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Palaeontology

Ontogeny and the fossil record: what, if anything, is an adult dinosaur?

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Identification of the ontogenetic status of an extinct organism is complex, and yet this underpins major areas of research, from taxonomy and systematics to ecology and evolution. In the case of the non-avian dinosaurs, at least some were reproductively mature before they were skeletally mature, and a lack of consensus on how to define an 'adult' animal causes problems for even basic scientific investigations. Here we review the current methods available to determine the age of non-avian dinosaurs, discuss the definitions of different ontogenetic stages, and summarize the implications of these disparate definitions for dinosaur palaeontology. Most critically, a growing body of evidence suggests that many dinosaurs that would be considered 'adults' in a modern-day field study are considered 'juveniles' or 'subadults' in palaeontological contexts.

1. Introduction

The non-avian members of the Dinosauria (hereafter simply 'dinosaurs') were a diverse group of terrestrial archosaurian tetrapods that dominated global terrestrial environments during most of the Mesozoic era. In recent decades, major research advances have reframed our understanding of these animals, including their evolution, ecology, development, functional morphology and behaviour. Nonetheless, fundamentals about their biology remain problematic: most notably, the question of 'What is an adult dinosaur?'. Within extant sauropsids, an adult or mature individual is usually implicitly or explicitly defined as one that has reached sexual maturity (i.e. it is capable of reproduction). This is sometimes assessed directly, but frequently is inferred from proxies such as body size, coloration or skeletal characteristics (e.g. [1,2]). Because sexual maturity can only be indirectly inferred for a handful of specimens in most extinct dinosaurs, numerous other morphological criteria have been used (table 1 and figure 1). Yet, due to discordances in the timing of life events, an adult under one definition may be juvenile under another. Additionally, it is rarely practical or even possible to evaluate a fossil individual under all potential criteria for adulthood.

Reconciling these contradictions is critical to advancing understanding of dinosaur palaeobiology. Many studies presume to sample individuals that are adult or close to adult status, representing the 'adult' (typically an idealized 'final' ontogenetic stage) of a taxon. As commonly implied by dinosaur palaeontologists (although rarely stated outright), fully adult animals are those that display the 'ultimate' derived morphology for a taxon, with the complete development of autapomorphies and unique character combinations that define a taxon (e.g. [10]). A violation of this assumption has potentially enormous implications—juveniles and adults of the same taxon may be misidentified as adults of different species, affecting taxonomic and phylogenetic hypotheses (e.g. [15,16]). The work built on these assumptions, such as assessments of evolutionary rates of anatomical traits, in turn becomes questionable. Similarly, our ability to use data

Table 1. This is not an exhaustive list of terms used or definitions given. Age classes are given as used in the original sources and the definitions or reasoning for the assignment to this age class are direct quotes from the text. Additional details are often provided in the respective sources for assigning age classes, but these quotes are intended to be representative and not overarching. EFS, external fundamental system.

age class	definition	source
embryo	these occur both <i>in situ</i> and inside fragments of eggs exposed on erosional surfaces	[3]
perinate	we use the term 'perinate' ('around birth')	[4]
small nestling	the bone tissue that forms the shafts of the longer limb bones is . . . composed of vascular canals surrounded by an undifferentiated mineralized bone matrix	[5]
large nestling	in cross section, the shafts of the long bones generally have a cortex that is well differentiated from the marrow cavity	[5]
young	^a numerous differences in cranial and postcranial morphology given between 'young' and 'adult' <i>Protoceratops</i>	[6]
juvenile	a bone that is less than one-half the size of that of a typical adult specimen	[7]
juvenile	all vertebrae in embryonic and neonate ornithischian material, unsurprisingly, preserve open neurocentral sutures	[8]
juvenile	histological section of the tibia shows well-vascularized, woven and parallel-fibred primary cortical bone typical of juvenile ornithopods	[4]
subadult	the frontoparietal fontanelle is open in late-stage subadults	[9]
subadult	individuals of adult or virtually adult size, with additional characters indicating pre-adult status . . . but individuals lack several adult characters	[10]
subadult	a bone between one-half and two-thirds the size of that of a typical adult specimen	[7]
subadult	the individuals in this stage have both 'young' and 'adult' characters	[11]
subadult or young adult	^a neurocentral sutures have closed, partial fusion of scapula and coracoid and of the ilium and ischium, fusion of some cranial elements	[12]
adult	fully grown individuals with full expression of adult characters, often including fusion of skull elements	[10]
adult	a bone that is approximately the size of that of a typical adult specimen	[7]
adult	this histology is typical of an EFS and indicates that the individual was fully grown	[13]
old adult	nearly all of the cranial sutures are obliterated by co-ossification	[14]

^aThe quotations have been compressed for brevity.

from extant taxa and ecosystems to reconstruct the biology of ancient animals relies upon identification of age classes that are meaningfully equivalent.

Here we assess diverse concepts of ontogenetic status in dinosaurs and the associated problems with determining the life stage of a given specimen. We provide suggested definitions of different classes of ontogeny that attempt to align multiple current concepts and permit easier comparisons between disparate ideas about dinosaur growth. Importantly, and as part of a growing consensus in the field, we posit that a clear statement of criteria used for determining ontogenetic stage (already done in many studies) is necessary at all times. This not only enables unambiguous communication, but also allows discussion of the implications of the range of ages in dinosaurs and how this affects current ideas about their biology.

2. Institutional abbreviations

AMNH, American Museum of Natural History, New York, NY, USA; BYU, Brigham Young University, Provo, UT, USA; CM, Carnegie Museum of Natural History, Pittsburgh, PA, USA; FMNH, Field Museum of Natural History, Chicago, IL, USA; GMNH, Gunma Museum of Natural History, Gunma, Japan; MBR, Museum für Naturkunde, Humboldt Universität, Berlin, Germany; NMST, Division of Vertebrate Paleontology,

National Science Museum, Tokyo, Japan; YPM, Yale Peabody Museum of Natural History, NH, USA; ZPAL, Institute of Palaeobiology, Polish Academy of Sciences, Warsaw, Poland.

3. Methods of assessing ontogenetic status

The first key distinction in dinosaur ontogeny is that between adults and non-adults (figure 2). Fully adult animals are ideally the basic unit of alpha taxonomy (and by extension, systematic work), and presumed adult morphology is also often the ideal basis of most functional and ecological analyses. For example, non-adult animals often have traits that match the presumed ancestral condition [21], and thus the inclusion of non-adult animals in an analysis may lead to the recovery of an incorrect position for such a taxon [22]. Therefore, perhaps the most fundamental question with respect to ontogeny is: At what point does an animal become an adult?

Note that hereafter we use the term 'mature' or 'maturity' simply to mean that the animal has reached adult status under a particular criterion, and 'immature' that it has not reached this threshold. We do not mean to imply that maturity based on histology is, for example, the same as that based on skeletal fusion. In particular, 'maturity' does not necessarily imply that the animal is capable of reproducing,



Figure 1. A tableau of *Tyrannosaurus rex* skeletal reconstructions, on display at the Natural History Museum of Los Angeles County. The largest individual represents typical adult size for the taxon—current mainstream scientific consensus considers them all different ontogenetic stages of *T. rex* but the smaller specimens were originally referred to different genera. Photo: DWEH. (Online version in colour.)

although this is a standard use of the term in extant animals (further complicating comparisons).

(a) Body size

Although the youngest individuals are undoubtedly smaller than the oldest individuals in a population, absolute body size generally is a poor indicator of adult status in most taxa (e.g. [23]) although it has been used (e.g. [7]) for some dinosaurs and is often a (sometimes unreliable) proxy for maturity in studies of extant reptiles (e.g. [2,24]). Indeed, at least some specimens are extremely large by any standard yet do not appear to have stopped growing or reached osteological maturity (see also below). Even within extant animals with determinate growth, maximum sized individuals may be considerably larger than a more typical animal (e.g. the savannah elephant, *Loxodonta africana*, has a male height at adulthood between 3.2 and 4.0 m [25]). Thus, large size in one individual is not necessarily an indication of immaturity in another smaller one. Even if an adult has been diagnosed by multiple different criteria, a similarly sized animal from the same species may not be mature. Perhaps the best-case scenarios are represented by large samples of the hadrosaur dinosaurs *Maiasaura* and *Shantungosaurus*, which apparently represent standing populations for a wide size range of individuals [26,27]. The size distribution suggests that the largest individuals are indeed adults, corroborated by histology for *Maiasaura* [26].

(b) Osteological fusion

The fusion of major skeletal elements is often cited as a key indicator of adult status in dinosaurs ('skeletal maturity'; [28]). For instance, fusion of the sacral vertebrae to each

other and to the ilia is seen across many lineages as ontogeny progresses. Other elements show similar co-ossification, including the fusion of cranial ossifications to the underlying skull bones in some dinosaurs [10,29].

A widely cited criterion of maturity in dinosaurs concerns fusion between the neural arches of the vertebrae with their respective centra. This process often begins posteriorly, such that the posterior-most vertebrae will become fused before those that are more anterior. 'Adult' animals are presumed to have fully obliterated synchondroses in all vertebrae. Although this pattern of vertebral fusion is seen in extant crocodylians [8,30] and at least appears to generally follow in dinosaurs, the situation is complex (table 2). As with the example above, animals may fuse their vertebrae and be considered adult even while considerably smaller than other known individuals of the genus. In extant lizards [31], homologous elements show extensive interspecific variation in fusion relative to sexual maturity, and dinosaurs were likely similarly variable.

(c) Osteohistology

At a microscopic level, bone tissue undergoes considerable modelling and remodelling through the course of development. This phenomenon is increasingly well documented in modern species, permitting applications in extinct dinosaurs [22,32–34].

The smallest, and presumed youngest, individuals have limb bones characterized by unmodified primary bone. As an individual grows, this primary bone is replaced and remodelled as secondary bone, with clear differences visible in thin section. Appositional growth occurs around the circumference of a given element; pauses in this growth (often over an annual cycle) produce visible lines of arrested

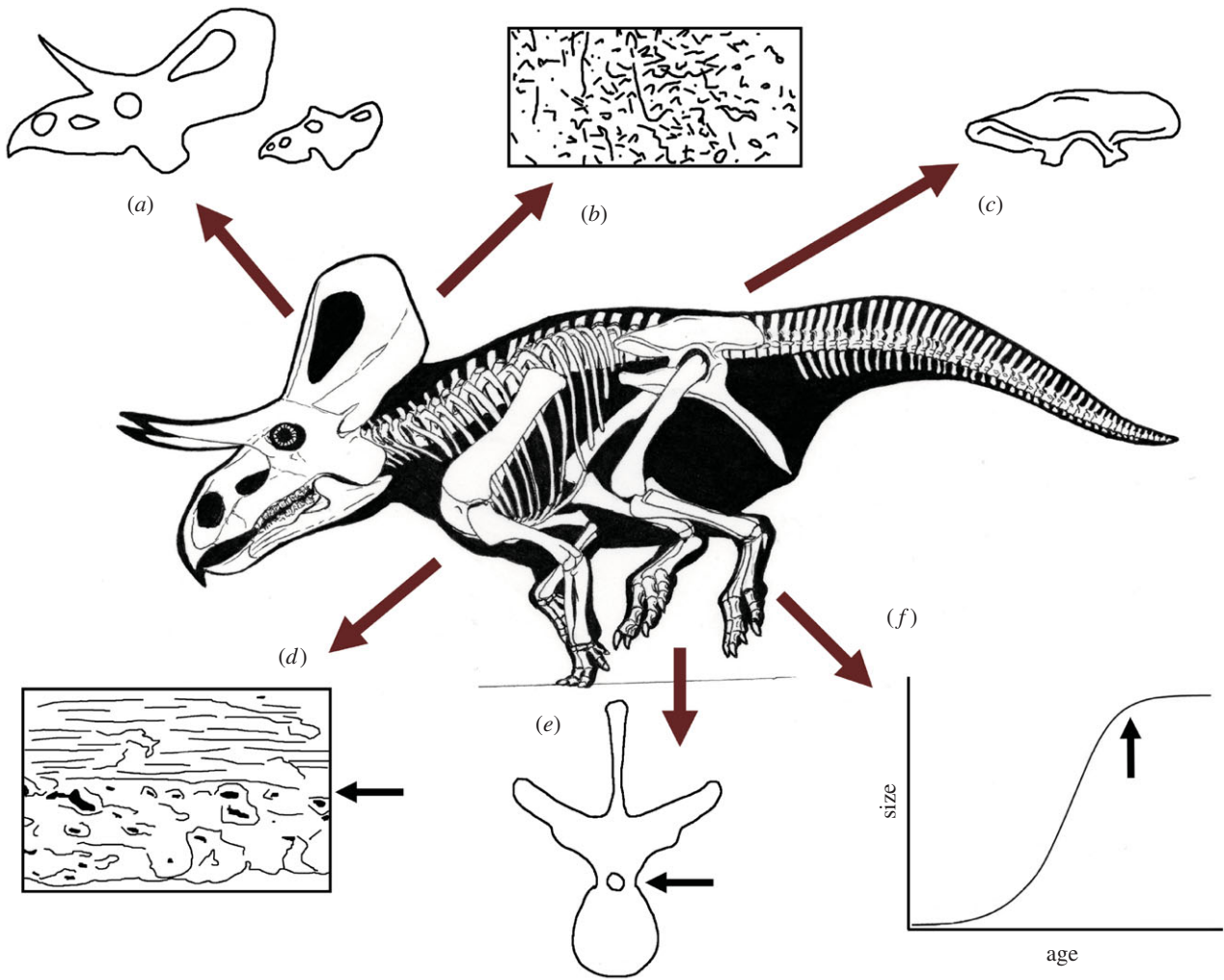


Figure 2. Various methods that may be used to determine the age/ontogenetic status of a given dinosaur specimen. Central image is a reconstruction of the skeleton of an adult ceratopsian *Zuniceratops*, with surrounding indications of maturity (taken from multiple sources and do not necessarily relate to this taxon). (a) Development of sociosexual signals (adult left, juvenile right—modified from [9]), (b) surface bone texture (traced from [17]), (c) large size, represented here by an ilium of the same taxon that is considerably larger than that of a known adult specimen, (d) reproductive maturity, here based on the presence of medullary bone here shown below the black arrow (traced from [18]), (e) fusion of the neurocentral arch—location of the obliterated synchondrosis indicated by black arrow (traced from [19]), (f) asymptote of growth based on multiple species indicated by black arrow (based on [20]). Central image by Julius Csotonyi, used with permission. (Online version in colour.)

growth (LAGs—also ‘annuli’ or ‘resting lines’). As growth slows over the course of a lifetime, the spacing between LAGs becomes closer. When LAGs are closely spaced within avascular bone of the peripheral cortex, these are termed an external fundamental system (EFS; [34,35]). An EFS is presumed to indicate cessation of overall growth and an unambiguous adult (even if it would already have been adult by other criteria), or even a senescent adult [26].

In some dinosaurs, mechanically important parts of the skeleton are cartilaginous, only ossifying late in ontogeny or not at all [36]. A prime example is the olecranon process of the stegosaur *Kentrosaurus*, which is only ossified in the largest individuals [37]; smaller individuals lack an olecranon entirely.

(d) Bone surface texture

Gross texture of the bone surface changes through ontogeny, mirroring microscopic remodelling [38]. Perhaps the best example within dinosaurs concerns the skulls of horned dinosaurs, which change surface texture from lightly striated to deeply rugose during ontogeny [10,17].

(e) Growth curves

Because LAGs and similar features are often assumed to be annual in deposition, they have been used to create growth curves for an individual [32]. The shapes of the growth curves (which illustrate changes in growth rate), through comparison with extant taxa, are in turn used to mark ontogenetic milestones such as reproductive maturity (associated with the initial slow-down of growth) and the effective cessation of growth during full adulthood [20]. These sorts of analyses and observations have been important for recognizing individuals that were reproductively mature but not skeletally mature (e.g. [39,40]).

(f) Reproductive maturity

Reproductive (i.e. sexual) maturity is the ability to produce offspring and could be indicated by the presence of eggs inside the body cavity of an animal [41] or the possession of medullary bone [18], either of which indicates a female that is able to lay eggs. Additional support could come from an animal preserved brooding on a nest or in the

Table 2. The timing of macroscopic changes in sauropod skeletons over ontogeny is not consistent among taxa. 'yes' indicates fusion to the adjacent respective spines or centra, 'no' indicates lack of fusion, and a blank indicates that the relevant material is not preserved. Modified from Wedel & Taylor [23, table 1].

taxon	specimen	sacral 1 spine	sacral 4 spine	sacral 1 centrum	sacral 5 centrum	sacral 1 rib	sacral 5 rib	all cervical ribs fused	scapula and coracoid fused
<i>Apatosaurus ajax</i>	YPM 1860			no	no	no	no	yes	no
<i>Apatosaurus ajax</i>	NMST-PV 20375	no	yes	yes	yes	yes	yes	yes	yes
<i>Brontosaurus excelsus</i>	YPM 1981			yes	no	yes	no		
<i>Brontosaurus excelsus</i>	YPM 1980	no	yes	yes	yes	yes	yes	yes	no
<i>Diplodocus carnegii</i>	CM 84/94	no	no	yes	yes	no	yes	yes	yes
<i>Barosaurus lentus</i>	AMNH 6341	no		no	yes			no	yes
<i>Haplocanthosaurus delphi</i>	CM 879	yes	yes	no	no	no	no	no	no
<i>Haplocanthosaurus delphi</i>	CM 572	yes	no	yes	yes	yes	yes	yes	
<i>Camarasaurus grandis</i>	GMMH-PV 101	no	no					no	no
<i>Camarasaurus lewisi</i>	BYU 9047	yes	no	yes	yes	yes	yes	yes	
<i>Camarasaurus supremus</i>	AMNH 5761	no	yes	no	yes	no	yes	no	no
<i>Brachiosaurus altithorax</i>	FMNH P 25107	no	no	yes	yes	yes	yes	no	no
<i>Opisthocoelocaudia skarzynskii</i>	ZPAL MgD-I/48			yes	yes	yes	yes	yes	yes

company of small conspecifics. The latter indicators are based on the assumption that the larger animal in question is a parent or of similar age, and other interpretations are possible. For example, species where juveniles assist adults in rearing the young (e.g. 'helper at the nest') could lead to immature animals that are not parents becoming fossilized alongside nests or younger animals. Mixed age class aggregations are also known for many dinosaurs, and there seems to be bias in some aggregations that favours sampling of smaller individuals [42]. It is therefore possible that in some contexts, apparent aggregations of juveniles with 'adults' may in fact be two sizes of juveniles. Importantly, some studies have identified individuals that are reproductively mature but not skeletally or histologically mature, nor at full 'adult' size [39,40].

(g) Development of sociosexual dominance characteristics

In association with reproductive maturity comes the full development of additional sociosexual characteristics that are linked to reproduction. Animals that are not yet capable of reproducing are unlikely to need these often large and costly ornaments and weapons. Thus, the allometric growth of such features probably indicates that they are used, at least as one function, in sexual or social dominance contests and that the animals are capable of reproducing [43]. For growth series within a single taxon, adults are identified as the individuals with full development of ornamentation. Noted examples in dinosaurs include the facial horns and frills of ceratopsids (e.g. [6,10]) or the cranial crests of hadrosaurids [44]. Because structures involved with sociosexual dominance tend to be exaggerated, they will typically form part of the package of autapomorphies and synapomorphies by which specimens are both identified and sorted by phylogenetic analysis. This measure of ontogeny has the potential to confound our understanding more than any other, since juveniles and subadults that lack 'extreme' display structures can be difficult to even recognize as members of the same taxon.

4. Reconciliation of definitions

Clearly, there are both contradictions and overlap for various definitions of maturity (table 1), and these create problems for researchers. A specimen that is mature on the basis of cranial ornamentation may be immature on the basis of osteohistology or skeletal fusion, for instance. This is further complicated if the basis of assigning maturity within a particular research paper is not noted, and complicated still further by a lack of comparability with sometimes conflicting definitions of maturity (usually reproductive) in extant taxa.

In addition, many fossils simply cannot be compared with one another. Histological samples cannot be taken from every specimen owing to preservation, fragility or equipment availability, and such detailed sampling is not practical for large and wide-ranging studies that may cover hundreds of specimens. Thus, definitions must be flexible enough to cover the variation seen in not just the growth of dinosaurs, but also the available information. Here we create a set of definitions for the fundamental life stages of dinosaurs that will be broadly applicable in most situations. We consider these only a starting point and encourage different definitions to be used as

appropriate to the situation. However, we caution that terms even as simple as 'adult' or 'juvenile' be accompanied by a definition (and/or appropriate citation) or description of the characteristics used to define such life stages.

Within a given genus or species, the relative ages of specimens may be somewhat simple to determine, with different animals exhibiting different sizes, or differing levels of acquisition of adult characteristics, and histological thin sections may simply show different LAG counts. However, such comparisons are of little value between species (including close relatives), and may even vary greatly within a species hence the suggestions here for reconciliation of definitions.

Additional subdivisions can be identified in some cases (e.g. [5]), and should not be automatically subsumed. However, because the intention here is to provide simple definitions that can work across multiple taxa with different biologies and differing types of evidence, we restrict ourselves to broad definitions of largely unambiguous life stages that are common across all dinosaurs.

(a) Adult

The identity of this age class is critical because definitions of other classes often rely on it (e.g. [7]). Adult animals may be diagnosed through any of the above-described criteria (size, asymptote of growth, osteological fusion, etc.), but may also be confounded by conflicting signals (e.g. sexually mature animals that have not yet acquired all morphological features that characterize a taxon). Ideally, therefore, multiple overlapping criteria should be used, and researchers should explicitly state which regime they employ (histological, fusion, etc.). A definition of an adult dinosaur is therefore: *An animal that has reached a point in life commensurate with the cessation of rapid growth as indicated by osteological and histological features, in addition to reproductive maturity.* Animals that fall primarily under this definition may be considered adult.

(b) Subadult

Those individuals that are transitioning between juvenile and adult status are subadult, and thus any definition should encompass this shift. Therefore, we define subadults as: *An animal that combines features of juveniles and adults, lacking definitive adult characteristics (e.g. an EFS or final form of ornamentation) but possessing features that do not correspond to the juvenile condition (e.g. numerous fused elements, large body size).* Because sexual maturity can occur well before adult status under some criteria (e.g. an EFS), this is one area where a given individual might be considered both reproductively mature and yet still osteologically subadult.

(c) Juvenile

These may be considered as: *Any animal that does not show any signs of impending maturity that would place it as an adult or subadult animal (i.e. little or no skeletal fusion, poorly developed ornamentation, few or no LAGS, no medullary bone, etc.).* Note that some characters do appear very early in the ontogeny of some taxa (e.g. the incipient frill present in even very young ceratopsians [45]). We subsume the oft-used categories of 'hatchling', 'neonate' and 'nestling' into 'juvenile'. Although they are useful descriptors from a behavioural and taphonomic perspective, they represent a very limited stretch of life for most animals (and for potentially only a matter of minutes).

(d) Embryo

An embryo is here considered: *Any specimen preserved within the confines of an egg or likely to have been so, representing an individual prior to hatching.* An egg is not required as part of this definition because examples of embryos apparently preserved without an egg [46, p. 211] are known. Note that we also consider Horner *et al.*'s [4] 'perinate' a useful alternative, because it is not always possible to distinguish between an embryo and a newly hatched animal.

5. Discussion and implications

Many of the ways by which 'adult' dinosaurs have previously been recognized imply that numerous individual dinosaurs had not actually reached maturity when they died. Even very large animals may exhibit a lack of fusion across multiple elements or lack an EFS, indicating potential for considerable growth. This is true even for some specimens exhibiting fully developed sociosexual characteristics, occurrence within normal population distributions or the presence of medullary bone implying that they are reproductively mature.

As a result, studies of dinosaurs may make assumptions about the ontogenetic status of a given specimen without regard to the variations known. Although ontogenetic trajectories have been studied in detail for a handful of taxa, allowing solid interpretations of the likely intersection of features such as size, asymptote of growth and fusion of various sutures (thus allowing maturity to be judged in other specimens from limited data [15,18,20] for *Tyrannosaurus*), most are not. Reasonable assumptions can be made in many cases about the likely age of various specimens, but nevertheless we urge researchers to be more explicit in stating under which criteria they are defining specimens as various ontogenetic stages, particularly adults. A lack of explicit information about such identifications does not inherently mark an assignment incorrect, but does potentially limit confidence in the referral and the repeatability of any analysis or use of the data. Ontogenetic sequence analysis holds some promise in this regard, particularly in formalizing definitions of ontogenetic stages and documentation of individual variation [47].

Correct identification of the ontogenetic status of a specimen (or at least a clear statement on the basis for the assignment) is critical to ensuring that specimens and/or taxa are comparable in large analyses where body size is relevant to the data. For example, studies on browsing height, giantism and biomass of populations may all be profoundly influenced if specimens are identified as adult when they are not.

Similarly, given the ontogenetic changes that can occur to major characters, it is critical to both taxonomic and cladistic studies that the life stage of a given specimen is correctly

identified. This is not to suggest that non-adult specimens should be excluded from such assessments and analyses, or that single small changes or incongruencies in, for example, patterns of osteological fusion, should be used to assign a specimen to a particular life stage or rule out another. Many important taxa are known from only definitively non-adult specimens [48]. Although caution is warranted in their identification and use in studies, they are often identifiable as distinct taxa and should not be *a priori* ignored.

The questions of 'When in ontogeny can you recognize a species as distinct from closely related species?' and 'When in ontogeny can you correctly place a species in its evolutionary position?' are separate, but related (and often conflated) points. This is exemplified by the case of hadrosaurid dinosaurs, in which genus or species-diagnostic features are observable within juveniles of many taxa, despite major morphological changes through ontogeny (e.g. [49]). Nonetheless, the preponderance of 'primitive' features in juveniles still renders them difficult to place 'correctly' in a phylogeny (e.g. [50]).

Correct identification of life stage also is relevant to fundamentals of evolution—if the onset of sexual reproduction substantially preceded cessation of growth in dinosaurs then the 'adult' phenotype may not have been the primary target of selection. In fact, once juveniles or subadults are capable of reproducing, it is conceivable a population could exist with potentially no individuals making it through the survivorship gauntlet into 'adulthood' and close to maximum body size. The occasional hints from the fossil record of anomalously large sauropods like *Bruhathkayosaurus* [51], and the Broome trackmaker [52] might be explained if many sauropods were primarily 'subadult' reproducers, and thus extremely large adults were actually vanishingly rare. This is a rather extreme hypothesis, but not an impossible one, and it raises the issue that some well-known species may not actually be represented by fully adult individuals under any of the criteria suggested above. Similarly, the apparent lack of sexual dimorphism common in taxa with large ornaments could relate to mutual sexual selection [53] but might also be because few individuals reached a 'final' stage where dimorphism was clear between the two ornamented sexes.

Indeed, the whole concept of an 'adult' may not be directly comparable in any meaningful sense between extant tetrapods and extinct dinosaurs. This issue is ripe for study, both in extant and in extinct taxa.

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References

- Ralph CJ, Geupel G, Pyle P, Martin T, DeSante D. 1993 *Handbook of field methods for monitoring landbirds. General Technical Report PSW-GTR-144-www.* Albany, CA: Pacific Southwest Research Station, Forest Service U.S. Department of Agriculture.
- Dunham AE, Morin PJ, Wilbur HM. 1988 Methods for the study of reptile populations. In *Biology of the Reptilia*, vol. 16. *Ecology B: defense and life history*, pp. 331–386. New York, NY: Alan R. Liss, Inc.
- Chiappe L, Dingus L, Jackson F, Grellet-Tinner G, Aspínall R, Clarke J, Coria R, Garrido AC, Loope D. 2000 Sauropod eggs and embryos from the Late Cretaceous of Patagonia. In *First Int. Symp. on Dinosaur Eggs and Babies, Isona I Conca Dellà, Catalonia, Spain. Extended Abstracts* (eds AM Bravo, T Reyes), pp. 23–29. Lleida: Impremta Provincial de la Diputació de Lleida.
- 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)
5. 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. *J. Vertebr. Paleontol.* **20**, 115–129. (doi:10.1671/0272-4634(2000)020[0115:LBHOTH]2.0.CO;2)
 6. Brown B, Schlaikjer EM. 1940 The structure and relationships of *Protoceratops*. *Ann. NY Acad. Sci.* **40**, 133–266. (doi:10.1111/j.1749-6632.1940.tb57047.x)
 7. Ryan MJ, Russell AP, Eberth DA, Currie PJ. 2001 The taphonomy of a *Centrosaurus* (Ornithischia, Ceratopsidae) bone bed from the Dinosaur Park Formation (Upper Campanian), Alberta, Canada, with comments on cranial ontogeny. *Palaios* **16**, 482–506. (doi:10.1669/0883-1351(2001)016<0482:TTOACO>2.0.CO;2)
 8. Irmis RB. 2007 Axial skeleton ontogeny in the Parasuchia (Archosauria: Pseudosuchia) and its implications for ontogenetic determination in archosaurs. *J. Vertebr. Paleontol.* **27**, 350–361. (doi:10.1671/0272-4634(2007)27[350:ASOITP]2.0.CO;2)
 9. Scannella JB, Fowler DW, Goodwin MB, Horner JR. 2014 Evolutionary trends in *Triceratops* from the Hell Creek Formation, Montana. *Proc. Natl Acad. Sci. USA* **111**, 10 245–10 250. (doi:10.1073/pnas.1313334111)
 10. Sampson SD, Ryan MJ, Tanke DH. 1997 Craniofacial ontogeny in centrosaurine dinosaurs (Ornithischia: Ceratopsidae): taxonomic and behavioral implications. *Zool. J. Linn. Soc.* **121**, 293–337. (doi:10.1111/j.1096-3642.1997.tb00340.x)
 11. Handa N, Watabe M, Tsogtbaatar K. 2012 New specimens of *Protoceratops* (Dinosauria: Neoceratopsia) from the Upper Cretaceous in Udyn Sayr, southern Gobi area, Mongolia. *Paleontol. Res.* **16**, 179–198. (doi:10.2517/1342-8144-16.3.179)
 12. Sereno PC, Tan L, Brusatte SL, Kriegstein HJ, Zhao X, Cloward K. 2009 Tyrannosaurid skeletal design first evolved at small body size. *Science* **326**, 418–422. (doi:10.1126/science.1177428)
 13. Sander MP, Mateus O, Laven T, Knötschke N. 2006 Bone histology indicates insular dwarfism in a new Late Jurassic sauropod dinosaur. *Nature* **441**, 739–741. (doi:10.1038/nature04633)
 14. Farke AA. 2011 Anatomy and taxonomic status of the chasmosaurine ceratopsid *Nedoceratops hatcheri* from the Upper Cretaceous Lance Formation of Wyoming, U.S.A. *PLoS ONE* **6**, e16196. (doi:10.1371/journal.pone.0016196)
 15. Carr TD. 1999 Craniofacial ontogeny in Tyrannosauridae (Dinosauria, Coelurosauria). *J. Vertebr. Paleontol.* **19**, 497–520. (doi:10.1080/02724634.1999.10011161)
 16. Scannella JB, Horner JR. 2010 *Torosaurus* Marsh, 1891, is *Triceratops* Marsh, 1889 (Ceratopsidae: Chasmosaurinae): synonymy through ontogeny. *J. Vertebr. Paleontol.* **30**, 1157–1168. (doi:10.1080/02724634.2010.483632)
 17. Brown CM, Russell AP, Ryan MJ. 2009 Pattern and transition of surficial bone texture of the centrosaurine frill and their ontogenetic and taxonomic implications. *J. Vertebr. Paleontol.* **29**, 132–141. (doi:10.1671/039.029.0119)
 18. Schweitzer MH, Wittmeyer JL, Horner JR. 2005 Gender-specific reproductive tissue in ratites and *Tyrannosaurus rex*. *Science* **308**, 1456–1460. (doi:10.1126/science.1112158)
 19. Hatcher JB, Marsh OC, Lull RS. 1907 *The Ceratopsia*. *US Geol. Surv. Monogr.* **49**, 1–300. (doi:10.5962/bhl.title.60500)
 20. Erickson GM, Rogers KC, Yerby SA. 2001 Dinosaurian growth patterns and rapid avian growth rates. *Nature* **412**, 429–433. (doi:10.1038/35086558)
 21. Rozhdestvensky AK. 1965 Growth changes in Asian dinosaurs and some problems of their taxonomy. *Paleontol. Ž.* **3**, 95–109.
 22. Fowler DW, Woodward HN, Freedman EA, Larson PL, Horner JR. 2011 Reanalysis of ‘*Raptorex kriegsteini*’: a juvenile tyrannosaurid dinosaur from Mongolia. *PLoS ONE* **6**, e21376. (doi:10.1371/journal.pone.0021376)
 23. Wedel MJ, Taylor MP. 2013 Neural spine bifurcation in sauropod dinosaurs of the Morrison Formation: ontogenetic and phylogenetic implications. *Palarch’s J. Vertebr. Paleontol.* **10**, 1–34.
 24. Halliday TR, Verrell PA. 1988 Body size and age in amphibians and reptiles. *J. Herpetol.* **22**, 253–265. (doi:10.2307/1564148)
 25. Laursen L, Bekoff M. 1978 *Loxodonta africana*. *Mamm. Species* **92**, 1–8. (doi:10.2307/3503889)
 26. Woodward HN, Freedman Fowler EA, Farlow JO, Horner JR. 2015 *Maiasaura*, a model organism for extinct vertebrate population biology: a large sample statistical assessment of growth dynamics and survivorship. *Paleobiology* **41**, 503–527. (doi:10.1017/pab.2015.19)
 27. Hone DWE, Sullivan C, Zhao Q, Wang K, Xu X. 2014 Body size distribution in a colossal hadrosaurid death assemblage from the Upper Cretaceous of Zhucheng, Shandong Province, China. In *Hadrosaurids* (eds DA Eberth, DC Evans), pp. 524–531. Bloomington, IN: Indiana University Press.
 28. O’Connor PM. 2007 The postcranial axial skeleton of *Majungasaurus crenatissimus* (Theropoda: Abelisauridae) from the Late Cretaceous of Madagascar. *Soc. Vertebr. Paleontol. Mem.* **8**, 127–162. (doi:10.1671/0272-4634(2007)27[127:TPASOM]2.0.CO;2)
 29. Goodwin MB, Horner JR. 2004 Cranial histology of pachycephalosaurs (Ornithischia: Marginocephalia) reveals transitory structures inconsistent with head-butting behavior. *Paleobiology* **30**, 253–267. (doi:10.1666/0094-8373(2004)030<0253:CHOPOM>2.0.CO;2)
 30. Brochu CA. 1996 Closure of neurocentral sutures during crocodylian ontogeny: Implications for maturity assessment in fossil archosaurs. *J. Vertebr. Paleontol.* **16**, 49–62. (doi:10.1080/02724634.1996.10011283)
 31. Maisano JA. 2002 Terminal fusions of skeletal elements as indicators of maturity in squamates. *J. Vertebr. Paleontol.* **22**, 268–275. (doi:10.1671/0272-4634(2002)022[0268:TFOSEA]2.0.CO;2)
 32. Erickson GM. 2014 On dinosaur growth. *Annu. Rev. Earth Planet. Sci.* **42**, 675–697. (doi:10.1146/annurev-earth-060313-054858)
 33. Padian K, Lamm E-T (eds). 2013 *Bone histology of fossil tetrapods: issues, methods, and databases*. Berkeley, CA: University of California Press.
 34. Huttenlocker AK, Woodward HN, Hall BK. 2013 The biology of bone. In *Bone histology of fossil tetrapods: issues, methods, and databases* (eds K Padian, E-T Lamm), pp. 13–34. Berkeley, CA: University of California Press.
 35. Klein N, Sander M. 2008 Ontogenetic stages in the long bone histology of sauropod dinosaurs. *Paleobiology* **34**, 247–263. (doi:10.1666/0094-8373(2008)034[0247:OSITLB]2.0.CO;2)
 36. Jensen JA. 1988 A fourth new sauropod dinosaur from the Upper Jurassic of the Colorado Plateau and sauropod bipedalism. *Gt Basin Nat.* **48**, 121–145.
 37. Mallison H. 2010 CAD assessment of the posture and range of motion of *Kentrosaurus aethiopicus* Hennig 1915. *Swiss J. Geosci.* **103**, 211–233. (doi:10.1007/s00015-010-0024-2)
 38. Tumarkin-Deratzian AR. 2010 Histological evaluation of ontogenetic bone surface texture changes in the frill of *Centrosaurus apertus*. In *New Perspectives on Horned Dinosaurs, the Royal Tyrrell Museum Ceratopsian Symp.*, pp. 251–263. Bloomington, IN: Indiana University Press.
 39. 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. *Biol. Lett.* **3**, 558–561. (doi:10.1098/rsbl.2007.0254)
 40. Lee AH, Werning S. 2008 Sexual maturity in growing dinosaurs does not fit reptilian growth models. *Proc. Natl Acad. Sci. USA* **105**, 582–587. (doi:10.1073/pnas.0708903105)
 41. Sato T, Cheng Y-N, Wu X, Zelenitsky DK, Hsiao Y. 2005 A pair of shelled eggs inside a female dinosaur. *Science* **308**, 375. (doi:10.1126/science.1110578)
 42. Varricchio DJ. 2011 A distinct dinosaur life history? *Hist. Biol.* **23**, 91–107. (doi:10.1080/08912963.2010.500379)
 43. Knell RJ, Naish D, Tomkins JL, Hone DWE. 2013 Sexual selection in prehistoric animals: detection and implications. *Trends Ecol. Evol.* **28**, 38–47. (doi:10.1016/j.tree.2012.07.015)
 44. Dodson P. 1975 Taxonomic implications of relative growth in lambeosaurine hadrosaurids. *Syst. Zool.* **24**, 37–54. (doi:10.2307/2412696)
 45. Fastovsky DE, Weishampel DB, Watabe M, Barsbold R, Tsogtbaatar K, Narmandakh P. 2011 A nest of *Protoceratops andrewsi* (Dinosauria, Ornithischia). *J. Paleontol.* **85**, 1035–1041. (doi:10.1666/11-008.1)
 46. Carpenter K. 1999 *Eggs, nests, and baby dinosaurs: a look at dinosaur reproduction*. Bloomington, IN: Indiana University Press.
 47. Colbert MW, Rowe T. 2008 Ontogenetic sequence analysis: using parsimony to characterize developmental sequences and sequence

- polymorphism. *J. Exp. Zool.* **310B**, 398–416. (doi:10.1002/jez.b.21212)
48. Dal Sasso C, Maganuco S. 2011 *Scipionyx samniticus* (Theropoda: Compsognathidae) from the Lower Cretaceous of Italy. *Mem. Della Soc. Ital. Sci. Nat. E Mus. Civ. Storia Nat. Milano* **37**, 1–283.
49. Farke AA, Chok DJ, Herrero A, Scolieri B, Werning S. 2013 Ontogeny in the tube-crested dinosaur *Parasaurolophus* (Hadrosauridae) and heterochrony in hadrosaurids. *PeerJ* **1**, e182. (doi:10.7717/peerj.182)
50. Campione NE, Brink KS, Freedman EA, McGarrity CT, Evans DC. 2013 '*Glishades ericksoni*', an indeterminate juvenile hadrosaurid from the Two Medicine Formation of Montana: implications for hadrosauroid diversity in the latest Cretaceous (Campanian-Maastrichtian) of western North America. *Palaebiodiv. Palaeoenviron.* **93**, 65–75. (doi:10.1007/s12549-012-0097-1)
51. Yadagiri P, Ayyasami K. 1987 A carnosaurian dinosaur from the Kallamedu Formation (Maastrichtian horizon), Tamil Nadu. *Geol. Surv. India Spec. Publ.* **11**, 523–528.
52. Thulborn RA, Hamley T, Foulkes P. 1994 Preliminary report on sauropod dinosaur tracks in the Broome Sandstone (Lower Cretaceous) of Western Australia. *Gaia* **10**, 85–96.
53. Hone DWE, Naish D, Cuthill IC. 2012 Does mutual sexual selection explain the evolution of head crests in pterosaurs and dinosaurs? *Lethaia* **45**, 139–156. (doi:10.1111/j.1502-3931.2011.00300.x)