/10.7717/peerj.12160#supplemental-information.

REFERENCES
Aramburú RM. 1997. Descripción y desarrollo del pichón de la cotorra Myiopsitta monachus monachus (Aves: Psittacidae) en una población silvestre de Argentina. Revista Chilena de Historia Natural 70:53–58.
Atterholt J. 2011. Phylogenetic mapping of traits of the avian altricial-precocial spectrum, and its implications for inferring early avialan life history. Journal of Vertebrate Paleontology 31:64–65.
Atterholt J, Poust AW, Erickson GM, O’Connor JK. 2021. Intraskeletal osteohistovariability reveals complex growth strategies in a late Cretaceous enantiornithine. Frontiers in Earth Science 9:118.
Bell AK, Chiappe LM, Erickson GM, Suzuki S, Watabe M, Barsbold R, Tsogbaatar K. 2010. Description and ecologic analysis of Hollanda luceria, a late Cretaceous bird from the Gobi Desert (Mongolia). Cretaceous Research 31(1):16–26 DOI 10.1016/j.cretres.2009.09.001.
Blomberg S, Garland T, Ives A. 2003. Testing for phylogenetic signal in comparative data: behavioral traits are more labile. Evolution 57:717–745.
Bonucci E, Gherardi G. 1975. Histochemical and electron microscope investigations on medullary bone. Cell and Tissue Research 163:81–97.
Bourdon E, Castanet J, De Ricqlès A, Scofield P, Tennyson A, Lamrous H, Cubo J. 2009. Bone growth marks reveal protracted growth in New Zealand kiwi (Aves, Apterygidae). *Biology Letters* 5:639–642.

Cambra-Moo O, Delgado Buscalioni A, Cubo J, Castanet J, Loth M-M, De Margerie E, de Ricqlès A. 2006. Histological observations of enantiornithine bone (Saurischia, Aves) from the lower Cretaceous of Las Hoyas (Spain). *Comptes Rendus Palevol* 5(5):685–691 DOI 10.1016/j.crpv.2005.12.018.

Carrier DR, Auriemma J. 1992. A developmental constraint on the fledging time of birds. *Biological Journal of the Linnean Society* 47:1–17.

Carrier DR, Leon LR. 1990. Skeletal growth and function in the California gull (*Larus californicus*). *Journal of Zoology* 222:375–389.

Carril J, Tambussi CP, Rasskin-Gutman D. 2021. The network ontogeny of the parrot: altriciality, dynamic skeletal assemblages, and the avian body plan. *Evolutionary Biology* 48(1):1–13.

Case TJ. 1978. Speculations on the growth rate and reproduction of some dinosaurs. *Paleobiology* 4(3):320–328.

Castanet J, Grandin A, Abourachid A, de Ricqlès A. 1996. Expression de la dynamique de croissance dans la structure de l’os périostique chez *Anas platyrhynchos*. *Comptes Rendus de l’Académie des Sciences-Series III-Sciences de la Vie* 319:301–308.

Castanet J, Rogers KC, Cubo J, Jacques-Boisard J. 2000. Periosteal bone growth rates in extant ratites (ostriche and emu). Implications for assessing growth in dinosaurs. *Comptes Rendus de l’Académie des Sciences-Series III-Sciences de la Vie* 323:543–550.

Castanet J, Smirina E. 1990. Introduction to the skeletochronological method in amphibians and reptiles. In: *Annales des Sciences Naturelles Zoologie et Biologie Animale*. Amsterdam, Netherlands: ScienceDirect, 191–196.

Cerda IA, Pol D, Chinsamy A. 2014. Osteohistological insight into the early stages of growth in *Mussaurus patagonicus* (Dinosauria, Sauropodomorpha). *Historical Biology* 26:110–121.

Chinsamy A, Angst D, Canoville A, Göhlich UB. 2020a. Bone histology yields insights into the biology of the extinct elephant birds (Aepyornithidae) from Madagascar. *Biological Journal of the Linnean Society* 130:268–295.

Chinsamy A, Chiappe LM, Dodson P. 1994. Growth rings in Mesozoic birds. *Nature* 368:196–197.

Chinsamy A, Chiappe LM, Dodson P. 1995. Mesozoic avian bone microstructure: physiological implications. *Paleobiology* 21:561–574.

Chinsamy A, Chiappe LM, Marugán-Lobón J, Chunging G, Fengjiao Z. 2013. Gender identification of the Mesozoic bird *Confuciusornis sanctus*. *Nature Communications* 4:1381.

Chinsamy A, Elzanowski A. 2001. Evolution of growth pattern in birds. *Nature* 412:402–403.

Chinsamy A, Marugán-Lobón J, Serrano FJ, Chiappe L. 2020b. Osteohistology and life history of the basal pygostylian, *Confuciusornis sanctus*. *The Anatomical Record* 303:949–962.

Cordeiro-Spinetti E, Taichman RS, Balduino A. 2015. The bone marrow endosteal niche: how far from the surface? *Journal of Cellular Biochemistry* 116:6–11.

Cubo J, Baudin J, Legendre L, Quilhac A, De Buffrénil V. 2014. Geometric and metabolic constraints on bone vascular supply in diapsids. *Biological Journal of the Linnean Society* 112(4):668–677 DOI 10.1111/bij.12331.

Cubo J, Legendre P, de Ricqlès A, Montes LL, de Margerie E, Castanet J, Desseives Y. 2008. Phylogenetic, functional, and structural components of variation in bone growth rate of amniotes. *Evolution & Development* 10(2):217–227 DOI 10.1111/j.1525-142X.2008.00229.x.
Cubo J, Ponton F, Laurin M, De Margerie Ed, Castanet J. 2005. Phylogenetic signal in bone microstructure of sauropsids. *Systematic Biology* 54(4):562–574 DOI 10.1080/10635150591003461.

Currey J. 2003. The many adaptations of bone. *Journal of Biomechanics* 36(10):1487–1495 DOI 10.1016/S0021-9290(03)00124-6.

D’Emic MD, Benson RBJ. 2013. Measurement, variation, and scaling of osteocyte lacunae: a case study in birds. *Bone* 57(1):300–310 DOI 10.1016/j.bone.2013.08.010.

Dacke CG, Arkle S, Cooke DJ, Wormstone IM, Jones S, Zaidi M, Bascal ZA. 1993. Medullary bone and avian calcium regulation. *Journal of Experimental Biology* 184:63–88.

de Margerie E. 2006. Biomechanical function of bone microstructure in birds. *Comptes Rendus Palevol* 5:619–628.

-de Margerie E, Cubo J, Castanet J. 2002. Bone typology and growth rate: testing and quantifying ‘Amprino’s rule’ in the mallard (Anas platyrhynchos). *Comptes Rendus Biologies* 325:221–230.

-de Margerie E, Robin JP, Verrier D, Cubo J, Groscolas R, Castanet J. 2004. Assessing a relationship between bone microstructure and growth rate: a fluorescent labelling study in the king penguin chick (*Aptenodytes patagonicus*). *Journal of Experimental Biology* 207(5):869–879 DOI 10.1242/jeb.00841.

-de Margerie E, Sanchez S, Cubo J, Castanet J. 2005. Torsional resistance as a principal component of the structural design of long bones: comparative multivariate evidence in birds. *The Anatomical Record* 282:49–66.

-de Ricqlès A. 1975. Recherches paléohistologiques sur les os longs de tétrapodes VII—sur la classification, la signification fonctionnelle et l’histoire des tissus osseux des tétrapodes (première parti). *Annales de Paléontologie (Vertébrés)* 61:51–129.

-de Ricqlès A, Bourdon E, Legendre LJ, Cubo J. 2016. Preliminary assessment of bone histology in the extinct elephant bird Aepyornis (Aves Palaeognathae) from Madagascar. *Comptes Rendus Palevol* 15:197–208.

-de Ricqlès A, Mateus O, Antunes MT, Taquet P. 2001. Histomorphogenesis of embryos of upper Jurassic theropods from Lourinhã (Portugal). *Comptes Rendus de l’Académie des Sciences-Series IIA-Earth and Planetary Science* 332:647–656.

-de Ricqlès A, Meunier J, Castanet FJ, Francillon-Vieillot H. 1991. Comparative microstructure of bone. In: Hall BK, ed. *Bone Volume 3: Bone Matrix and Bone Specific Products*. Boca Raton, Florida: CRC Press.

-de Ricqlès A, Padian K, Horner J, Lammm E, Myhrvold N. 2003. Osteohistology of *Confuciusornis sanctus* (Theropoda: Aves). *Journal of Vertebrate Paleontology* 23:373–386.

-Dial K. 2003. Evolution of avian locomotion: Correlates of flight style, locomotor modules, nesting biology, body size, development, and the origin of flapping flight. *Auk* 120:941–952.

-Dial TR, Carrier DR. 2010. Precocial hindlimbs and altricial forelimbs of developing Mallard ducks: a study of locomotor performance and morphometrics. *Integrative and Comparative Biology* 50:E43.

-Enlow D, Brown S. 1957. A comparative histological study of fossil and recent bone tissues, Part 2. *Texas Journal of Science* 9:186–214.

-Erickson GM, Rauhut OW, Zhou Z, Turner AH, Inouye BD, Hu D, Norell MA. 2009. Was dinosaurian physiology inherited by birds? Reconciling slow growth in Archaeopteryx. *PLOS ONE* 4:e7390.

-Eroschenko V. 2008. di Fiore’s atlas of histology with functional correlations: lippincott.
Francillon H. 1980. Mise en évidence expérimentale du caractère annuel des lignes d’arrêt de croissance (LAC) chez le triton crêté, *Triturus cristatus* (LAUR.). *Bulletin of the Zoological Society, France* 105:343–346.

Francillon-Vieillot H, de Buffrénil V, Castanet J, Géraudie J, Meunier F, Sire J, Zylberberg L, de Ricqlès A. 1990. Microstructure and mineralization of vertebrate skeletal tissues. In: Carter JG, ed. *Skeletal Biomineralization: Patterns, Processes, and Evolutionary Trends*. New York: Van Nostrand Reinhold, 471–530.

González R, Cerda IA, Filippi LS, Salgado L. 2020. Early growth dynamics of titanosaur sauropods inferred from bone histology. *Palaeogeography, Palaeoclimatology, Palaeoecology* 537:109404.

Gould SJ. 2002. *The structure of evolutionary theory*. Harvard: Harvard University Press.

Hall BK. 2005. *Bones and cartilage: developmental and evolutionary skeletal biology*. Cambridge: Academic Press.

Houde P. 1987. Histological evidence for the systematic position of Hesperornis (Odontornithes: Hesperornithiformes). *Auk* 104(1):125–129 DOI 10.2307/4087243.

Houde P, Olson SL. 1986. Small arboreal nonpasserine birds from the early Tertiary of western North America. In: *Acta XIX Congressus Internationalis Ornithologici*. 2031–2036.

Huttenlocker AK, Woodward HN, Hall BK. 2013. Biology of bone. In: Padian K, Lamm E-T, eds. *Bone Histology of Fossil Tetrapods*. Berkeley: University of California Press.

Kastschenko N. 1880. Ueber die genese und architectur der batrachierknochen. *Archiv für Mikroskopische Anatomie* 19(1):1–52 DOI 10.1007/BF02952689.

Kirkwood JK, Duignan PJ, Kember NF, Bennett PM, Price DJ. 1989. The growth rate of the tarsometatarsus bone in birds. *Journal of Zoology, London* 217(3):403–416 DOI 10.1111/j.1469-7998.1989.tb02498.x.

Kuehn AL, Lee AH, Main RP, Simons EL. 2019. The effects of growth rate and biomechanical loading on bone laminarity within the emu skeleton. *PeerJ* 7(3):e7616 DOI 10.7717/peerj.7616.

Lee AH, Simons EL. 2015. Wing bone laminarity is not an adaptation for torsional resistance in bats. *PeerJ* 3:e823 DOI 10.7717/peerj.823.

Legendre L, Le Roy N, Martinez-Maza C, Montes L, Laurin M, Cubo J. 2013. Phylogenetic signal in bone histology of amniotes revisited. *Zoologica Scripta* 42:44–53.

Legendre LJ, Bourdon E, Scofield RP, Tennyson AJ, Lamrous H, de Ricqlès A, Cubo J. 2014. Bone histology, phylogeny, and palaeognathous birds (Aves: Palaeognathae). *Biological Journal of the Linnean Society* 112:688–700.

Lehman TM, Woodward HN. 2008. Modeling growth rates for sauropod dinosaurs. *Paleobiology* 34:264–281.

McGuire RS, Ourfalian R, Ezell K, Lee AH. 2020. Development of limb bone laminarity in the homing pigeon (*Columba livia*). *PeerJ* 8:e9878.

Meister W. 1962. Histological structure of long bones in penguins. *The Anatomical Record* 143:377–387.

Miller SC, Bowman BM. 1981. Medullary bone osteogenesis following estrogen administration to mature male Japanese quail. *Developmental Biology* 87(1):52–63 DOI 10.1016/0012-1606(81)90060-9.

Montes L, Castanet J, Cubo J. 2007. Relationship between bone growth rate and bone vascular density in amniotes: a first test of Amprino’s rule in a phylogenetic context. *Journal of Morphology* 268:1108.
Montes L, Castanet J, Cubo J. 2010. Relationship between bone growth rate and bone tissue organization in amniotes: first test of Amprino’s rule in a phylogenetic context. *Animal Biology* 60(1):25–41 DOI 10.1163/157075610X12610595764093.

Montes L, de Margerie E, Castanet J, de Ricqlès A, Cubo J. 2005. Relationship between bone growth rate and the thickness of calcified cartilage in the long bones of the Galloanserae (Aves). *Journal of Anatomy* 206(5):445–452 DOI 10.1111/j.1469-7580.2005.00410.x.

Nice MM. 1962. *Development of behavior in precocial birds*. New York: Linnaean Society.

O’gorman JP, Talevi M, Fernández MS. 2016. Osteology of a perinatal aristonectine (Plesiosauria; Elasmosauridae). *Antarctic Science* 1(1):1–12.

O’Connor JK, Wang M, Zheng X-T, Wang X-L, Zhou Z-H. 2014. The histology of two female early Cretaceous birds. *Vertebrata PalAsiatica* 52:112–128.

O’Connor JK, Wang M, Zhou S, Zhou Z. 2015. Osteohistology of the lower Cretaceous Yixian formation ornithuromorph (Aves) Iteravis huchzermeyeri. *Palaeontology Electronica* 18:1–11.

O’Keefe F, Sander P, Winrich T, Werning S. 2019. Ontogeny of polycotylid long bone microanatomy and histology. *Integrative Organismal Biology* 1:oby007.

Padian K, de Ricqlès A. 2020. Inferring the physiological regimes of extinct vertebrates: methods, limits and framework. *Philosophical Transactions of the Royal Society B* 375:20190147.

Padian K, de Ricqlès A, Horner JR. 2001. Dinosaurian growth rates and bird origins. *Nature* 412:405–408.

Padian K, Horner J, de Ricqlès A. 2004. Growth in small dinosaurs and pterosaurs: the evolution of archosaurian growth strategies. *Journal of Vertebrate Paleontology* 24:555–571.

Prondvai E, Godefroit P, Adriaens D, Hu D-Y. 2018. Intraskeletal histovariability, allometric growth patterns, and their functional implications in bird-like dinosaurs. *Scientific Reports* 8(1):1–16 DOI 10.1038/s41598-017-18218-9.

Prondvai E, Witten PE, Abourachid A, Huysseune A, Adriaens D. 2020. Extensive chondroid bone in juvenile duck limbs hints at accelerated growth mechanism in avian skeletogenesis. *Journal of Anatomy* 236(3):463–473 DOI 10.1111/joa.13109.

Prum R, Brush A. 2002. The evolutionary origin and diversification of feathers. *The Quarterly Review of Biology* 77(3):261–295 DOI 10.1086/341993.

Prum RO, Berv JS, Dornburg A, Field DJ, Townsend JP, Lemmon EM, Lemmon AR. 2015. A comprehensive phylogeny of birds (Aves) using targeted next-generation DNA sequencing. *Nature* 526(7574):569–573 DOI 10.1038/nature15697.

Rath N, Balog J, Huff W, Huff G, Kulkarni G, Tierce J. 1999. Comparative differences in the composition and biomechanical properties of tibiae of seven-and-seventy-two-week-old male and female broiler breeder chickens. *Poultry Science* 78(8):1232–1239 DOI 10.1093/ps/78.8.1232.

Ricklefs RE. 1968. Patterns of growth in birds. *Ibis* 110(4):419–451 DOI 10.1111/j.1474-919X.1968.tb00058.x.

Ricklefs RE. 1973. Patterns of growth in birds. II. Growth rate and mode of development. *The Ibis* 115(2):177–200 DOI 10.1111/j.1474-919X.1973.tb02636.x.
Ricklefs RE. 1979. Adaptation, constraint, and compromise in avian postnatal development. *Biological Review* 54(3):269–290 DOI 10.1111/j.1469-185X.1979.tb01013.x.

Rozenblut B, Ogielska M. 2005. Development and growth of long bones in European water frogs (Amphibia: Anura: Ranidae), with remarks on age determination. *Journal of Morphology* 265:304–317 DOI 10.1002/(ISSN)1097-4687.

Seilacher A. 1970. Arbeitskonzept zur konstruktions-morphologie. *Lethaia* 3(4):393–396 DOI 10.1111/j.1502-3931.1970.tb00830.x.

Simons EL, O’connor PM. 2012. Bone laminarity in the avian forelimb skeleton and its relationship to flight mode: testing functional interpretations. *Anatomical Record* 295(3):386–396 DOI 10.1002/ar.22402.

Skedros JG, Hunt KJ. 2004. Does the degree of laminarity correlate with site-specific differences in collagen fibre orientation in primary bone? An evaluation in the turkey ulna diaphysis. *Journal of Anatomy* 205(2):121–134 DOI 10.1002/(ISSN)1097-4687.

Starck JM. 1989. Zeitmuster der ontogenesen bei nestflüchtenden und neststockenden vögeln. *Cour Forsch-Inst Senckenberg* 114:1–319.

Starck JM. 1993. Evolution of avian ontogenies. *Current Ornithology* 10:275–366.

Starck JM. 1994. Quantitative design of the skeleton in bird hatchlings: does tissue compartmentalization limit posthatching growth rates? *Journal of Morphology* 222:113–131 DOI 10.1002/(ISSN)1097-4687.

Starck JM, Ricklefs RE. 1998. Patterns of development: the altricial-precocial spectrum. In: Starck JM, Ricklefs RE, eds. *Avian Growth and Development: Evolution within the Altricial-Precocial Spectrum*. Oxford: Oxford University Press, 3–30.

Symonds MR, Blomberg SP. 2014. A primer on phylogenetic generalised least squares. In: Garamszegi LZ, ed. *Modern Phylogenetic Comparative Methods and their Application in Evolutionary Biology*. Berlin: Springer, 105–130.

Turvey S, Holdaway R. 2005. Postnatal ontogeny, population structure, and extinction of the giant moa Dinornis. *Journal of Morphology* 265:70–86 DOI 10.1002/(ISSN)1097-4687.

Van de Velde J, Vermeiden J, Bloot A. 1985. Medullary bone matrix formation, mineralization, and remodeling related to the daily egg-laying cycle of Japanese quail: a histological and radiological study. *Bone* 6(5):321–327 DOI 10.1016/0350-114X(85)90322-9.

Voeten DF, Cubo J, De Margerie E, Röper M, Beyrand V, Bureš S, Tafforeau P, Sanchez S. 2018. Wing bone geometry reveals active flight in Archaeopteryx. *Nature Communications* 9:923 DOI 10.1038/s41467-018-03296-8.

Wang J, Hao X, Kundrát M, Liu Z, Uesugi K, Jurašeková Z, Guo B, Hoshino M, Li Y, Monfroy Q. 2019. Bone tissue histology of the Early Cretaceous bird Yanornis: evidence for a diphyletic origin of modern avian growth strategies within Ornithuromorpha. *Historical Biology* 32(1):1–13.

Wang M, O’Connor JK, Bailleul AM, Li Z. 2020. Evolution and distribution of medullary bone: evidence from a new early Cretaceous enantiornithine bird. *National Science Review* 7(6):1068–1078 DOI 10.1093/nsr/nzw214.

Watanabe J. 2018. Ontogeny of surface texture of limb bones in modern aquatic birds and applicability of textural ageing. *The Anatomical Record* 301:1026–1045.
Wilson LE, Chin K. 2014. Comparative osteohistology of *Hesperornis* with reference to pygoscelid penguins: the effects of climate and behaviour on avian bone microstructure. *Royal Society Open Science* 1:140245.

Woodward HN, Tremaine K, Williams SA, Zanno LE, Horner JR, Myhrvold N. 2020. Growing up Tyrannosaurus rex: Osteohistology refutes the pygmy “Nanotyrannus” and supports ontogenetic niche partitioning in juvenile Tyrannosaurus. *Science Advances* 6(1):p.eaax6250.