Integrating gross morphology and bone histology to assess skeletal maturity in early dinosauromorphs: new insights from Dromomeron (Archosauria: Dinosauromorpha)

Christopher T. Griffin¹, Lauren S. Bano², Alan H. Turner³, Nathan D. Smith⁴, Randall B. Irmis⁵,⁶ and Sterling J. Nesbitt¹

¹ Department of Geosciences, Virginia Tech, Blacksburg, VA, USA
² Department of Biology, Virginia Tech, Blacksburg, VA, USA
³ Department of Anatomical Sciences, Stony Brook University, Stony Brook, NY, USA
⁴ The Dinosaur Institute, Natural History Museum of Los Angeles County, Los Angeles, CA, USA
⁵ Natural History Museum of Utah, University of Utah, Salt Lake City, UT, USA
⁶ Department of Geology and Geophysics, University of Utah, Salt Lake City, UT, USA

ABSTRACT

Understanding growth patterns is central to properly interpreting paleobiological signals in tetrapods, but assessing skeletal maturity in some extinct clades may be difficult when growth patterns are poorly constrained by a lack of ontogenetic series. To overcome this difficulty in assessing the maturity of extinct archosaurian reptiles—crocodilians, birds and their extinct relatives—many studies employ bone histology to observe indicators of the developmental stage reached by a given individual. However, the relationship between gross morphological and histological indicators of maturity has not been examined in most archosaurian groups. In this study, we examined the gross morphology of a hypothesized growth series of Dromomeron romeri femora (96.6–144.4 mm long), the first series of a non-dinosauriform dinosauromorph available for such a study. We also histologically sampled several individuals in this growth series. Previous studies reported that D. romeri lacks well-developed rugose muscle scars that appear during ontogeny in closely related dinosauromorph taxa, so integrating gross morphology and histological signal is needed to determine reliable maturity indicators for early bird-line archosaurs. We found that, although there are small, linear scars indicating muscle attachment sites across the femur, the only rugose muscle scar that appears during ontogeny is the attachment of the M. caudofemoralis longus, and only in the largest-sampled individual. This individual is also the only femur with histological indicators that asymptotic size had been reached, although smaller individuals possess some signal of decreasing growth rates (e.g., decreasing vascular density). The overall femoral bone histology of D. romeri is similar to that of other early bird-line archosaurs (e.g., woven-bone tissue, moderately to well-vascularized, longitudinal vascular canals). All these data indicate that the lack of well-developed femoral scars is autapomorphic for this species, not simply an indication of skeletal immaturity. We found no evidence of the high intraspecific variation present in early dinosaurs and other dinosauromorphs, but a limited sample

How to cite this article Griffin CT, Bano LS, Turner AH, Smith ND, Irmis RB, Nesbitt SJ. 2019. Integrating gross morphology and bone histology to assess skeletal maturity in early dinosauromorphs: new insights from Dromomeron (Archosauria: Dinosauromorpha). PeerJ 7:e6331 DOI 10.7717/peerj.6331

OPEN ACCESS
size of other early bird-line archosaur growth series make this tentative. The evolutionary history and phylogenetic signal of gross morphological features must be considered when assessing maturity in extinct archosaurs and their close relatives, and in some groups corroboration with bone histology or with better-known morphological characters is necessary.

**Subjects**  Evolutionary Studies, Paleontology

**Keywords**  Ontogeny, Skeletal maturity, Histology, Dinosauromorph, Triassic, Bone scar

**INTRODUCTION**

Assessing the skeletal maturity of individuals of extinct taxa is critical to interpreting the paleobiology of their constituent species and clades. Interpretations of the growth stages of a taxon impact evaluations of species synonymy (Abdala & Giannini, 2000; Vucetich et al., 2005; Cartelle & De Iuliis, 2006; Billet, de Muizon & Mamani Quispe, 2008; Scannella & Horner, 2010; Shelton et al., 2012), potential sexual dimorphism (Raath, 1990; Klein & Sander, 2008; Saitta, 2015), maximum body size (e.g., Huttenlocker & Rega, 2012; Turner & Nesbitt, 2013), intraspecific variation (Brinkman, 1988; Raath, 1990; Griffin & Nesbitt, 2016a, 2016b), and ecological role (Codron et al., 2012), among others (Hone, Farke & Wedel, 2016). Reliable skeletal maturity indicators are present in many extant vertebrate groups, and these characteristics have been used successfully to estimate skeletal maturity and growth stage in their extinct members. For example, in placental mammals, the closure of epiphyseal sutures indicates cessation of growth (e.g., Dawson, 1925; Morris, 1972; Ogden, Conlogue & Rhodin, 1981; Storå, 2002; Nilsson & Baron, 2004; Köhler & Moyà-Solà, 2009; Moran et al., 2015) and the presence of permanent teeth can give an indication that the individual is not among the earliest stages of growth, when deciduous (or milk) teeth are present (e.g., Morris, 1972; Spinage, 1973; Smith, Crummett & Brandt, 1994). Among reptiles, many squamates also fuse epiphyses during ontogeny, which provides an indicator of body size at skeletal maturity in taxa with determinate growth (e.g., Haines, 1969; Chinsamy-Turan, 2005; Frýdlová et al., 2017). However, it remains challenging to assess the skeletal maturity of individuals from clades that are highly disparate or possess long evolutionary histories with highly modified living members. This difficulty is exemplified in archosaurs (crocodylians, birds, and their extinct relatives), which possess a larger range of body sizes than any other clade of terrestrial animals (Carrano, 2006; Benson et al., 2014; Turner & Nesbitt, 2013). In this clade, many indicators of skeletal maturity in other living reptiles are absent or of limited use (e.g., epiphyseal fusion, Chinsamy-Turan, 2005; Frýdlová et al., 2017), making determining skeletal maturity difficult (Hone, Farke & Wedel, 2016).

To overcome this dilemma, many studies have employed a variety of assessment techniques derived from a combination of observations in living taxa and deductions made from extinct taxa. The use of bone histology in extinct archosaurs (e.g., Horner, de Ricqlès & Padian, 2000; Padian, Horner & de Ricqlès, 2004; Erickson, 2014) aids in determining an individual’s skeletal maturity through the presence or absence of lines of
arrested growth (LAGs) as well as the pattern of tissue organization (e.g., collagen fiber/hydroxyapatite crystal density, fiber orientation) and vascularity present in the bone (reviewed in Huttenlocker, Woodward & Hall, 2013). In addition to its success in providing indicators that asymptotic size has been reached (e.g., Woodward, Horner & Farlow, 2011; Erickson et al., 2004; 2007; Erickson, 2014), studies of bone histology have contributed to answering a diverse array of questions, including determining the thermometabolism of extinct archosaurs (e.g., Legendre et al., 2016), hypothesizing sex-specific tissues (e.g., Schweitzer, Wittmeyer & Horner, 2005; 2016; Lee & Werning, 2008) and estimating age of sexual maturity in dinosaurs (Lee & Werning, 2008), estimation of growth rates (e.g., Cubo et al., 2012), and hypothesizing behavioral implications of embryonic and prenatal development (Horner, Padian & de Ricqlès, 2001). However, creating osteohistological thin sections is time consuming and permanently alters original bone morphology through consumptive sampling, and so this method may not be viable for rare species and for specimens with exceptional preservation. Furthermore, some specimens comprise a taphonomic mode that may not preserve internal structure (e.g., Jans, 2008; Pewkliang, Pring & Brugger, 2008). Depending on the taxon, osteohistological thin sections may also lack cyclical indicators of numerical ontogenetic age (e.g., LAG), making these data ambiguous or of limited utility when used independently of other methods (e.g., the silesaurid Asilisaurus kongwe, Griffin & Nesbitt, 2016a; several sauropod dinosaurs, Sander et al., 2004). Therefore, although osteohistology is a powerful tool for understanding growth both in terms of absolute and relative ontogenetic age, its potential drawbacks necessitate other means of constraining maturity assessment.

In contrast with osteohistological age assessment, gross morphological indicators of skeletal maturity provide an immediate and nondestructive assessment of the ontogenetic stage of an individual. Skeletal proportions (Bennett, 2001; Piechowski, Taldana & Dzik, 2014), periosteal surface texture (Sampson, Ryan & Tanke, 1997; Tumarkin-Deratzian, Vann & Dodson, 2006, 2007; Brown, Russell & Ryan, 2009), suture closure (Brochu, 1996; Irmis, 2007; Bailleul et al., 2016), muscle scarring (Raath, 1990; Tumarkin-Deratzian, Vann & Dodson, 2006, 2007; Griffin & Nesbitt, 2016a, 2016b), and clade specific indicators (e.g., growth of ceratopsian ornamentation, Hone, Farke & Wedel, 2016) have all been used to assess skeletal maturity in extinct archosaurs, especially non-avian dinosaurs. However, these methods are not always applicable across clades, which can produce results difficult to assess independently. For example, variable timing of the closure of sutures, a commonly used method for assessing skeletal maturity in archosaurs (e.g., vertebral neurocentral sutures, Brochu, 1996; Irmis, 2007) can be influenced by changes in biomechanics (e.g., in cranial sutures, Bailleul et al., 2016), and the sequence of postcranial co-ossification events can be variable both between clades (Irmis, 2007) and within a single species (Griffin, 2018). Therefore, skeletal maturity assessment using morphological characteristics alone may result in ambiguous age determinations. In addition, taken alone, these methods are nearly always relative age indicators; without independent evidence from other sources (e.g., histology), they can only indicate whether one individual is older or younger than another.
The strength of a well-supported assessment of skeletal maturity lies in the combination of external morphological and histological data. Yet the relationship between histological and morphological maturity has not been critically examined in most archosaurian clades (but see Woodruff, Fowler & Horner, 2017 for this combination of methods in diplodocids). This integration is especially important in examining growth of early dinosaurs and their close relatives, which possess high intraspecific variation in growth (Griffin & Nesbitt, 2016a, 2016b). This high variation has been almost entirely observed through the lens of gross morphology (Griffin & Nesbitt, 2016a, 2016b; Barta, Nesbitt & Norell, 2017; Griffin, 2018), whereas histology has only been touched on in a small number of taxa (Griffin & Nesbitt, 2016a). Therefore, the origin of this variation and the relationship between gross morphological and histological ontogenetic changes is poorly constrained by a lack of integration between methods of maturity assessment in early bird-line archosaurs.

In this study, we examined the histology and gross morphology of the first available femoral growth series of a non-dinosauriform dinosauromorph to better understand the origin of the dinosaurian growth pattern. We assess the skeletal maturity of well-preserved femora of Dromomeron romeri, an early-diverging dinosauromorph from the Late Triassic Period of northern New Mexico, USA (Irmis et al., 2007; Nesbitt et al., 2009a), using well-developed bone scars, which typically indicate muscle, ligament, and tendon attachment sites. These femoral features are important morphological characteristics that have been used for determining skeletal maturity in previous studies of early dinosauromorphs (Raath, 1990; Nesbitt et al., 2009a; Piechowski, Talanda & Dzik, 2014; Griffin & Nesbitt, 2016a, 2016b). Although these scars appear sequentially throughout ontogeny (Nesbitt et al., 2009a; Griffin & Nesbitt, 2016a, 2016b), how the development of bone scars relates to absolute ontogenetic age—and how widespread the utility of these scars are for assessing skeletal maturity among early bird-line archosaurs—is poorly constrained by a lack of ontogenetic series. Previous studies indicate that D. romeri lacks well-developed bone scars that other closely related archosaurs develop in the proximal region of the femur (Irmis et al., 2007; Nesbitt et al., 2009a; Martínez et al., 2016; Cabreira et al., 2016). In this study, we determine whether this lack of femoral scarring is indicative of the skeletal immaturity of the specimens (Bennett, 2015), or if this absence is an evolutionary novelty (i.e., autapomorphy). In doing so we integrated histological and gross morphological signals to better understand their relationship between bone scarring, ontogenetic age, and evolutionary history in early archosaurs.

**MATERIALS AND METHODS**

**Taxonomic justification**

We consider all femoral specimens examined in this study to be assignable to D. romeri based on the shared presence of diagnostic character states of the referred material and the holotype femur (e.g., absence of a distinct ridge for the attachment of the M. caudofemoralis longus (this absence is not ontogenetic, see below); presence of a ridge on the anteromedial edge of the distal end; Irmis et al., 2007; Nesbitt et al., 2009a) and these specimens’ provenance (all were collected from the type locality—Hayden Quarry, ...
Ghost Ranch, NM, USA). The tibia and fibula of GR 238 are identifiable as *D. romeri* because they were found articulated with each other and with the femur of an individual diagnosable as *D. romeri*, even though the proximal and distal ends of these elements are not preserved.

*Bennett (2015)* recently suggested that the holotype of *D. romeri*, an isolated femur (GR 218), is a skeletally immature individual of the theropod *Tawa hallae* based on femoral size, the absence of a fourth trochanter interpreted as indicative of skeletal immaturity, and the suggestion that the morphology of the proximal and distal ends of the femur are incompletely ossified. However, the smallest *Tawa hallae* femur (116.3 mm in length; GR 244) is smaller than the larger *D. romeri* femora in our study (e.g., 144.4 mm in length, GR 1038; 135.3 mm in length, GR 238), and all recent cladistic analyses including early dinosauromorphs have recovered *D. romeri* as a lagerpetid dinosauromorph (*Irmis et al., 2007; Nesbitt et al., 2009a; 2009b; Nesbitt, 2011; Martínez et al., 2016; Cabreira et al., 2016; Müller, Langer & Dias-da-Silva, 2018*). Additionally, the postcranium of an associated specimen of *D. romeri* (GR 1041) lacks dinosaurian and dinosauromorph synapomorphies, instead possessing apomorphic character states that pertain to lagerpetids—outside Dinosauriformes—and these character optimizations have been borne out in all phylogenetic analyses. Specifically, *D. romeri* lacks an ilium with a convex ventral acetabular margin (= “imperforate” acetabulum, character 273–0, *Nesbitt, 2011*; GR 1041), a fibular condyle of the tibia that is level with the medial condyle at its caudal border (character 201–1 from *Cabreira et al., 2016*; character from *Langer & Benton, 2006*), and a posterior groove present on the astragalus (character 217–1 from *Cabreira et al., 2016*; character from *Nesbitt, 2011*). In addition, *D. romeri* has been independently supported as a unique taxon of non-dinosaurian dinosauromorph based on the large number of differences in femoral characters between *D. romeri* and *Tawa hallae* compared to ontogenetic series of other early dinosauromorphs (*Müller, 2017*). Thus, all current evidence strongly supports the original division of the two taxa and we concur with *Müller (2017)* in considering *D. romeri* to be a non-dinosaurian dinosauromorph. Nonetheless, our examination here of the gross morphological and histological ontogeny of *D. romeri* provides further evidence to test *Bennett’s (2015)* hypothesis. We can falsify this hypothesis if any *Dromomeron* specimens are demonstrated to be near skeletal maturity.

**Gross morphological assessment and histological sampling**

Seven femora were included in our *Dromomeron romeri* sample, hypothesized to form an ontogenetic series (GR 218, 234, 238, 1036–1039; femur length between 96.6 and 144.4 mm; *Table 1*). We use the term ‘well-developed bone scar’ to refer to rugose scars of muscle, tendon, or ligament attachment that are distinct from the shaft or main body of the bone. The term ‘poorly developed bone scar’ refers to scars of attachment which do not comprise rugose processes or features distinct form the shaft, such as thin lineations or grooves left by muscle attachments on bone. Poorly developed bone scars can be useful for hypothesizing muscle attachment sites but are not as informative for assessing maturity (e.g., *Griffin & Nesbitt, 2016a*). We evaluated the femora for presence of muscle
scars by comparing the position of femoral muscle origins and insertions in extant Aves and Crocodylia, those inferred in dinosaurs and relatives (Rowe, 1986; Hutchinson, 2001), and in closely related taxa [i.e., studies on Dromomeron gregorii (Nesbitt et al., 2009a); D. gigas (Martínez et al., 2016); Ixalerpeton polesinensis (Cabreira et al., 2016); Asilisaurus kongwe (Griffin & Nesbitt, 2016a); Teleocrater rhadinus (Nesbitt et al., 2017)]. We whitened specimens for high-contrast photography of bone scars using removable ammonium chloride following standard techniques (Feldmann, 1989; Green, 2001; Hegna, 2010; Shelburne & Thompson, 2016). After coating the specimen in a thin layer of B-72 glue dissolved in acetone, we heated the ammonium chloride in a glass test tube and held the specimen under the resulting vapor stream until the area of interest was satisfactorily whitened. Each specimen was photographed with a Canon EOS Rebel t4i digital camera. Following photography, the ammonium chloride was removed with a gentle stream of water. We used a macro lens (Tamron AF 90mm f/2.8 Di SP AF/MF 1:1 Macro Lens) for close-up photography of these bone scars, and photographs taken in stacks of different focal planes were blended in Adobe Photoshop CS6 v. 13.0.6.

One isolated femur (GR 1036), a tibia and fibula from the largest articulated individual (GR 238), and the largest femur (GR 1038) were histologically sampled as close to the midshaft region as possible. The midshaft region of the bone is generally consistent in the type of tissue present and does not remodel as much as other parts of the bone; the periosteal surface is also older, and therefore has more recorded information when compared to the distal and proximal ends (Padian, Lamm & Werning, 2013). To preserve the original gross morphology of the specimens before destructive sampling, we molded (using Smooth-On Mold Star 15 SLOW platinum silicone rubber) and casted (using Smooth-On Smooth Cast 30 liquid plastic resin) each specimen before thin sectioning. The tibia and fibula of GR 238 were sampled because the femora of this individual were not available for histological sectioning. Because the largest femur (GR 1038) was from the largest known D. romeri individual, and was well-preserved and complete, we were unable to section this element fully. To collect data on the histological maturity of this individual, we instead removed a portion of the midshaft which included the outermost cortex, rather than cut the entire element in half. Although we were not sure if this sample contained a complete record of the cortex when we sampled

| Specimen | Femoral length | Maximum proximal width | Maximum distal width |
|----------|----------------|------------------------|----------------------|
| GR 1037  | 96.6           | 20.1                   | 18.4                 |
| GR 218   | 96.9           | 18.7                   | 18.3                 |
| GR 1036* | 103.9          | 18.9                   | 18.5                 |
| GR 234   | 126.8          | 23.1                   | 23.3                 |
| GR 238*  | 135.3          | 23.3                   | 26.7                 |
| GR 1039  | 135.5          | 30                     | 26.7                 |
| GR 1038* | 144.4          | 28.5                   | 26.3                 |

Note: Asterisks (*) indicate specimens that were histologically sampled.
this element, we have evidence for the presence of endosteal lamellae in the internalmost portion of the sample (see Results), indicating that at least a portion of this sampled bone fragment spans the entire cortex. We created two histological slides from this portion, with slide 1 being the more proximal section of the two.

We embedded the sampled bone in Castolite AC clear polyester resin (Eager Polymers) and placed it in a vacuum chamber for 2–3 min to eliminate bubbles before curing overnight. Thin wafers (1.5 mm) were cut from the original sample perpendicular to the proximodistal axis with an IsoMet 1000 precision saw. We ground down one side, starting with a 1200 grit grinding disc and ending with a 2400 grit disk; the wafer was then polished with 0.3 micron slurry powder. We then glued the polished side face down on a slide and the opposite side was ground and polished to the desired thickness. Finally, we examined the slides with an Olympus BX51 petrographic microscope with plane- and cross-polarized light, using a gypsum wedge (530 nm) to better observe the orientations of hydroxyapatite crystallites (which are in turn influenced by the orientation of the original collagen fibrils). Images were captured with an Infinity1 camera and associated software. We also captured high-resolution images of the full slides using a Nikon Eclipse LV100ND transmitted and reflected light microscope, imaging them using a Nikon DS-Fi2 camera and Digital sight DS-U3 interface together with a Prior ProScan III automated microscope stage and digitally assembled using Nikon NIS-Element Basic Research v. 4.40.00 (Build 1084). These full-slide images were captured in three light regimes: plane-polarized light, cross-polarized light, and cross-polarized light with a gypsum plate (530 nm). The exception to this was the histology of the largest femur, GR 1038, which was examined with an Olympus BX51 research microscope under plane- and cross-polarized light (with and without a gypsum plate), and images were captured with Lumenera Infinity capture imaging software. Large high-resolution whole-slide images have been uploaded onto the online repository Morphobank (Project Number 3127: http://morphobank.org/permalink/?P3127).

RESULTS

Gross morphology

Well-developed scars just distal to the proximal articular surface of the femur do not appear in our sample of *Dromomeron romeri* individuals (Figs. 1 and 2). Instead, thin lineations of striated bone (i.e., poorly developed scars) mark what we hypothesize to be the insertion of the *M. iliotrochantericus caudalis* on the anterolateral surface of the proximal end of the femur. This muscle has been interpreted to form the anterior trochanter in many other early bird-line archosaurs (Hutchinson, 2001; Maidment & Barrett, 2011), which appears during ontogeny in several early dinosauromorphs (D. gregorii, Nesbitt et al., 2009a; Asilisaurus kongwe, Griffin & Nesbitt, 2016a). This poorly developed scar is in a similar location as the interpreted insertion of the *M. iliofemoralis externus* in the non-ornithodiran avemetatarsalian *Teleocrater rhadinus* (Nesbitt et al., 2017; note that the insertions of the *M. iliofemoralis externus* [= trochanteric shelf] and the *M. iliotrochantericus caudalis* [= anterior trochanter] are mistakenly reversed in
Nesbitt et al., 2017, but corrected in Nesbitt et al., 2018). Larger femora of D. gregorii also possess a continuous lightly striated structure which is hypothesized to correspond to both the anterior trochanter and trochanteric shelf (Nesbitt et al., 2009a).

Unlike the other scars on the proximal end of the femur, the attachment site morphology of the M. caudofemoralis longus insertion (= fourth trochanter in most archosaurs,
Hutchinson, 2001; Maidment & Barrett, 2011; Schachner, Manning & Dodson, 2011) does vary in our sample of *D. romeri* (Figs. 1 and 2C). All individuals in this series lack the ridge-like fourth trochanter found in many other lagerpetid taxa (*D. gregorii*, Nesbitt et al., 2009a; *Lagerpeton chanarensis*, Sereno & Arcucci, 1994; *Ixalerpeton polesinensis*, Cabreira et al., 2016). However, the largest femur in our sample (GR 1038) possesses a rugose scar marking the insertion of the *M. caudofemoralis longus* (Fig. 2D), with similar morphology to the homologous scar in *D. gigas* (Martínez et al., 2016). However, unlike *D. gigas*, *D. romeri* shows no evidence of a posteromedial scar (sensu Martínez et al., 2016), which may be homologous with the ‘obturator ridge’ of early dinosauriforms and dinosaurs (Griffin, 2018). Nonetheless, *D. romeri* does possess poorly developed scars on this portion of the femur, which probably mark the attachment of the muscle(s) or ligament(s) that form the ‘obturator ridge’ of *D. gigas* and dinosauriforms. Therefore, only a single well-developed muscle scar, the insertion of the *M. caudofemoralis longus*, appears during ontogeny in *D. romeri*; it does not form a flange or ridge-like fourth

**Figure 2** Bone scars of the largest femur of *Dromomeron romeri* in (A) anterolateral and (B) posteromedial views. (C) The attachment of the *M. iliofemoralis* in standard view, whitened, and in line drawing. (D) The attachment of the *M. femorotibialis externus* in standard view, whitened, and in line drawing. (E) The ‘obturator ridge’ in standard view, whitened, and in line drawing. (F) The well-developed attachment scar of the *M. caudofemoralis longus* in standard view, whitened, and in line drawing. (G) The attachment of the *M. gastrocnemius externus* in standard view, whitened, and in line drawing. Thin lineations in all line drawings indicate muscle scars. GR 1308 is a right femur. Scale bars are 1 cm. All photographs and line drawings by Christopher Griffin.

Griffin et al. (2019), *PeerJ*, DOI 10.7717/peerj.6331/fig-2
trochanter as present in almost all other early diverging dinosauromorphs and early archosaurs (cf. Nesbitt, 2011: Figs. 37–39).

The distal end of the femur also possesses lineations that indicate muscle attachment sites (Figs. 1 and 2). The anterior of the distalmost portion of the femur, just proximal to the unfinished bone of the distal articular surface, possesses a rough, linearly grooved area that marks the attachment of *M. femorotibialis externus* (following the hypothesis of Nesbitt et al., 2009a). This scar is present in all femora of *D. romeri* for which its presence can be evaluated, including the smallest specimen (GR 218; Fig. 1), and is also present in larger individuals of *D. gregorii* (Nesbitt et al., 2009a) as well as in *D. gigas* (Martínez et al., 2016); this scar has been hypothesized to be a synapomorphy of the clade *Dromomeron* (Nesbitt et al., 2009a; Müller, Langer & Dias-da-Silva, 2018: character state 194-1).

Another poorly developed scar extends in an anteroproximal to posterodistal orientation across the posterior side of the femur, just proximal to the popliteal surface (Fig. 1D). This scar is subtle and unable to be evaluated in all specimens of *D. romeri* because of preservation quality; however, two of the smaller individuals lack this scar (GR 218, 1037; Fig. 1B), and it is present in the two largest individuals for which it can be evaluated (GR 234, 1038; Figs. 1D and 2E). This suggests that this scar may appear during ontogeny, although the condition in all *D. romeri* specimens greatly differs from the well-developed scars typical of archosaurian ontogeny (e.g., anterior trochanter, fourth trochanter, trochanteric shelf; Raath, 1990; Tumarkin-Deratzian, Vann & Dodson, 2006, 2007; Griffin & Nesbitt, 2016a, 2016b). This small posterior femoral scar appears to also be present in the femora of other lagerpetids, including *D. gregorii* (Nesbitt et al., 2009a: Fig. 2B) and *D. gigas* (Martínez et al., 2016: Fig. 2.2), and potentially *Lagerpeton chanarensis* (Nesbitt et al., 2009a: Fig. 3B). Based on its location in comparison to *Alligator* and *Aves* (Carrano & Hutchinson, 2002), we hypothesize that this scar is an attachment for *M. gastrocnemius externus* (= *Mm. gastrocnemii pars lateralis* of *Aves*; Carrano & Hutchinson, 2002).

**Histology**

For osteohistological descriptions, we use the terminology of Werning (2013), who expanded on and provided diagnostic criteria for the terminology of Francillon-Vieillot et al. (1990). Because of uncertainty in identification given the intermediate level of tissue organization present in many of these sections (see below) and the fact that we have no longitudinally sampled slides to diagnose woven-fibered bone with the highest certainty (Stein & Prondvai, 2014; Prondvai et al., 2014), we do not refer to woven- or parallel-fibered tissue with lamellated primary osteons as a ‘fibrolamellar complex’, but instead describe the tissue organization and vascular orientation/style separately.

The smaller of the two histologically sampled femora (GR 1036) is partially crushed, but sections of cortical bone preserving the entire endosteal-periosteal cross section are present. This bone comprises woven-fibered bone with primary osteons in the interiormost one-half to two-thirds of the cortex (in what is often termed a ‘fibrolamellar complex’, but see Stein & Prondvai, 2014). The apatite crystals in this region are more
Figure 3  Bone histology of three individuals of *Dromomerom romeri*. Silhouettes are modified from Cabreira et al. (2016: Fig. 1A). (A) Femoral histology of the smaller sampled individual (GR 1036) under cross-polarized light with a gypsum waveplate, showing two LAGs near the periosteal surface. (B) Femoral histology of GR 1036 under plane-polarized and cross-polarized light showing the transition from less- to more-organized tissue in the external third of the cortex. (C) Femoral histology of GR 1036 showing tissues typical of the less-organized primary bone of the internal two-thirds of the cortex, under plane-polarized and cross-polarized light. (D) Fibular histology of the mid-sized sampled individual (GR 238) under cross-polarized light with a gypsum waveplate, showing the boundary of the medullary bone with endosteal lamellae, secondary osteons, and highly organized primary bone. (E) Fibular histology of GR 238 under cross-polarized light with a gypsum waveplate showing two LAGs and the periosteal surface. (F) Fibular histology of GR 238 under cross-polarized light with a gypsum waveplate showing two LAGs and the periosteal surface. (G) Tibial histology of the mid-sized sampled individual.
organized than stereotypical woven-fibered bone (Figs. 3A–3C), and may be considered intermediate between ‘classic’ woven-fibered bone and highly organized parallel-fibered bone (sensu Stein & Prondvai, 2014). Consistent with this level of organization, the osteocyte lacunae in this region are preferentially oriented circumferentially in the interior portion of the cortex, although many lacunae possess different orientations throughout this region. The internal, woven-fibered portion of the cortex is well-vascularized with primary osteons possessing one to two thin bands of concentric lamellae. These vascular canals are small (~10–30 μm in diameter) and circular, although some are slightly circumferentially oblong, and almost all are longitudinally oriented in the interiormost portion of the bone. Some radially oriented canals are present in the more external portions of the woven-fibered bone, and these canals are oriented at a slight angle (~25–30°) to the classic radial orientation, in which canals extend directly towards the subperiosteal surface. All these radial canals are angled in the same direction and to the same degree. Endosteal lamellae are present where the endosteal border is preserved. Roughly one-half to two-thirds of the distance to the subperiosteal surface, the woven-fibered bone smoothly transitions to highly organized parallel-fibered bone, and after this transition vascular density noticeably decreases to moderately well vascularized, or even poorly vascularized just internal to the subperiosteal surface (Figs. 3A and 3B). These vascular canals are primary osteons, and most are longitudinally oriented in this region with only one layer of lamellar bone ringing the canal. All osteocyte lacunae are elongate and oriented circumferentially in bands following the orientation of the collagen fibers. A LAG is present ~100 μm beneath the subperiosteal surface, and in portions of the bone this presents as an annulus or a double LAG (Fig. 3A). However, in this double LAG the interior LAG is more distinct than the outer LAG. There is no external fundamental system (EFS, a group of tightly spaced LAGs at the subperiosteal surface indicative that asymptotic size has been reached), and poorly vascularized parallel-fibered bone is present between the LAG and the subperiosteal surface (Fig. 3A).

The cortical bone of the tibia of GR 238 is broadly similar to that of the external tissues of the smaller femoral specimen (GR 1036) in that it comprises parallel-fibered primary
bone with longitudinally oriented osteons (Figs. 3G and 3H). However, unlike the smaller femur (GR 1036), there are no radial canals. The vascular canals are similarly sized (~10–30 µm in diameter) throughout and possess one to two lamellae. Vascular density decreases in the external portion of the cortex, shifting from well-vascularized to between well- and moderately-well vascularized about three-fourths of the distance to the subperiosteal surface. A portion of the cortex on the medial side of the tibia differs from the rest of the sample bone; here, the parallel-fibered bone is less organized and less well-vascularized, the vascular canals are both circular and elliptical in shape, and the vascular density (i.e., number of vascular canals in a given area) decreases whereas the size of the vascular canals increases (40–100 µm in long-axis diameter; see online supplementary full-slide images on Morphobank). Osteocyte lacunae are denser than in the small femur, and only a small majority of lacunae are preferentially oriented circumferentially (and these preferentially oriented lacunae are in the internalmost cortex; Fig. 3G). The size and preferred orientation of osteocyte lacunae does not change between internal and external cortical bone. There are between one and two LAGs present in the sections of this element. The interior LAG is the most clearly defined, and is only ~150 µm away from the medullary border because of cortical drift; it has been eliminated by cortical drift in many portions of the cortex. The external LAG is less clearly defined (Fig. 3H) and is not easily traceable to different regions of the cortex. There is highly disorganized woven-fibered bone with large (100–150 µm in diameter) primary osteons internal to the deepest primary cortical bone on the medial and lateral sides of the element (Fig. 3G). Although the large osteons may superficially resemble secondary or Haversian bone, this is compacted coarse cancellous bone (CCC; e.g., Chinsamy, 1995; Reid, 1996; Chinsamy-Turan, 2005) and is the result of cortical drift, as evidenced by a lack of cement lines around osteons (compare with cortical drift in the tibial histology of the silesaurid A. kongwe, Griffin & Nesbitt, 2016a; Fig. 4G). A clear boundary separates the CCC from the more external primary cortical bone (Fig. 3G); however, in most regions this boundary is a simple difference in tissue types, with only one portion of the boundary is formed by a band of avascular, lacunae-rich bone (similar to an annulus). Osteocyte lacunae in the CCC are large relative to the rest of the cortex and have no preferred orientation. Some trabecular fragments are preserved in the medullary cavity.

The sampled portion of the right fibula of GR 238 is comprised of predominantly primary parallel-fibered bone with primary osteons (generally with one lamellae), of overall similar composition, vascular density, and apatite crystallite organization to the tibia of this individual (Figs. 3D–3F). Unlike in the histology of the tibia, anastomosing vascular canals are not uncommon, although longitudinal canals still predominate, and these anastomoses impart a radial presentation to the canals (Figs. 3E and 3F). There are at least two LAGs that are traceable around the cortex except for a portion of the slide that has been ground too thin, and a portion where the internalmost LAG has been destroyed by medullary cavity migration/expansion. External to the second LAG, vascular density and connectivity decreases, and collagen fibers become more organized, with osteocytes oriented circumferentially along lamellae, although the cortex remains...
vascularized all the way to the edge of the subperiosteal surface (Figs. 3E and 3F). In the posterior and posteromedial portions of the cortex this decrease in vascular density and increase in organization is dramatic, with almost no vascular canals present, two additional LAGs visible, and highly organized parallel-fibered or even lamellar bone tissue containing abundant Sharpey’s fibers (Fig. 3D; Sharpey’s fibers not visible in figure, which focuses on the internal cortex; see online supplementary full-slide images on Morphobank). In addition to the LAGs, there are several weakly developed growth lines in the fibular cortex that do not persist around the entire circumference of the cortex (Figs. 3E and 3F); these may be either weak annual cycles or growth subcycles within a year constrained by true LAGs. The endosteal margin in this region also preserves several bands of avascular bone separating the medullary cavity from the cortical bone (Fig. 3D), although in other regions medullary cavity expansion or migration has apparently destroyed this avascular bone (Fig. 3E). A few secondary osteons are present just external to the endosteal margin on the posteromedial portion of the cortex (Fig. 3D).

The overall histology of the largest femur (GR 1038) is similar to that of the other sampled elements (Figs. 3I–3K). The majority of the sampled cortex is comprised of parallel-fibered primary bone, but the tissues possess an overall greater degree of organization than do those of the other elements. For example, most osteocyte lacunae are preferentially oriented in a circumferential direction except those of the internalmost sampled cortex (Figs. 3I–3K). Almost all vascular canals are ringed by a single layer of lamellar bone, and the vascular canals are almost exclusively longitudinal. Although the piece of cortex sampled from GR 1038 was an overhanging portion of the crushed midshaft, we hypothesize that the entire cortex, from endosteal lamellae to periosteal surface, is present in this sample. The internalmost portion of the sample in both slides 1 and 2 of GR 1038 comprises dense lamellar bone with large, globular osteocyte lacunae (Fig. 3J). This lamellar bone contains some simple vascular canals, and the interiormost surface is roughly parallel to the outermost surface where this lamellar bone is preserved. In slide 2, there is a small patch of more disorganized woven-fibered bone that abuts this lamellar bone on the internal side of the patch and grades into the normal cortical bone on the other (Fig. 3J). This lamellar bone possesses morphology consistent with that of endosteal lamellae. The vascular canals contained within the lamellar bone are also consistent with endosteal lamellae, as with several examples of vascularized endosteal lamellae that have recently been reported (Chinsamy, Cerda & Powell, 2016; Canoville et al., 2017; Prondvai et al., 2018). The line between the lamellar bone and the rest of the cortex is scalloped, indicating it was a resorption line on which the endosteal lamellae were then deposited, not smooth as would be expected in a LAG. In the outermost cortex the bone shifts to a highly organized parallel-fibered bone with decreased vascularization and circumferentially orientedapatite crystallites (Figs. 3I and 3K). Osteocyte lacunae are exclusively circumferentially oriented and elongate. There are two closely spaced LAGs within this parallel-fibered bone in the outermost region of the cortex, similar to GR 1036, and there appears to be a third LAG just beneath the subperiosteal surface (Fig. 3I). There are no secondary osteons in the sampled region of GR 1038, and no LAGs observed in the main body of the cortex.
DISCUSSION

Growth of *Dromomeron romeri*

There are no large, robust muscle scars present across the sampled size range of
*Dromomeron romeri* femora, although there are clear lineations showing the locations of
muscle attachments. The sole exception to this is the attachment of the *M. caudofemoralis*
musculature, which in most other early archosaurs forms a large flange- or wing-like
structure (= fourth trochanter; *Hutchinson, 2001*; *Nesbitt, 2011*) but in *D. romeri*
only forms a small scar, and only in the largest individual. Therefore, we hypothesize that
*D. romeri* did form a well-developed femoral scar through ontogeny, although only
one muscle attachment—that of the *M. caudofemoralis*—changed during development.
There is no evidence for the development of other well-developed femoral scars in
*D. romeri*; however, without considering histological data this absence could merely
indicate skeletal immaturity in the individuals sampled.

The histology of the three individuals sampled suggest that *D. romeri* reached the larger
sizes in our sample within at least one to 2 years of growth, although earlier LAGs
may easily have been destroyed by medullary cavity expansion, or not recorded (i.e., no
slow-down in growth), making this estimate a lower bound on the absolute ontogenetic age
of these individuals. Neither of the smaller individuals had reached asymptotic size
and were still growing, as indicated by a lack of an EFS in all elements sampled (*Cormack,
1987*; *Horner, de Ricqlès & Padian, 1999*; *Chinsamy-Turan, 2005*; *Huttenlocker,
*Woodward & Hall, 2013*). However, based on the increasing organization of more external
tissues and a decrease in vascular density and anastomoses in these regions, both
individuals had slowed linear growth from more rapid rates earlier in ontogeny, reaching
the stage of depressed growth rates prior to asymptotic size within a few years,
consistent with other studies of *D. romeri* histology (*Werning, 2013*). The size, histology,
and ontogenetic ages of these smaller individuals is similar to other individuals of
*D. romeri* that have been histologically sampled (*Werning, 2013*), suggesting that the
relationship between size and histological signal is consistent across this taxon.
The histology of the largest femur (GR 1038) suggests that this individual had nearly
reached full-size. The subperiosteal portion of the cortex is composed of poorly
vascularized parallel-fibered or lamellar bone with two to three closely spaced LAGs. If this
is an EFS this may indicate that asymptotic size had been reached in this individual.
However, the uneven spacing of the LAGs may suggest that this is not an EFS, but that the
individual deposited a triple LAG shortly before death. Regardless, it is clear that
linear growth had greatly slowed, if not completely ceased in this individual. Using these
data in conjunction with the bone scar data (see below) we hypothesize that the size
of the largest femur (GR 1038) provides a rough estimate on the maximum femoral size
reached by *D. romeri* at skeletal maturity, although the individual may have grown
slightly larger before reaching maximum size (especially if the three LAGs are not the onset
of an EFS). The smaller femur also possesses some histological indicators of slowed
growth (e.g., decreased vascularization), which could be indicative of intraspecific
variation in maximum body size in this taxon, or that growth had only slowed in this
individual as part of an annual cycle that ended with the deposition of a double LAG (Fig. 3A).

Given that the M. iliofemoralis externus, the insertion of which is hypothesized to form the trochanteric shelf of dinosauriforms (Hutchinson, 2001), does not form a distinct structure in D. romeri, the ‘trochanteric shelf’ of D. gregorii may have independently evolved in D. gregorii and dinosauriforms. Conversely, the trochanteric shelf may have been present in all skeletally mature lagerpetids except D. romeri, but the immaturity of all other known lagerpetid individuals prevents its observation. The muscles forming the anterior trochanter and trochanteric shelf (Mm. iliotrochantericus caudalis and iliofemoralis externus, respectively) have been hypothesized to be homologous with the crocodylian M. iliofemoralis, splitting from a single muscle into two at Dinosauriformes (Hutchinson, 2001). If the M. iliofemoralis insertion did shift proximally on the femur at the common ancestor of Dinosauria, and only split into two heads in Dinosauriformes (Nesbitt et al., 2009a), then this may be supported by the simultaneous appearance of the anterior trochanter and ‘trochanteric shelf’ in the ontogeny of D. gregorii, as well as the continuous structure formed by this scar(s). However, the non-ornithodiran avemetatarsalian Teleocrater rhadinus and other aphanosaurs possess a proximally located attachment for M. iliotrochantericus caudalis and a more distally located attachment for M. iliofemoralis externus (Nesbitt et al., 2017). Therefore, the split of the ancestral M. iliofemoralis has already occurred in this taxon, with only one portion of the split shifting proximally. This could indicate that 1) the split of M. iliofemoralis occurred at Aphanosauria + Dinosauria, and by Dinosauria, the M. iliofemoralis externus had also shifted proximally to insert next to M. iliotrochantericus caudalis in lagerpetids, with the continuous structure in Dromomeron representing the insertion of two muscles, or that 2) the split and proximal movement of M. iliofemoralis evolved independently in Teleocrater and Dinosauriformes. In the latter hypothesis, the anterior trochanter-trochanteric shelf structure of D. gregorii (and corresponding linear scars of other lagerpetids) would be the insertion of a single muscle, M. iliofemoralis. Both scenarios are plausible based on the phylogenetic relationships and optimizations of character states.

Based on these data, we hypothesize that the absence of an anterior trochanter and other scars of the proximal end of the femur in D. romeri is an autapomorphic feature of this taxon, and not simply indicative of the morphological immaturity of known individuals of D. romeri, falsifying the hypothesis (Bennett, 2015) that D. romeri is an immature Tawa hallae. The absence of a fourth trochanter (i.e., ridge for attachment of M. caudifemoralis) is especially striking. This feature is widely present among early archosaurs and is one of the first well-developed muscle scars to form during ontogeny, present before all other femoral scars in early avemetatarsalians (e.g., Teleocrater, Nesbitt et al., 2017; D. gregorii, Nesbitt et al., 2009a; Ixalerpeton, Cabreira et al., 2016; Silesaurus, Piechowski, Talanda & Dzik, 2014; Asilisaurus, Griffin & Nesbitt, 2016a) and pseudosuchians (e.g., hatchling Alligator, Brochu, 1992). Furthermore, the anterior trochanter consistently appears early in post-natal ontogeny among early bird-line archosaurs (Nesbitt et al., 2009a; Griffin & Nesbitt, 2016a; Griffin, 2018), so even though
the sampled individuals of *D. romeri* had not completely ceased growth, the absence of this feature in individuals that are close to reaching histological maturity suggests that it is absent in all individuals of this taxon, regardless of maturity.

**Comparison with other dinosauromorphs**

Although many immature individuals of early bird-line archosaurs lack an anterior trochanter (e.g., *Dromomeron gregorii*, Nesbitt et al., 2009a; *Asilisaurus*, Griffin & Nesbitt, 2016a), the absence of an anterior trochanter and fourth trochanter throughout ontogeny is unique to *D. romeri* among known Triassic dinosauromorphs. *Lagerpeton chanarensis* possesses both a large, aliform fourth trochanter and an anterior trochanter in the form of a low rugosity (Sereno & Arcucci, 1994). Larger individuals of *D. gregorii* (Nesbitt et al., 2009a) possess a fourth trochanter and a ridge that is probably homologous with the ‘obturator ridge’ of early theropods and *Asilisaurus* (Griffin, 2018), as well as an anterior trochanter. *D. gigas* possesses a large anterior trochanter and ‘obturator ridge’, as well as a large scar marking the insertion of the *M. caudofemoralis longus*; only the proximal end of this scar is preserved, but it was interpreted as lacking a true fourth trochanter (Martínez et al., 2016), possessing a flat scar more similar to what we have described in *D. romeri*. Although *Ixalerpeton polesinensis* lacks an anterior trochanter, it possesses a robust aliform fourth trochanter, and a small scar similar in location (Cabreira et al., 2016: Fig. S1; ULBRA-PVT059) to the insertion of *M. caudofemoralis brevis* in dinosauriforms (Griffin & Nesbitt, 2016a; Griffin, 2018). Unfortunately, growth series are only known from *D. romeri* and *D. gregorii*, so broader comparisons between ontogenetic changes in femoral histology and gross morphology among lagerpetids are unavailable.

Intuitively, the morphology, size, and surface area of a muscle attachment would seem to be related to activity levels of the muscle, with larger or more robust scars indicating increased stress. However, this assumption has little empirical support: studies indirectly testing this assumption are often inconclusive and focused on humans (Foster, Buckley & Tayles, 2014), and the only experimental test of this hypothesis, in adult sheep, found no relationship between muscle activity and scar size (Zumwalt, 2006). Muscle activity is known to play a major role in the formation of overall bone morphology early in development, with paralyzed chick embryos possessing splayed and distorted limb bones (e.g., Hall & Herring, 1990; Hosseini & Hogg, 1991; Lamb et al., 2003), but this is a distinct process from the post-natal formation of well-developed bone scars, and the two may not be related. How muscle activity relates to the formation of muscle scars in non-mammalian taxa, or during ontogeny, remains poorly understood. Therefore, the relationship between these factors and archosauromian bone scars is unclear, so this lack of scarring in *D. romeri* is not evidence of a difference in activity level or body mass between this taxon and other lagerpetids. Whether differences in muscle scars relate to disparity in locomotion remains to be tested.

To date, there have been no histological investigations of any lagerpetids besides *D. romeri* (Werning, 2013; this study) because of a scarcity of material. However, the bone histology of the aphanosaur *Teledocrater* (Nesbitt et al., 2017), the early pterosaurs...
Eudimorphodon, Dimorphodon, and Dorygnathus (Padian, Horner & de Ricqlès, 2004), the silesaurids Asilisaurus (Griffin & Nesbitt, 2016a) and Silesaurus (Fostowicz-Frelık & Sulej, 2010), the possible dinosaur or close dinosaurian relative Nyasasaurus (Nesbitt et al., 2013), and many early dinosaurs (e.g., Herrerasaurus ichnigraphastensis, de Ricqlès, Padian & Horner, 2003; Padian, Horner & de Ricqlès, 2004; Tawa hallae, Werning, 2013; Coelophysis bauri, Colbert, 1995; Nesbitt et al., 2006; Megapnosaurus rhodesiensis, Raath, 1977; Chinsamy, 1990, Werning, 2013; Lesothosaurus, Knoll, Padian & de Ricqlès, 2010; Scutellosaurus, Padian, Horner & de Ricqlès, 2004) have all been described, allowing for broad comparison of the growth of D. romeri with other early avemetatarsalians.

The histology of Teleocrater (fibula and humerus) and D. romeri (femora, tibia, and fibula) are broadly similar: both possess cortices consisting of woven bone with longitudinal vascular canals and primary osteons and are more similar to each other than to stem archosaurs (Botha-Brink & Smith, 2011; Werning, 2013). However, much of the outer cortex of D. romeri possesses more organized collagen fibers, osteocyte lacunae preferentially oriented circumferentially, lower vascularization, fewer anastomoses, and are more similar to the histology of stem archosaurs (e.g., Botha-Brink & Smith, 2011). In contrast, the cortex of sampled elements of Teleocrater possesses more disorganized woven-fibered bone with a higher number of anastomoses, especially in the humerus (Nesbitt et al., 2017). Additionally, unlike both Teleocrater and other descriptions of D. romeri histology (Werning, 2013), the fibula of the largest individual of D. romeri possesses secondary osteons, although this could be related to the element sampled (fibulae of Dromomeron have not been previously sampled), or the ontogenetic age of the individual. The histology of Teleocrater suggests a more sustained high rate of growth over a longer time. Whereas much of the outer cortex of D. romeri possesses features that suggest slowing growth (e.g., more organized collagen fibers, osteocyte lacunae preferentially oriented circumferentially, lower vascularization, fewer anastomoses), the cortex of sampled elements of Teleocrater possesses more disorganized and vascularized woven-fibered bone (Nesbitt et al., 2017). All these features suggest that, although at their highest rates of growth D. romeri and Teleocrater present similar histology, this rate of growth appears to be more sustained in Teleocrater. This shorter period of sustained growth in D. romeri may be related to its smaller body size—that is, D. romeri simply did not have to grow for as long to reach maximum size. This shorter and lower rate of sustained growth is also true in comparison with the histology of silesaurids, which possess larger regions of disorganized woven bone and more anastomosing vascular canals than are present in D. romeri, although the vascularization level of silesaurids is similar to that of the inner cortex of D. romeri (Fostowicz-Frelık & Sulej, 2010; Griffin & Nesbitt, 2016a).

The Late Triassic pterosaur Eudimorphodon possesses more organized bone tissues than D. romeri, with highly organized parallel-fibered tissue and a moderate level of vascularization with rarely anastomosing longitudinal canals (Padian, Horner & de Ricqlès, 2004; Werning, 2013), all indicating a slower rate of growth. However, because of the thin pterosaurian bone cortex, the presumably faster-growing tissue from earlier in ontogeny has been eroded, and confidently assessing the maturity of the sampled individual is difficult (Werning, 2013). Dimorphodon, a pterosaur from the Early Jurassic,
possesses mainly disorganized woven-fibered bone similar to *Dromomeron romeri*, with some areas of more parallel-fibered bone. The degree of vascularization is slightly lower than that of *D. romeri*, but with a larger number of anastomosing vascular canals and looser tissue organization (*de Ricqlès et al., 2000; Padian, Horner & de Ricqlès, 2004; Werning, 2013*). These bone tissues suggest a higher rate of growth than that of *D. romeri*. The bone histology of the dinosauriform *Nyasasaurus* shows much higher levels of tissue disorganization, level of vascularity, anastomoses, and osteocyte density (*Nesbitt et al., 2013*) which is similar to that of early dinosaurs (e.g., *Herrerasaurus ischigualastensis, de Ricqlès, Padian & Horner, 2003; Padian, Horner & de Ricqlès, 2004; Tawa hallae, Werning, 2013; Coelophysis bauri, Colbert, 1995; Nesbitt et al., 2006; Megapnosaurus rhodesiensis, Raath, 1977; Chinsamy, 1990, Werning, 2013; Lesothosaurus, Knoll, Padian & de Ricqlès, 2010; Scutellosaurus, Padian, Horner & de Ricqlès, 2004; see Werning, 2013 for a comparative review of early dinosaur osteohistology*). Therefore, the bone histology of sampled early dinosaurs and their close relatives also suggests sustained growth proceeding more rapidly than in *D. romeri*.

All taxa discussed, including *D. romeri* and excepting *Eudimorphodon*, shift from predominantly disorganized woven-fibered to highly organized parallel-fibered bone tissue later in ontogeny as growth slows, and all but *Asilisaurus, Nyasasaurus*, and the pterosaurs deposit LAGs. Based on this, *Dromomeron romeri* appears to have possessed a growth rate most similar to *Teleocrater rhadinus*, although sustained over a shorter period of time, with a maximum rate of growth that was slower than most pterosaurs, dinosauriforms, and dinosaurs but faster than what is known for *Eudimorphodon*.

**Implications for assessing skeletal maturity in early avemetatarsalians**

Although *Dromomeron romeri* lacks many femoral bone scars that are present in skeletally mature individuals of many other early avemetatarsalians, one scar—the attachment of *M. caudofemoralis longus*—does appear during post-natal ontogeny in this taxon. This is consistent with the general trend of increasing number and size of muscle scars with increasing skeletal maturity observed in other early avemetatarsalians (*Nesbitt et al., 2009a; Piechowski, Talanda & Dzik, 2014; Griffin & Nesbitt, 2016a, 2016b; Griffin, 2018*) and in their closest living relatives (*Brochu, 1992, 1996; Tumarkin-Deratzian, Vann & Dodson, 2006; 2007; Griffin & Nesbitt, 2016b*). Therefore, the sampled series of *D. romeri* provides more evidence that the presence of bone scars is a reliable indicator of increasing skeletal maturity among archosaurs in general, and especially for early avemetatarsalians.

That said, the presence of a single bone scar or even a small set of scars may not be indicative that full skeletal maturity (i.e., cessation of linear growth; presence of all morphological features of maturity) has been reached in a given individual. Because bone scars appear in a stepwise pattern during ontogeny, and not simultaneously (*Brochu, 1992; Tumarkin-Deratzian, Vann & Dodson, 2006, 2007; Nesbitt et al., 2009a; Piechowski, Talanda & Dzik, 2014; Griffin & Nesbitt, 2016a, 2016b; Griffin, 2018*), and because an individual may develop bone scars before histological maturity and attainment of asymptotic size (*Griffin & Nesbitt, 2016a*; this study), the presence of a single scar is a better
indicator of relative skeletal maturity in comparison to conspecific individuals or those of close relatives, rather than as an absolute method of determining skeletal maturity. For example, an individual of *D. gregorii* possessing an anterior trochanter can be reasonably assumed to have progressed further in ontogeny than an individual lacking this scar. Because both taxa are closely related, and both are known to develop the anterior trochanter, an individual of *D. gigas* possessing this same scar can likewise be assumed to have attained a higher level of maturity than an individual of *D. gregorii* that lacks this scar. However, this does not indicate that all individuals of *D. gregorii* and *D. gigas* that possess the anterior trochanter are at the same ontogenetic stage or level of skeletal maturity.

Characters useful for maturity assessment vary across clades, and our confidence in assessment based on characters should vary with phylogenetic proximity to well-taxa with established growth patterns (e.g., *Irmis, 2007*). The anterior trochanter appears early in the ontogeny of dinosauriforms (*Griffin, 2018*), but in lagerpetids appears later in ontogeny (*D. gregorii, Nesbitt et al., 2009a*), or potentially not at all (*Ixalerpeton, Cabreira et al., 2016*), although this individual could be immature; *D. romeri*, this study), and is apparently absent in aphanosaurs (with a low mound marking this muscle insertion instead, *Nesbitt et al., 2017*). Therefore, the absence of an anterior trochanter in an individual may: 1) be uninformative for assessing skeletal maturity (in aphanosaurs); 2) indicate that the individual is skeletally immature, but with histological corroboration necessary (in lagerpetids); or 3) be strongly indicative that the individual is in the earliest stages of skeletal maturity (in dinosauriforms). This high degree of inter- and intraspecific variation in the anterior trochanter, trochanteric shelf, and potentially the fourth trochanter among early avemetatarsalians may be suggestive of a ‘zone of developmental variability’ (a complex pattern of character gains and losses before character fixation; see *Bever, Gauthier & Wagner, 2011; Fig. 4*). However, more work must be done to determine whether the absence of these scars in some lagerpetids is simply from the immaturity of individuals, or if there is true variability in the presence of these characters. Therefore, although the presence of muscle scars is indicative of increasing skeletal maturity, the phylogenetic history of each feature must be considered when assessing relative maturity (Fig. 4), and in some clades corroboration with other characters or osteohistology may be necessary.

Intraspecific variation in ontogenetic patterns may also complicate assessing maturity via gross morphology. There appear to be multiple sequences by which morphological characters progress from immature to mature states among silesaurids and early dinosaurs (*Griffin & Nesbitt, 2016a, 2016b; Barta, Nesbitt & Norell, 2017; Griffin, 2018*), so that the presence of a muscle scar or scars may not indicate the same level of maturity in all individuals, because the relative timing and body size at scar appearance may be widely disparate. However, whether this high intraspecific variation is constrained to Dinosauriformes or is more widespread among early diverging avemetatarsalians has not been explored. We did not observe similar levels of intraspecific variation among the growth series of *D. romeri*, either in gross morphology or histology as they relate to
femoral size, and such variation has also not been reported in D. gregorii, the only other lagerpetid known from a growth series. Additionally, the five femoral specimens of Lagerpeton chanarensis are of roughly equivalent sizes, but also show no variation like that present in similarly sized early dinosauriform individuals (Nesbitt et al., 2009a). Therefore, we tentatively hypothesize that this high intraspecific variation in growth patterns is restricted to Dinosauriformes, although larger sample sizes of lagerpetid growth series will provide better testing of this hypothesis. Alternatively, if the intraspecific variation reported in some later pterosaurs (e.g., Rhamphorhynchus, Prondvai et al., 2012) is homologous with that in early dinosauriforms, this would imply that this high variation extends to at least the common ancestor of pterosaurs, lagerpetids, and dinosaurs.

CONCLUSIONS

Unlike many other early avemetatarsalians, even the largest known individuals of Dromomeron romeri do not possess large, robust femoral bone scars. Because bone histology—generally consisting of more organized woven-fibered and parallel-fibered primary bone, longitudinal vascular canals, and one to two preserved LAGs—indicates that these large individuals are close to reaching asymptotic size, and are therefore near skeletal maturity, this absence of bone scars is not the result of the immaturity of the individuals in question. Although these features are important for assessing maturity in many groups, phylogenetic history must be considered when utilizing femoral scars to assess skeletal maturity. Complicating the matter is intraspecific variation—although there is currently no evidence in lagerpetids for levels of intraspecific variation as high as has been observed in dinosauriforms, small sample sizes may be hindering observation of this signal.
INSTITUTIONAL ABBREVIATIONS
GR  Ghost Ranch Ruth Hall Museum of Paleontology, Abiquiu, NM, USA.
TMM  Vertebrate Paleontology Laboratory, Texas Natural Science Center, Austin, TX, USA.
ULBRA  Universidade Luterana do Brasil, Coleção de Paleovertebrados, Canoas, Rio Grande do Sul, Brazil.

ACKNOWLEDGEMENTS
We thank the many Ghost Ranch field volunteers and students for participating in excavations that recovered these Dromomeron romeri specimens, S. Werning for helpful discussion and use of microscope and K. Stein, Z. Morris, and J. Olori for discussion, and we thank A. Downs for facilitating many aspects of this work. This study was greatly improved by the input of three anonymous reviewers.

ADDITIONAL INFORMATION AND DECLARATIONS
Funding
This work was supported by National Science Foundation (NSF) EAR 1349667 (awarded to Sterling J. Nesbitt), EAR 1349654 (awarded to Alan H. Turner), EAR 1349554 (awarded to Nathan D. Smith), EAR 1349650 (awarded to Randall B. Irmis), the University of Utah (to Randall B. Irmis), an NSF Graduate Research Fellowship (awarded to Christopher T. Griffin), and a Geological Society of America Southeastern Section Undergraduate Student Research Grant (awarded to Lauren S. Bano). 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:
National Science Foundation (NSF): EAR 1349667, EAR 1349654, EAR 1349554, EAR 1349650.
University of Utah.
NSF Graduate Research Fellowship.
Geological Society of America Southeastern Section Undergraduate Student Research Grant.

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

Author Contributions
• Christopher T. Griffin conceived and designed the experiments, performed the experiments, analyzed the data, prepared figures and/or tables, authored or reviewed drafts of the paper, approved the final draft.
• Lauren S. Bano conceived and designed the experiments, performed the experiments, analyzed the data, prepared figures and/or tables, authored or reviewed drafts of the paper, approved the final draft.
• Alan H. Turner authored or reviewed drafts of the paper, approved the final draft.
• Nathan D. Smith authored or reviewed drafts of the paper, approved the final draft.
• Randall B. Irmis authored or reviewed drafts of the paper, approved the final draft.
• Sterling J. Nesbitt conceived and designed the experiments, performed the experiments, analyzed the data, contributed reagents/materials/analysis tools, authored or reviewed drafts of the paper, approved the final draft.

Field Study Permissions
The following information was supplied relating to field study approvals (i.e., approving body and any reference numbers):
Fieldwork at Ghost Ranch was conducted under permission from Ghost Ranch Conference Center.

Data Availability
The following information was supplied regarding data availability:
Morphobank (Project Number 3127: http://morphobank.org/permalink/?P3127).

REFERENCES
Abdala F, Giannini NP. 2000. Gomphodont cynodonts of the Chañares Formation: the analysis of an ontogenetic sequence. Journal of Vertebrate Paleontology 20(3):501–506 DOI 10.1671/0272-4634(2000)020[0501:GCOTCA]2.0.CO;2.
Bailleul AM, Scannella JB, Horner JR, Evans DC. 2016. Fusion patterns in the skulls of modern archosaurs reveal that sutures are ambiguous maturity indicators for the Dinosauria. PLOS ONE 11(2):e0147687 DOI 10.1371/journal.pone.0147687.
Barta DE, Nesbitt SJ, Norell MA. 2017. The evolution of the manus of early theropod dinosaurs is characterized by high inter- and intraspecific variation. Journal of Anatomy 232(1):80–104 DOI 10.1111/joa.12719.
Bennett SC. 2001. The osteology and functional morphology of the Late Cretaceous pterosaur Pteranodon. Palaeontographica Abteilung A 260:113–153.
Bennett SC. 2015. An external mandibular fenestra and other archosauriform characters in basal pterosaurs re-examined. Historical Biology 27(6):796–814 DOI 10.1080/08912963.2015.1036257.
Benson RBJ, Campione NE, Carrano MT, Mannion PD, Sullivan C, Upchurch P, Evans DC. 2014. Rates of dinosaur body mass evolution indicate 170 million years of sustained ecological innovation on the avian stem lineage. PLOS Biology 12(5):e1001853 DOI 10.1371/journal.pbio.1001853.
Bever GS, Gauthier JA, Wagner GP. 2011. Finding the frame shift: digit loss, developmental variability, and the origin of the avian hand. Evolution & Development 13(3):269–279 DOI 10.1111/j.1525-142X.2011.00478.x.
Billet G, de Muizon C, Mamani Quispe B. 2008. Late Oligocene mesotheriids (Mammalia, Notoungulata) from Salla and Lacayani (Bolivia): implications for basal mesotheriid phylogeny Griffin et al. (2019), PeerJ, DOI 10.7717/peerj.6331
and distribution. Zoological Journal of the Linnean Society 152(1):153–200
DOI 10.1111/j.1096-3642.2007.00388.x.

Botha-Brink J, Smith RMH. 2011. Osteohistology of the Triassic archosauromorphs Prolacerta, Proterosuchus, Euparkeria, and Erythrosuchus from the Karoo Basin of South Africa. Journal of Vertebrate Paleontology 31(6):1238–1254 DOI 10.1080/07224634.2011.621797.

Brinkman D. 1988. Size-independent criteria for estimating relative age in Ophiacodon and Dimetrodon (Reptilia, Pelycosauria) from the Admiral and lower Belle Plains formations of west-central Texas. Journal of Vertebrate Paleontology 8(2):172–180 DOI 10.1080/02724634.1988.10011695.

Brochu CA. 1992. Ontogeny of the postcranial in crocodylomorph archosaurs Master's Master's thesis. University of Texas at Austin.

Brochu CA. 1996. Closure of neurocentral sutures during crocodilian ontogeny: implications for maturity assessment in fossil archosaurs. Journal of Vertebrate Paleontology 16(1):49–62 DOI 10.1080/02724634.1996.10011283.

Brown CM, Russell AP, Ryan MJ. 2009. Pattern and transition of surficial bone texture of the centrosaurine frill and their ontogenetic and taxonomic implications. Journal of Vertebrate Paleontology 29(1):132–141 DOI 10.1080/02724634.1996.10011283.

Cabreira SF, Kellner AWA, Dias-de-Silva S, LRd Silva, Bronzato M, JCDa Marsola, Müller RT, JSD Bittencourt, Batista BJ, Raugust T, Carrilho R, Brodt A, Langer MC. 2016. A unique Late Triassic dinosauromorph assemblage reveals dinosaur ancestral anatomy and diet. Current Biology 26(22):3090–3095 DOI 10.1016/j.cub.2016.09.040.

Canoville A, Yang T-R, Zanno L, Zheng W, Schweitzer MH. 2017. Paleohistology of a gravid oviraptorosaurian dinosaur from the Upper Cretaceous Nanxiong Formation, China, with an enigmatic endosteal tissue. Journal of Vertebrate Paleontology, Programs and Abstracts 37:93.

Carrano MT. 2006. Body-size evolution in the dinosauria. In: Carrano MT, Gaudin TJ, Blob RW, Wible JR, eds. Amniote Paleobiology: Perspectives on the Evolution of Mammals, Birds, and Reptiles: a Volume Honoring James Allen Hopson. Chicago: University of Chicago Press, 225–268.

Carrano MT, Hutchinson JR. 2002. Pelvic and hindlimb musculature of Tyrannosaurus rex (Dinosauria: Theropoda). Journal of Morphology 253(3):207–228 DOI 10.1002/jmor.10018.

Cartelle C, De Iuliis G. 2006. Eremotherium laurillardi (Lund) (Xenarthra, Megatheriidae), the Panamerican giant ground sloth: taxonomic aspects of the ontogeny of skull and dentition. Journal of Systematic Palaeontology 4(2):199–209 DOI 10.1017/S14777201905001781.

Chinsamy A. 1990. Physiological implications of the bone histology of Syntarsus rhodesiensis (Saurischia: Theropoda). Palaeontologia Africana 27:77–82.

Chinsamy A. 1995. Ontogenetic changes in the bone histology of the Late Jurassic ornithopod Dryosaurus lettowvorbecki. Journal of Vertebrate Paleontology 15(1):96–104 DOI 10.1080/02724634.1995.10011209.

Chinsamy A, Cerda I, Powell J. 2016. Vascularised endosteal bone tissue in armoured sauropod dinosaurs. Scientific Reports 6(1):24858 DOI 10.1038/srep24858.

Chinsamy-Turan A. 2005. The microstructure of dinosaur bone. Baltimore: Johns Hopkins University Press.

Codron D, Carbone C, Müller DWH, Clauss M. 2012. Ontogenetic niche shifts in dinosaurs influenced size, diversity and extinction in terrestrial vertebrates. Biology Letters 8(4):620–623 DOI 10.1098/rsbl.2012.0240.

Colbert EH. 1995. The little dinosaurs of ghost ranch. New York: Columbia University Press.
Cormack D. 1987. *Ham's histology*. New York: Lippincott.

Cubo J, Roy NL, Martinez-Maza C, Montes L. 2012. Paleohistological estimation of bone growth rate in extinct archosaurs. *Paleobiology* 38(2):335–349 DOI 10.1666/08093.1.

Dawson AB. 1925. The age order of epiphyseal union in the long bones of the albino rat. *Anatomical Record* 31(1):1–17 DOI 10.1002/ar.1090310102.

de Ricqlès AJ, Padian K, Horner JR, Francillon-Vieillot H. 2000. Palaeohistology of the bones of pterosaurs (Reptilia: Archosauria): anatomy, ontogeny, and biomechanical implications. *Zoological Journal of the Linnean Society* 129(3):349–385 DOI 10.1111/j.1096-3642.2000.tb00016.x.

Erickson GM. 2014. On dinosaur growth. *Annual Review of Earth and Planetary Sciences* 42(1):675–697 DOI 10.1146/annurev-earth-060313-054858.

Erickson GM, Mackovicky PJ, Currie PJ, Norell MA, Yerby SA, Brochu CA. 2004. Gigantism and comparative life-history parameters of tyrannosaurid dinosaurs. *Nature* 430(7001):772–775 DOI 10.1038/nature02699.

Erickson GM, Rogers KC, Varricchio DJ, Norell MA, Xu X. 2007. Growth patterns in brooding dinosaurs reveals the timing of sexual maturity in non-avian dinosaurs and genesis of the avian condition. *Biology Letters* 3(5):558–561 DOI 10.1098/rsbl.2007.0254.

Feldmann RM. 1989. Whitening fossils for photographic purposes. *Paleontological Society Special Publication* 4:342–346 DOI 10.1017/S0753-3969(03)00005-3.

Foster A, Buckley H, Tayles N. 2014. Using enthesis robusticity to infer activity in the past: a review. *Journal of Archaeological Method and Theory* 21(3):511–533 DOI 10.1007/s10816-012-9156-1.

Fostowicz-Frelik Ł, Sulej T. 2010. Bone histology of *Silesaurus opolensis* Dzik, 2003 from the Late Triassic of Poland. *Lethaia* 43(2):137–148 DOI 10.1111/j.1502-3931.2009.00179.x.

Frýdlová P, Nutilová V, Dudák J, Žemlička J, Němec P, Velenský P, Jirásek T, Frynta D. 2017. Patterns of growth in monitor lizards (Varanidae) as revealed by computed tomography of femoral growth plates. *Zoomorphology* 136(1):95–106 DOI 10.1007/s00435-016-0338-3.

Green OR. 2001. *A manual of practical laboratory and field techniques in palaeobiology*. Dordrecht: Kluwer Academic Publishers, 471–530.

Griffin CT. 2018. Developmental patterns and variation among early theropods. *Journal of Anatomy* 232(4):604–640 DOI 10.1111/joa.12775.

Griffin CT, Nesbitt SJ. 2016a. The histology and femoral ontogeny of the Middle Triassic (?late Anisian) dinosauriform *Asilisaurus kongwe* and implications for the growth of early dinosaurs. *Journal of Vertebrate Paleontology* 36(3):e111224 DOI 10.1080/07224634.2016.1111224.

Griffin CT, Nesbitt SJ. 2016b. Anomalously high variation in postnatal development is ancestral for dinosaurs but lost in birds. *Proceedings of the National Academy of Sciences of the United States of America* 113:14757–14762 DOI 10.1073/pnas.1613813113.

Haines RW. 1969. Epiphyses and sesamoids. In: Gans C, AdA Bellairs, Parson TS, eds. *Biology of the Reptilia Morphology A*. Vol. 1. London: Academic Press, 81–115.
Hall BK, Herring SW. 1990. Paralysis and growth of the musculoskeletal system in the embryonic chick. *Journal of Morphology* 206(1):45–56 DOI 10.1002/jmor.1052060105.

Hegna TA. 2010. Photography of soft-bodied crustaceans via drying, whitening, and splicing. *Journal of Crustacean Biology* 30(3):351–356 DOI 10.1651/09-3253.1.

Hone DWE, Farke AA, Wedel MJ. 2016. Ontogeny and the fossil record: what, if anything, is an adult dinosaur? *Biology Letters* 12(2):20150947 DOI 10.1098/rsbl.2015.0947.

Horner JR, de Ricqlès A, Padian K. 1999. Variation in dinosaur skeletochronology indicators: implications for age assessment and physiology. *Paleobiology* 25(03):295–304 DOI 10.1017/S0094837300021308.

Horner JR, de Ricqlès A, Padian K. 2000. Long bone histology of the hadrosaurid dinosaur *Maiasaura peeblesorum*: growth dynamics and physiology based on an ontogenetic series of skeletal elements. *Journal of Vertebrate Paleontology* 20(1):115–129 DOI 10.1671/0272-4634(2000)020[0115:LBHOTH]2.0.CO;2.

Horner JR, Padian K, de Ricqlès A. 2001. Comparative osteohistology of some embryonic and perinatal archosaurs: developmental and behavioral implications for dinosaurs. *Paleobiology* 27:39–58 DOI 10.1666/0094-8373(2001)027<0039:COOSEA>2.0.CO;2.

Hosseini A, Hogg DA. 1991. The effects of paralysis on skeletal development in the chick embryo. I. General effects. *Journal of Anatomy* 177:159–168.

Hutchinson JR. 2001. The evolution of femoral osteology and soft tissues on the line to extant birds (Neornithes). *Zoological Journal of the Linnean Society* 131(2):169–197 DOI 10.1111/j.1096-3642.2001.tb01314.x.

Huttenlocker AK, Rega E. 2012. The paleobiology and bone microstructure of pelycosaurian-grade synapsids. In: Chinsamy-Turan A, ed. *Forerunners of Mammals*. Bloomington: Indiana University Press, 90–119.

Huttenlocker AK, Woodward H, Hall BK. 2013. The biology of bone. In: Padian K, Lamm E-T, eds. *Bone Histology of Fossil Tetrapods: Advancing Methods, Analysis, and Interpretation*. Berkeley: University of California Press, 13–34.

Irmis RB. 2007. Axial skeleton ontogeny in the Parasuchia (Archosauria: Pseudosuchia) and its implications for ontogenetic determination in archosaurs. *Journal of Vertebrate Paleontology* 27(2):350–361 DOI 10.1671/0272-4634(2007)27[350:ASOITP]2.0.CO;2.

Irmis RB, Nesbitt SJ, Padian K, Smith ND, Turner AH, Woody D, Downs A. 2007. A Late Triassic dinosauromorph assemblage from New Mexico and the rise of dinosaurs. *Science* 317(5836):358–361 DOI 10.1126/science.1143325.

Jans MM. 2008. Microbial bioerosion of bone – a review. In: Wisshak M, Tapanila L, eds. *Current Developments in Bioerosion*. Berlin: Springer, 397–413.

Klein N, Sander PM. 2008. Ontogenetic stages in the long bone histology of sauropod dinosaurs. *Paleobiology* 34(2):247–263 DOI 10.1666/0094-8373(2008)034[0247:OSITLB]2.0.CO;2.

Knoll F, Padian K, de Ricqlès A. 2010. Ontogenetic change and adult body size of the early ornithischian dinosaur *Lesothosaurus diagnosticus*: Implications for basal ornithischian taxonomy. *Gondwana Research* 17(1):171–179 DOI 10.1016/j.gr.2009.03.010.

Köhler M, Moya-Sola S. 2009. Physiological and life history strategies of a fossil large mammal in a resource-limited environment. *Proceedings of the National Academy of Sciences of the United States of America* 106:20354–20358.

Lamb KE, Lewthwaite JC, Lin J-P, Simon D, Kavanagh E, Wheeler-Jones CPD, Pitsillides AA. 2003. Diverse range of fixed positional deformities and bone growth restraint provoked by flaccid paralysis in embryonic chicks. *International Journal of Experimental Pathology* 84(4):191–199 DOI 10.1046/j.1365-2613.2003.00353.x.
Langer MC, Benton MJ. 2006. Early dinosaurs: a phylogenetic study. *Journal of Systematic Palaeontology* 4(4):309–358 DOI 10.1017/S1477201906001970.

Lee AH, Werning S. 2008. Sexual maturity in growing dinosaurs does not fit reptilian growth models. *Proceedings of the National Academy of Sciences of the United States of America* 105:582–587.

Legendre LJ, Guénard G, Botha-Brink J, Cubo J. 2016. Palaeohistological evidence for ancestral high metabolic rate in archosaurs. *Systematic Biology* 65(6):989–996 DOI 10.1093/sysbio/syw033.

Maidment SCR, Barrett PM. 2011. The locomotor musculature of basal ornithischian dinosaurs. *Journal of Vertebrate Paleontology* 31(6):1265–1291 DOI 10.1080/02724634.2011.606857.

Martínez RN, Apaldetti C, Correa GA, Abelín D. 2016. A Norian lagerpetid dinosauromorph from the Quebrada del Barro Formation, northwestern Argentina. *Ameghiniana* 53(1):1–13 DOI 10.5710/AMGH.21.06.2015.2894.

Moran MM, Bajpai S, George JC, Suydam R, Usip S, Thewissen JGM. 2015. Intervertebral and epiphyseal fusion in the postnatal ontogeny of cetaceans and terrestrial mammals. *Journal of Mammal Evolution* 22(1):93–109 DOI 10.1007/s10914-014-9256-7.

Morris P. 1972. A review of mammalian age determination methods. *Mammal Review* 2(3):69–104 DOI 10.1111/j.1365-2907.1972.tb00160.x.

Müller RT. 2017. Are the dinosauromorph femora from the Upper Triassic of Hayden Quarry (New Mexico) three stages in a growth series of a single taxon?. *Anais da Academia Brasileira de Ciências* 89(2):835–839 DOI 10.1590/0001-3765201720160583.

Müller RT, Langer MC, Dias-da-Silva S. 2018. Ingroup relationships of Lagerpetidae (Avemetatarsalia: Dinosauromorpha): a further phylogenetic investigation on the understanding of dinosaur relatives. *Zootaxa* 4392(1):149–158 DOI 10.11646/zootaxa.4392.1.7.

Nesbitt SJ. 2011. The early evolution of archosaurs: relationships and the origin of major clades. *Bulletin of the American Museum of Natural History* 352:1–292 DOI 10.1206/352.1.

Nesbitt SJ, Barrett PM, Werning S, Sidor CA, Charig AJ. 2013. The oldest dinosaur? A Middle Triassic dinosauriform from Tanzania. *Biology Letters* 9(1):20120949 DOI 10.1098/rsbl.2012.0949.

Nesbitt SJ, Butler RJ, Ezcurra MD, Barrett PM, Stocker MR, Angielczyk KD, Smith RMH, Sidor CA, Niedźwiedzki G, Sennikov AG, Charig AJ. 2017. The earliest bird-line archosaurs and the assembly of the dinosaur body plan. *Nature* 544(7651):484–487 DOI 10.1038/nature22037.

Nesbitt SJ, Butler RJ, Ezcurra MD, Charig AJ, Barrett PM. 2018. The anatomy of *Teleocrater rhadinus*, an early avemetatarsalian from the lower portion of the Lufia Member of the Manda Beds (~Middle Triassic). In: Sidor CA, Nesbitt SJ, eds. *Vertebrate and Climatic Evolution in the Triassic Rift Basins of Tanzania and Zambia*, 142–177.

Nesbitt SJ, Irmis RB, Parker WG, Smith ND, Turner AH, Rowe T. 2009a. Hindlimb osteology and distribution of basal dinosauromorphs from the Late Triassic of North America. *Journal of Vertebrate Paleontology* 29(2):498–516 DOI 10.1671/039.029.0218.

Nesbitt SJ, Smith ND, Irmis RB, Turner AH, Downs A, Norell MA. 2009b. A complete skeleton of a Late Triassic saurischian and the early evolution of dinosaurs. *Science* 326(5959):1530–1533 DOI 10.1126/science.1180350.

Nesbitt SJ, Turner AH, Erickson GM, Norell MA. 2006. Prey choice and cannibalistic behaviour in the theropod *Coelophysis*. *Biology Letters* 2(4):611–614 DOI 10.1098/rsbl.2006.0524.

Nilsson O, Baron J. 2004. Fundamental limits on longitudinal bone growth: growth plate senescence and epiphyseal fusion. *Trends in Endocrinology and Metabolism* 15(8):370–374 DOI 10.1016/S1043-2760(04)00185-7.
Ogden JA, Conlogue GJ, Rhodin AGJ. 1981. Roentgenographic indicators of skeletal maturity in marine mammals (Cetacea). *Skeletal Radiology* 7(2):119–123 DOI 10.1007/BF00347376.

Padian K, Horner JR, de Ricqlès A. 2004. Growth in small dinosaurs and pterosaurs: the evolution of archosaurian growth strategies. *Journal of Vertebrate Paleontology* 24(3):555–571 DOI 10.1671/0272-4634(2004)024[0555:GISDAP]2.0.CO;2.

Padian K, Lamm E-T, Werning S. 2013. Selection of specimens. In: Padian K, Lamm E-T, eds. *Bone Histology of Fossil Tetrapods: Advancing Methods, Analysis, and Interpretation*. Berkeley: University of California Press, 35–54.

Piewkliang B, Pring A, Brugger J. 2008. The formation of precious opal: clues from the opalization of bone. *Canadian Mineralogist* 46(1):139–149 DOI 10.3749/canmin.46.1.139.

Piewkliang B, Pring A, Brugger J. 2008. The formation of precious opal: clues from the opalization of bone. *Canadian Mineralogist* 46(1):139–149 DOI 10.3749/canmin.46.1.139.

Prondvai E, Godefroit P, Adriaens D, Hu D-Y. 2018. Intrasedial histovariability, allometric growth patterns, and their functional implications in bird-like dinosaurs. *Scientific Reports* 8(1):258 DOI 10.1038/s41598-017-18218-9.

Prondvai E, Stein KHW, de Ricqlès A, Cubo J. 2014. Development-based revision of bone tissue classification: the importance of semantics for science. *Biological Journal of the Linnean Society* 112(4):799–816 DOI 10.1111/bij.12323.

Prondvai E, Stein K, Ösi A, Sander MP. 2012. Life history of *Rhamphorhynchus* inferred from bone histology and the diversity of pterosaurian growth strategies. *PLOS ONE* 7(2):e31392 DOI 10.1371/journal.pone.0031392.

Raath MA. 1977. The anatomy of the Triassic theropod *Syntarsus rhodesiensis* (Sauruschia: Podokesauridae) and a consideration of its biology. PhD dissertation. Rhodes University.

Raath MA. 1990. Morphological variation in small theropods and its meaning in systematics: evidence from *Syntarsus rhodesiensis*. In: Carpenter K, Currie PJ, eds. *Dinosaur Systematics: Perspectives and Approaches*. Cambridge: Cambridge University Press, 91–105.

Reid REH. 1996. Bone histology of the Cleveland–Lloyd dinosaurs and of dinosaurs in general. I. Introduction to bone tissues. *Brigham Young University Geology Studies* 41:25–71.

Rowe T. 1986. Homology and evolution of the deep dorsal thigh musculature in birds and other reptilia. *Journal of Morphology* 189(3):327–346 DOI 10.1002/jmor.1051890310.

Saitta ET. 2015. Evidence for sexual dimorphism in the plated dinosaur *Stegosaurus mjosi* (Ornithischia, Stegosauria) from the Morrison Formation (Upper Jurassic) of western USA. *PLOS ONE* 10(4):e0123503 DOI 10.1371/journal.pone.0123503.

Sampson SD, Ryan MJ, Tanke DH. 1997. Craniofacial ontogeny in centrosaurine dinosaurs (Ornithischia: Ceratopsidae): taxonomic and behavioral implications. *Zoological Journal of the Linnean Society* 121(3):293–337 DOI 10.1111/j.1096-3642.1997.tb00340.x.

Sander MP, Klein N, Buffetaut E, Cuny G, Suteethorn V, Locuff JL. 2004. Adaptive radiation in sauropod dinosaurs: bone histology indicates rapid evolution of giant body size through acceleration. *Organisms Diversity & Evolution* 4(3):165–173 DOI 10.1016/j.ode.2003.12.002.

Scannella JB, Horner JR. 2010. *Torosaurus* Marsh, 1891, is Triceratops Marsh, 1889 (Ceratopsidae: Chasmosaurinae): synonymy through ontogeny. *Journal of Vertebrate Paleontology* 30(4):1157–1168 DOI 10.1080/02724634.2010.483632.

Schachner ER, Manning PL, Dodson P. 2011. Pelvic and hindlimb myology of the basal archosaur *Poposaurus gracilis* (Archosauria: Poposauroidea). *Journal of Morphology* 272(12):1464–1491 DOI 10.1002/jmor.10997.
Schweitzer MH, Wittmeyer JL, Horner JR. 2005. Gender-specific reproductive tissue in ratites and *Tyrannosaurus rex*. *Science* **308**(5727):1456–1460 DOI 10.1126/science.1112158.

Schweitzer MH, Zheng W, Zanno L, Werning S, Sugiyama T. 2016. Chemistry supports the identification of gender-specific reproductive tissue in *Tyrannosaurus rex*. *Scientific Reports* **6**(1):1–10 DOI 10.1038/srep23099.

Sereno PC, Arcucci AB. 1994. Dinosaurian precursors from the Middle Triassic of Argentina: *Lagerpeton chanarensis*. *Journal of Vertebrate Paleontology* **13**(4):385–399 DOI 10.1080/02724634.1994.10011522.

Shelburne ECH, Thompson AC. 2016. Specimen whitening: an assessment of methods of ammonium chloride smoke removal. *Collection Forum* **30**(1–2):63–72 DOI 10.14351/0831-4985-30.1.63.

Shelton CD, Sander PM, Stein K, Winklerhorst H. 2012. Long bone histology indicates sympatric species of *Dimetrodon* (Lower Permian, Sphenacodontidae). *Earth and Environmental Science Transactions of the Royal Society of Edinburgh* **103**(3–4):217–236 DOI 10.1017/S175569101300025X.

Smith BH, Crummett TL, Brandt KL. 1994. Ages of eruption of primate teeth: a compendium for aging individuals and comparing life histories. *American Journal of Physical Anthropology* **37**(S19):177–231 DOI 10.1002/ajpa.1330370608.

Spinage CA. 1973. A review of the age determination of mammals by means of teeth, with especial reference to Africa. *African Journal of Ecology* **11**(2):165–187 DOI 10.1111/j.1365-2028.1973.tb00081.x.

Stein K, Prondvai E. 2014. Rethinking the nature of fibrolamellar bone: an integrative biological revision of sauropod plexiform bone formation. *Biological Reviews* **89**(1):24–47 DOI 10.1111/brv.12041.

Storá J. 2002. Neolithic seal exploitation on the Åland Islands in the Baltic Sea on the basis of epiphyseal fusion data and metric studies. *International Journal of Osteoarchaeology* **12**(1):49–64 DOI 10.1002/oa.612.

Tumarkin-Deratzian AR, Vann DR, Dodson P. 2006. Bone surface texture as an ontogenetic indicator in long bones of the Canada goose *Branta canadensis* (Anseriformes: Anatidae). *Zoological Journal of the Linnean Society* **148**(2):133–168 DOI 10.1111/j.1096-3642.2006.00232.x.

Tumarkin-Deratzian AR, Vann DR, Dodson P. 2007. Growth and textural ageing in long bones of the American alligator *Alligator mississippiensis* (Crocodylia: Alligatoridae). *Zoological Journal of the Linnean Society* **150**(1):1–39 DOI 10.1111/j.1096-3642.2007.00283.x.

Turner AH, Nesbitt SJ. 2013. Body size evolution during the Triassic archosauriform radiation. In: Nesbitt SJ, Desojo JB, Irrnis RB, eds. *Anatomy, Phylogeny, and Palaeobiology of Early Archosaurs and Their Kin*. Vol. 379. London: Geological Society Special Publications, 573–597.

Vucetich MG, Deschamps CM, Olivares AI, Dozo MT. 2005. Cavybaras, size, shape, and time. *Acta Palaeontologica Polonica* **50**:259–272.

Werning S. 2013. Evolution of Bone Histological Characters in Amniotes, and the Implications for the Evolution of Growth and Metabolism. PhD dissertation. Berkeley: University of California.

Woodruff DC, Fowler DW, Horner JR. 2017. A new multi-faceted framework for deciphering diplodocid ontogeny. *Palaeontologia Electronica* **20**(3.43A):1–53.

Woodward HN, Horner JR, Farlow JO. 2011. Osteohistological evidence for determinate growth in the American Alligator. *Journal of Herpetology* **45**(3):339–342 DOI 10.1670/10-274.1.

Zumwalt A. 2006. The effect of endurance exercise on the morphology of muscle attachment sites. *Journal of Experimental Biology* **209**(3):444–454 DOI 10.1242/jeb.02028.

Griffin et al. (2019), *PeerJ*, DOI 10.7717/peerj.6331