A transitional species of *Daspletosaurus* Russell, 1970 from the Judith River Formation of eastern Montana

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ABSTRACT

Here we describe a new derived tyrannosaurine, *Daspletosaurus wilsoni* sp. nov., from Judithian strata (~76.5 Ma) intermediate in age between either of the previously described species of this genus. *D. wilsoni* displays a unique combination of ancestral and derived characteristics, including a cornual process of the lacrimal reduced in height relative to *D. torosus* and more basal tyrannosaurines, and a prefrontal with a long axis oriented more rostrally than in *D. horneri* and more derived tyrannosaurines. The description of this taxon provides insight into evolutionary mode in Tyrannosaurinae, lending strength to previous hypotheses of anagenesis within *Daspletosaurus* and increasing the resolution with which the evolution of this lineage can be reconstructed. Cladistic phylogenetic methods, stratigraphy, and qualitative analysis of the morphology of relevant taxa supports an anagenetic model for the origin of morphological novelty in this genus, highlighting the predominance of anagenetic evolution among contemporary dinosaur lineages.

INTRODUCTION

Since their naming at the turn of the 20th century (*Osborn, 1905*), tyrannosaurids have captivated public and scientific imagination alike, and are as a result among the best-studied groups of Cretaceous theropods (*Brusatte et al., 2010*). Perhaps the most successful group of tyrannosaurids were the latest-Cretaceous tyrannosaurines, including among them a diverse array of forms from the slender-snouted alioramins (*Lü et al., 2014*) to robust and deep-jawed taxa like *Teratophoneus* (*Carr et al., 2011*) and the eponymous *Tyrannosaurus rex* (*Carr & Williamson, 2010*). However, much of the diversity of derived tyrannosaurines remains understudied or poorly understood (*Paulina Carabajal et al., 2021*), hampering understanding of paleobiogeographic and evolutionary trends (*Loewen et al., 2013; Carr et al., 2017; Brusatte & Carr, 2016*).

The tyrannosaurine *Daspletosaurus* has been known from Campanian fossil deposits of northern Laurasia for over half a century. However, published work on the phylogeny and paleobiology of this genus is relatively scarce beyond its initial description (*Russell, 1970; Carr et al., 2017; Paulina Carabajal et al., 2021*). Several enigmatic tyrannosaurine specimens initially referred to the type species or simply to *Daspletosaurus sp.* (including...
the recently named *D. horneri*) have been noted as representing novel species by previous workers for several decades (*Carr, 1999; Currie, 2003; Carr et al., 2017; Horner, Varricchio & Goodwin, 1992; Paulina Carabajal et al., 2021*), indicating a more speciose genus than has currently been described. Filling this gap is especially pertinent to understanding rates and patterns of speciation in the Campanian of Laurasia, both within tyrannosaurs and among dinosaurs as a whole, as both described species of *Daspletosaurus* have been hypothesized to represent an anagenetic lineage (*Carr et al., 2017*), including this genus among the many contemporary dinosaur lineages for which anagenesis has been suggested (*Horner, Varricchio & Goodwin, 1992; Fowler & Fowler, 2020*).

Here we describe *Daspletosaurus wilsoni* (sp. nov.). This addition to Campanian tyrannosaurid diversity has the potential to refine existing hypotheses regarding tyrannosaurid evolution in the Late Cretaceous, and lends strength to the hypothesis of anagenesis as a predominant mode of evolution in *Daspletosaurus* (*Carr et al., 2017*).

**Etymology**

*wilsoni*, Latinization of “Wilson,” after John Wilson, the discoverer of the holotype specimen.

**Holotype**

BDM 107, preserving a partial disarticulated skull and postcranium, including both premaxillae, a right maxilla, jugal, lacrimal, quadratojugal, dentary, and splenial, and a left postorbital and squamosal. Also preserved are partial cervical, sacral, and caudal series, a rib, a chevron, and a first metatarsal. Cranial bones are very finely preserved, with intricate and detailed surface textures especially on the maxilla and postorbital, with teeth preserved in the maxilla, dentary, and one premaxilla. The sacral and caudal centra are preserved in a heavy and hard concretion and are not yet prepared. The holotype specimen is stored in the collections of the Badlands Dinosaur Museum (BDM) in Dickinson, North Dakota.

**Geological Setting**

BDM 107 was recovered from the site “Jack’s B2,” discovered in 2017 by John Wilson in exposures of the Judith River Formation near Glasgow (Valley County, MT, USA). This is significantly further east than classic ‘Judith’ localities (*Fig. 1*), and is sedimentologically atypical, representing distal floodplain and delta sediments deposited during the maximum Campanian regression of the Western Interior Seaway. Here, the Judith River Formation is up to ~48 m thick, with the “Jack’s B2” site occurring ~30 m below the contact with the overlying Bearpaw Shale.

Precise stratigraphic placement of this easternmost Judith is currently unclear, although an age of ~76.5 Ma seems most likely, which would correlate in time with the lower to middle part of the Dinosaur Park Formation, Alberta (*Eberth, Currie & Koppelhus, 2005*; Warshaw and Fowler (2022), *PeerJ*, DOI 10.7717/peerj.14461)
A youngest age limit of 75.64 Ma (Ogg & Hinnov, 2012) is delineated by ammonites tentatively identified as *Didymoceras stevensoni* (J. Slattery, 2020, personal communication) collected by BDM from local outcrops of the overlying Bearpaw Shale (although these were not at the base of the Bearpaw, so older ammonite specimens may be encountered during future prospecting). At present, more precise stratigraphic position can be inferred from the timing of the maximum regression of the Western Interior Seaway during the Campanian (correlated with the R8 regression of Kauffman (1977) and Rogers et al. (2016)). In Alberta and Saskatchewan, the Foremost, Oldman, and Dinosaur Park formations represent early to late subcycles (respectively) of the R8 regression, and of these, the Foremost (~80.5–79.5 Ma) and lower Oldman (~79.5–79.0 Ma; and regional equivalents) are restricted to the west (Alberta and west central Montana), and did not extend as far east as Saskatchewan or our study area in eastern Montana (Eberth, Currie & Koppelhus, 2005). During late R8, the upper Oldman (~77.5–77.0 Ma) and Dinosaur Park (~76.9–76.0 Ma) Formations were deposited much further to the east, with the lowermost Dinosaur Park recording the R8 maximum regression at ~76.9–76.4 Ma (Eberth, Currie & Koppelhus, 2005; Fowler, 2017). This correlates well with the Judith River Formation of Montana, where Rogers et al. (2016) show the maximum regression of R8 occurring shortly
before 76.2 Ma, based on radiometric dates acquired either side of the mid-Judith discontinuity. As such, it seems likely that the study section corresponds in age to the lower to middle part of the Dinosaur Park Formation (although not necessarily lithostratigraphically correlated). A radiometric analysis of a newly discovered volcanic ash is currently underway, and it is hoped that this will provide definitive stratigraphic placement.

Regardless of the precise age of BDM 107, it can be expected to lie intermediate stratigraphically between *D. torosus* (known from the upper Oldman Formation, ~77.0 Ma; *Paulina Carabajal et al., 2021*) and *D. horneri* (known from the Two Medicine Formation, ~75.0 Ma; *Carr et al., 2017*).

**Diagnosis**

*D. wilsoni* can be assigned to *Daspletosaurus* based on the following characteristics: extremely coarse subcutaneous surface of the maxilla with no elevated ridges or corresponding fossae (*Carr et al., 2017; Voris et al., 2020*); cornual process of the postorbital approaching the laterotemporal fenestra (*Carr et al., 2017*); dorsal postorbital process of the squamosal terminating caudal to the rostral margin of the laterotemporal fenestra (*Carr et al., 2017; Voris et al., 2019*); and extremely coarse symphyseal surface of the dentary (*Voris et al., 2020*).

*D. wilsoni* possesses a single autapomorphy: a rostrocaudally elongate and dorsoventrally narrow mylohyoid foramen of the splenial (this foramen is much deeper in other *Daspletosaurus*, *Carr et al., 2017*; see below), and can additionally be diagnosed by a unique combination of ancestral and derived *Daspletosaurus* characteristics. *D. wilsoni* and *D. torosus* share a pneumatic inflation of the lacrimal reaching the medial edge of the bone (this inflation does not reach the medial edge of the bone in the holotype of *D. horneri*, but this may represent an allometric, ontogenetic, or taphonomic bias; E. A. Warshaw, 2022, unpublished data; *Carr et al., 2017*), cornual process of the postorbital approaching the laterotemporal fenestra (this process terminates much more rostrally relative to the fenestra in *D. horneri, contra Carr et al., 2017*; see below), cornual process of the postorbital subdivided into two distinct processes (this subdivision is absent in *D. horneri*, E. A. Warshaw, 2022, personal observations; see Description); prefrontal oriented rostromedially (determined from the angle of the prefrontal articular surface on the lacrimal of the holotype of *D. wilsoni*, which does not preserve a prefrontal; the prefrontal of *D. horneri* is oriented mediolaterally), pneumatic excavation of the squamosal that does not undercut its rostromedial margin (entire margin undercut in *D. horneri; Carr et al., 2017*), and quadratojugal lacking a pneumatic foramen in its lateral surface (although the presence of this foramen is highly intraspecifically variable in both *D. horneri* and *Tyrannosaurus*, such that further discoveries of *D. wilsoni* individuals may reveal its presence in this taxon; *Carr et al., 2017; Carr, 2020*). *D. horneri* and *D. wilsoni* share, to the exclusion of *D. torosus*, a premaxillary tooth row oriented entirely mediolaterally, such that all but one premaxillary tooth is concealed in lateral view (rostromedial orientation in *D. torosus* and less derived tyrannosaurids), antorbital fossa of the maxilla terminating at the rostral limit of the external antorbital fenestra (this fossa extends ahead of this
boundary onto the subcutaneous surface of the maxilla in *D. torosus* and less derived tyrannosaurs; *Carr et al., 2017*; E. A. Warshaw, 2022, unpublished data), rostro-dorsal ala of the lacrimal inflated (uninflated in *D. torosus* and less derived tyrannosaurs), ventral ramus of the lacrimal longer than the rostral ramus (determined largely by the height of the postorbital bar in the reconstructed skull, given that the ventral ramus is largely unpreserved in the holotype of *D. wilsoni*; the rostral ramus of the lacrimal is longer than the ventral ramus in *D. torosus; Carr et al., 2017*), short cornual process of the lacrimal (tall in *D. torosus*, although this process is taller in *D. wilsoni* than *D. horneri* and may best be described as intermediate between the previously named species of this genus; *Carr et al., 2017*), and dorsal quadrate contact of the quadratojugal visible in lateral view (concealed in *D. torosus* and less derived tyrannosaurs).

**Description**

Given the wealth of detailed osteologies describing tyrannosaurine specimens (e.g., *Carr (1999)*; *Brochu (2003)* and *Hurum & Sabath (2003)*), our description of the holotype of *D. wilsoni* places heavy emphasis on characteristics (or combinations of characteristics) unique to this specimen, as well as those that are otherwise taxonomically or phylogenetically informative within Tyrannosaurinae, so as to avoid the reiteration of plesiomorphic tyrannosaurine morphologies (or synapomorphies of *Daspletosaurus*) already described by previous authors (e.g., *Carr et al. (2017)*; *Voris et al. (2019)* and *Voris et al. (2020)*).

**Ontogenetic Stage of BDM 107**

In order to facilitate comparison with other tyrannosaurine individuals of equivalent ontogenetic stages (and in doing so, to avoid the misattribution of a phylogenetic signal to ontogenetically derived characteristics), brief comment is warranted on the ontogenetic stage represented by BDM 107; two lines of evidence suggest that this specimen is of advanced ontogenetic age. Firstly, BDM 107 is among the largest known *Daspletosaurus* individuals (articulated skull length 105 cm; *D. torosus* holotype CMN 8506 skull length 104 cm, *Voris et al., 2019*; *D. horneri* holotype MOR 590 skull length 89.5 cm, *Carr et al., 2017*). Although *Carr (2020)* criticized the use of size as an indicator of ontogenetic status in *Tyrannosaurus*, this criticism was based on the absence of a correlation between size and maturity among adult individuals; all the largest specimens of this genus were unambiguously recovered as adult by *Carr’s (2020)* analysis (i.e., within the final stages of ontogenetic development), such that this feature remains ontogenetically informative in distinguishing adults from juveniles and subadults. Secondly, BDM 107 displays several morphologies known otherwise to characterize mature tyrannosaurines, including a deeply scalloped maxilla-nasal suture (*Carr & Williamson, 2004; Carr, 2020*), a maxillary fenestra positioned rostrally within the antorbital fossa (*Carr, 2020*), a cornual process of the lacrimal inflated and positioned dorsal to the ventral ramus (*Carr, 1999; Currie, 2003; Carr, 2020*), and a grossly exaggerated cornual process of the postorbital (*Carr, 1999; Currie, 2003; Voris et al., 2019; Carr, 2020*). The totality of this evidence supports an adult ontogenetic stage or later for BDM 107 (adult sensu *Carr, 2020*; ontogenetic Stage 4 sensu *Carr, 1999*); this hypothesis may be tested in future work through histological analysis and/
or comparison with further discoveries of *D. wilsoni* individuals of different ontogenetic stages, both of which lie outside of the scope of the present study.

**Premaxilla**

The premaxillae of *D. wilsoni* are similar to those of *D. horneri* (Carr et al., 2017, Fig. 1; Fig. 2), *Tarbosaurus* (Hurum & Sabath, 2003, Fig. 3), and *Tyrannosaurus* (Brochu, 2003, Fig. 4) in that the alveolar row is oriented largely mediolaterally, such that the rostrum of

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**Figure 2**  *Premaxillae of BDM 107*. Shown in lateral (A), medial (B), and rostral (C) views. Abbreviations are as follows: nf, neurovascular foramina; smp, symphysis. Scale is 10 cm. DOI: 10.7717/peerj.14461/fig-2
the skull is broad and the labial surfaces of the premaxillary teeth face rostrally. In *Tyrannosaurus* and similarly derived tyrannosaurines (*Tarbosaurus* and *D. horneri*), the premaxillary teeth largely overlap each other in lateral view such that only the distalmost tooth is clearly visible; the same would be true of the holotype of *D. wilsoni*, were more than a single premaxillary tooth preserved within its socket. Conversely, the premaxillary tooth row of *D. torosus* and less derived tyrannosauroids is oriented rostromedially, such that multiple teeth are clearly visible in lateral view (*Voris et al., 2019*, Fig. 6).

Although previous authors have regarded a mediolaterally oriented premaxillary tooth row as a synapomorphy of Tyrannosauridae or more inclusive groups (*e.g.*, Carr et al., 2017: character 15), this is in error; mature specimens of *Gorgosaurus* (UALVP 10, Voris et al., 2022, Fig. 1; AMNH 5458, Matthew & Brown, 1923, Fig. 2) and *Qianzhousaurus* (GM F10004, Foster et al., 2021, Fig. 2), have rostromedially oriented premaxillary tooth rows such that in specimens with preserved teeth, all premaxillary teeth are visible in lateral view (although all tyrannosaurids do have premaxillary tooth rows oriented more medially than basal tyrannosauroids; this is the phylogenetic signal recorded in character 15 of Carr et al. (2017)). Comparison with other tyrannosaurids is hampered by the absence of preserved premaxillae and/or published descriptions of this element for several species (*e.g.*, Thanatotheristes, Voris et al., 2020; Dynamoterror, McDonald, Wolfe &

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**Figure 3** Left maxilla of BDM 107. Shown in lateral (A) and medial (B) views. Abbreviations are as follows: af, antorbital fossa; aof, antorbital fenestra; ma, maxillary antrum; mxf, maxillary fenestra; pmr, promaxillary recess; pmx, promaxillary fenestra; ps, palatal shelf; nf, neurovascular foramina; ns, neurovascular sulci. Scale is 10 cm. DOI: 10.7717/peerj.14461/fig-3
Dooley, 2018; Nanuqsaurus, Fiorillo & Tykoski, 2014; Lythronax and Teratophoneus, Loewen et al., 2013, for which all published specimens lack premaxillae); however, D. wilsoni and more derived tyrannosaurines (D. horneri, Tarbosaurus, Tyrannosaurus) represent the greatest exaggeration of the medial inclination of the premaxillary tooth row among tyrannosaurids for which comparative material is available (although this condition, with only one clearly visible premaxillary tooth in lateral view, is present in at least one Gorgosaurus: TCMI 2001.89.1, Voris et al., 2022, Fig. 10). D. torosus is intermediate between the (presumably) ancestral rostromedial orientation and the mediolateral condition of later Daspletosaurus species; two to three premaxillary teeth are visible in lateral view in the holotype specimen, CMN 8506 (Carr & Williamson, 2004, Fig. 6; Voris et al., 2019, Fig. 6).

It should be noted that the orientation of the premaxillary tooth row is not necessarily equivalent to the orientation of the premaxillae themselves. In Tyrannosaurus AMNH 5027, for example, the premaxillae appear to be rostromedially oriented in dorsal view (Carr & Williamson, 2004, Fig. 7); however, the premaxillary alveoli are mediolaterally arranged when viewed ventrally (E. A. Warshaw, 2022, personal observations; Osborn, 1912, Fig. 5A; Molnar, 1991, Fig. 9A).

The taxonomic utility of this character is a hypothesis that will require further testing as individuals of D. wilsoni and other tyrannosaurids with preserved premaxillae are discovered; notably, two specimens previously referred to D. torosus display the derived condition (mediolateral orientation), sharing it with D. wilsoni and more derived tyrannosaurines: FMNH PR308 (Matthew & Brown, 1923, Fig. 5; Carr, 1999, Fig. 1) and TMP 2001.36.1 (Voris et al., 2019, Fig. 6). If these individuals were to represent D. torosus, the distinction between this species and D. wilsoni in the orientation of the premaxillary tooth row would be heavily undermined; however, both of these specimens have previously been noted as belonging to a novel taxon from the Dinosaur Park Formation (FMNH PR308, Currie, 2003; TMP 2001.36.1, Paulina Carabajal et al., 2021). Therefore, although relevant comparisons will be made with these specimens hereafter, they will be considered separately from D. torosus (and will be referred to below as the Dinosaur Park taxon).
precise taxonomic designation for these specimens is reserved for future work in accordance with comments by previous authors (Currie, 2003; Paulina Carabajal et al., 2021).

There is a small (~2 cm diameter) indentation in the nasal process of the right premaxilla of BDM 107; this is most likely pathological, as it is irregular in form and not present on the left premaxilla.

**Maxilla**

As in other *Daspletosaurus*, the subcutaneous surface of the maxilla in *D. wilsoni* is densely covered in anastomosing sulci extending from neurovascular foramina (Carr et al., 2017; Voris et al., 2020; Fig. 3). The degree of sculpturing of this surface in BDM 107 is
similar to CMN 8506 (*D. torosus*), although in the former, there is no smooth region rostral to the external antorbital fenestra indicating a rostral continuation of the antorbital fossa as *D. torosus* and alioramins (*Carr et al., 2017*). As in *Thanatotheristes* and other *Daspletosaurus* species, the shallow excavations that characterize the maxillae of the most derived tyrannosaurines (*Zhuchengtyrannus, Tyrannosaurus, Tarbosaurus; Hone et al., 2011; Voris et al., 2020*) are absent from the holotype maxilla of *D. wilsoni*. Also absent are the textural ridges present on the maxillae of *Zhuchengtyrannus* (*Hone et al., 2011*), *Tarbosaurus, Tyrannosaurus*, and *Thanatotheristes* (*Voris et al., 2020*), but not any *Daspletosaurus* species.

The rostral end of the maxilla of BDM 107 is bowed subtly medially towards its contact with the premaxilla and nasal; this may be a structural consequence of the greater medial inclination of the premaxillary tooth row (see above), as a similar condition characterizes

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**Figure 6** Left postorbital of BDM 107. Shown in lateral (A), dorsal (B), medial (C), caudal (D), and rostral (E) views. Abbreviations are as follows: cdt, caudodorsal tuberosity; dtf, dorsotemporal fossa; fc, frontal contact; lsc, laterosphenoid contact; sop, subocular process; sos, supraorbital shelf. Scale is 10 cm.
D. horneri (MOR 590, E. A. Warshaw, 2022, personal observations), Tarbosaurus (Hurum & Sabath, 2003, Fig. 15), and Tyrannosaurus (MOR 008, MOR 980, E. A. Warshaw, 2022, personal observations). Tyrannosaurids with more rostromedially inclined premaxillary tooth rows lack this bowing (e.g., D. torosus CMN 8506, J. T. Voris, 2022, personal communication).

The maxilla of BDM 107 is irregular relative to other species of Daspletosaurus in that it is proportionally elongate, being 64.1 cm in length and 24.8 cm in height (ratio of length to height = 2.6). This bone is 58.6 cm long rostrocaudally and 27.5 cm tall dorsoventrally in the holotype of D. horneri (ratio of length to height = 2.1; MOR 590, Carr et al., 2017). Given the broad range of variation in the proportions of this element in other tyrannosaurine species for which larger sample sizes are known (e.g., Tyrannosaurus; Carpenter, 1990; Paul, Persons & Van Raalte, 2022; E. A. Warshaw, 2022, personal observations), this characteristic was not included as an autapomorphy of D. wilsoni. Consistency in this trait across further discoveries of D. wilsoni individuals may require a reevaluation of the taxonomic utility of this character.

D. wilsoni possesses 15 maxillary alveoli, as in other species of Daspletosaurus (Carr et al., 2017). The 13th alveolus bears a swollen abscess in BDM 107, and the 15th maxillary tooth conceals a small replacement tooth within its root that is visible in medial (lingual) view. In general, the maxillary teeth are similar to those of other tyrannosaurid species in being labiolingually broad, although not to the degree present in more derived tyrannosaurines (e.g., Tyrannosaurus and Tarbosaurus), in which the labiolingual width of the maxillary teeth is subequal to their mesiodistal length (Carr et al., 2017). The first maxillary alveolus is not small and also bears an incrassate tooth (i.e., it does not bear a d-shaped crown similar to those present in the premaxillae, as in Gorgosaurus; Currie, 2003; Voris et al., 2022).
Jugal

The jugal of *D. wilsoni* is most similar to that of *D. torosus* among tyrannosaurines in that it has a mediolaterally thin ventral margin of the orbit (as opposed to a rounded margin as in *Thanatotheristes, Lythronax*, most *Tarbosaurus*, and some *Tyrranosaurus; Voris et al., 2020; Voris et al., 2022; J. T. Voris, 2022, personal communication*). A thin ventral margin of the orbit likely represents the ancestral tyrannosaurid condition, (as it is also present in *Bistahieversor, Albertosaurus, Gorgosaurus*, and *D. horneri*; J. T. Voris, 2022, personal communication) and does not bow medially along its rostrocaudal length (the jugals of *D. horneri, Tyrannosaurus*, and *Tarbosaurus* are angled rostromedially rostral to the orbit, such that the maxillae are medially inset from the orbitotemporal region; *D. horneri* MOR 590, E. A. Warshaw, 2022, personal observations; *Tyrannosaurus* AMNH 5027, *Molnar, 1991*, Fig. 9; *Tarbosaurus* GIN 107/1, *Hurum & Sabath, 2003*, Fig. 15; E. A. Warshaw, 2022, unpublished data).

As in *D. torosus*, the caudal portion of the lacrimal contact surface of the jugal is shallowly inclined (Fig. 4); this surface is very steep in *D. horneri*, as well as in *Albertosaurus* and *Gorgosaurus* (*Carr et al., 2017*). Although *Carr et al. (2017)* recovered this feature as unique to *D. horneri* among tyrannosaurines, it is also present in some *Tyrranosaurus* individuals (MOR 980, MOR 1125, AMNH 5027, E. A. Warshaw, 2022, personal observations).

Lacrimal

As in all tyrannosaurids except for *D. horneri, Tarbosaurus*, and *Tyrranosaurus*, the cornual process of the lacrimal in *D. wilsoni* is large and rises to a distinct apex along its dorsal margin (*Carr et al., 2017, Fig. 5*). This apex is situated directly dorsal to the lacrimal’s ventral ramus, as is characteristic of mature tyrannosaurines (*Currie, 2003; Carr,*
The cornual process of the lacrimal is shorter in *D. wilsoni* (5.2 cm from the dorsal margin of the lacrimal antorbital recess to the apex of the cornual process in BDM 107) than *D. torosus* (6.9 cm, CMN 8506; Voris et al., 2019, Fig. 6), but similar to the Dinosaur Park taxon (5.1 cm, TMP 2001.36.1; Voris et al., 2019, Fig. 6) (these three specimens are...
each within 2 cm of each other in skull length, such that measurements of this process need not be corrected for differences in absolute specimen size; see above; Voris et al., 2019, Fig. 6). The lacrimal cornual process of the D. horneri holotype MOR 590 is shorter still (3.7 cm; Carr et al., 2017, Fig. 1), although it should be noted that this specimen is also ~15% shorter in skull length than any of the specimens previously mentioned (Carr et al., 2017; Voris et al., 2019; see above), such that the difference in this feature between D. horneri and other Daspletosaurus is relatively less pronounced than isolated measurements of this process would suggest (scaled isometrically to the same skull length as MOR 590, however, BDM 107 would still have a taller cornual process of the lacrimal, at 4.4 cm).

Carr et al. (2017) regarded an accessory cornual process of the lacrimal as a synapomorphy of Daspletosaurus. However, this process is indistinguishable from the caudally directed supraorbital process of the lacrimal upon which it is purported to sit; the supraorbital processes of the lacrimals of Tyrannosaurus (MOR 555, MOR 980, MOR 1125, AMNH 5027, E. A. Warshaw, 2022, personal observations), Tarbosaurus (ZPAL MgD-1/4, Hurum & Sabath, 2003, Fig. 6), and Teratophoneus (UMNH VP 16690, Loewen et al., 2013, Fig. 3) are all morphologically identical to those of Daspletosaurus, although they are scored by Carr et al. (2017) as lacking an accessory cornual process. In lieu of any quantitative demonstration of this process’s presence in Daspletosaurus, the taxonomic utility of this character is rejected here.
The lacrimal antorbital recess differs in morphology from *D. torosus*, but is similar to that of *D. horneri*, *Tarbosaurus*, and *Tyrannosaurus* in that the rostrodorsal ala joining the rostral and ventral rami of the lacrimal is inflated into a cylindrical bar that is elevated in relief relative to the rest of the recess (this ala is inflated in *D. torosus*, but to a lesser degree such that no discrete bar is formed between the rostral and ventral rami; Carr et al., 2017, Fig. S2F) (E. A. Warshaw, 2022, personal observations). This feature is also present in the Dinosaur Park taxon (TMP 2001.36.1, Voris et al., 2019, Fig. 6). Also distinguishing the lacrimal of *D. wilsoni* from *D. torosus* is a ventrally directed antorbital fossa in the latter. The lacrimal antorbital fossa is laterally directed in other tyrannosaurids, including *D. wilsoni*, the Dinosaur Park taxon (TMP 2001.36.1, Voris et al., 2019, Fig. 6), *D. horneri* (MOR 590 and MOR 1130, Carr et al., 2017, Fig. 3), *Tyrannosaurus*, *Tarbosaurus*, *Albertosaurus*, and *Gorgosaurus* (Carr & Williamson, 2004, Fig. 10).

Rostrally, the ventral process of the lacrimal rostral ramus is unique in *D. wilsoni* in having a rounded distal end; this process comes to a pronounced tip in most tyrannosaurids (Carr, Williamson & Schwimmer, 2005, Fig. 8; Loewen et al., 2013, Fig. 3), with the possible exception of *D. horneri*, in which the holotype specimen MOR 590 has a pointed ventral process and that of the paratype MOR 1130 is rounded (E. A. Warshaw, 2022, personal observations; Carr et al., 2017, Figs. 2C and 3). Given the eminent possibility of taphonomic alteration of this feature (i.e., “rounding down” of a pointed ventral process into a rounded one by abrasion prior to burial), exaggerated by the small size of the ventral process of the lacrimal, this feature is excluded from consideration either as an autapomorphy of *D. wilsoni* or as uniting this species with *D. horneri*.

Caudodorsally, the prefrontal articular surface of the lacrimal can be used to determine the orientation of the long axis of the prefrontal. In *D. wilsoni* and *D. torosus*, this element is oriented rostrocaudally (Carr & Williamson, 2004, Fig. 8). This condition is shared with the Dinosaur Park taxon (TMP 2001.36.1, Paulina Carabajal et al., 2021, Fig. 2D), and is
also present in *Gorgosaurus* (UALVP 10, *Voris et al.*, 2022), *Teratophoneus* (UMNH VP 16690, *Loewen et al.*, 2013, Fig. 3), and *Qianzhousaurus* (GM F10004, *Foster et al.*, 2021, Fig. 3). Conversely, the prefrontal is oriented rostromedially or mediolaterally in *D. horneri* (MOR 590, E. A. Warshaw, 2022, personal observations; *Carr et al.*, 2017, Fig. 1).
Tarbosaurus (ZPAL MgD-I/4, Hurum & Sabath, 2003, Fig. 1), and Tyrannosaurus (AMNH 5027, E. A. Warshaw, 2022, personal observations; Carr & Williamson, 2004, Fig. 8), as well as at least one specimen of Albertosaurus (TMP 1981.10.1, Carr & Williamson, 2004, Fig. 8).

Postorbital

The postorbital of D. wilsoni is most similar to that of D. torosus and the Dinosaur Park taxon in bearing a massive cornual process that approaches the rostral margin of the laterotemporal fenestra caudally (Fig. 6; Carr et al., 2017; D. torosus CMN 8506, Voris et al., 2019, Fig. 6; TMP 2001.36.1, Voris et al., 2019, Fig. 4). Carr et al. (2017) proposed a cornual process of the postorbital approaching the laterotemporal fenestra as a synapomorphy of Daspletosaurus; however, the cornual process of the postorbital does not approach the laterotemporal fenestra in the holotype of D. horneri (MOR 590, E. A. Warshaw, 2022, personal observations), and is instead broadly separated from it as in Tyrannosaurus (MOR 980, MOR 1125, MOR 555, E. A. Warshaw, 2022, personal observations) and Tarbosaurus (ZPAL MgD-I/4, Hurum & Sabath, 2003, Fig. 1).

Also shared between D. wilsoni, D. torosus, and the Dinosaur Park taxon is the subdivision of the postorbital cornual process into two discrete processes: a supraorbital shelf protruding from the dorsal margin of the orbit and a caudodorsal tuberosity emerging more caudoventrally (Fig. 6; Voris et al., 2019, Fig. 4D), creating a sinusoidal relief when the postorbital is viewed rostrally or caudally. Both the supraorbital shelf and the caudodorsal tuberosity are situated upon a more ‘typical’ tyrannosaurine cornual process; that is, they lie lateral to a gross swelling of the postorbital similar to that present in other tyrannosaurines (e.g., Tyrannosaurus, MOR 1125, MOR 980, MOR 555, MOR 008, E. A. Warshaw, 2022, personal observations). The caudodorsal tuberosity overhangs its caudoventral base, creating a crease between this process and the underlying body of the postorbital; a similar condition is present in the postorbital cornual processes of Gorgosaurus, Teratophoneus, and Bistahieversor (Voris et al., 2022; J. T. Voris, 2022, personal communication), but not in D. horneri (MOR 590, Carr et al., 2017, Fig. 1), Tyrannosaurus (MOR 1125, MOR 980, MOR 555, MOR 008, E. A. Warshaw, 2022, personal observations), or Tarbosaurus (ZPAL MgD-I/4, Hurum & Sabath, 2003, Fig. 8). A similar crease forms between the body of the postorbital and the cornual process of Tyrannosaurus in specimens with an epipostorbital (sensu Carr, 2020; AMNH 5027, Molnar, 1991; Carr, 2020); however, no such element is present in the holotype of D. wilsoni (or any other Daspletosaurus specimens; E. A. Warshaw, 2022, personal observations).

The ventral ramus of the postorbital tapers ventrally to a point in D. wilsoni, as in other Daspletosaurus (Carr, 1999), including the Dinosaur Park taxon (Voris et al., 2019, Fig. 4), and in contrast to the enormous subocular process of the postorbital that projects rostrally in Tyrannosaurus (Carr, 2020), Tarbosaurus (Hurum & Sabath, 2003, Fig. 8), Gorgosaurus (Voris et al., 2022), Teratophoneus (Loewen et al., 2013, Fig. 3), and Albertosaurus (Currie, 2003). Although the subocular process is present in D. wilsoni (and other Daspletosaurus), it is small relative to those of other tyrannosaurids (Fig. 6).
Squamosal

The squamosal of *D. wilsoni* is indistinguishable from that of *D. torosus* in that the rostralmost extent of the postorbital contact surface terminates caudal to the rostral margin of the laterotemporal fenestra (also in *D. horneri*; Carr et al., 2017), the rostromedial margin of the pneumatic recess on the ventral surface is not undercut (Fig. 7), and the caudal process is pneumatized (as evidenced by pneumatic foramina in the process’s rostromedial surface; Carr et al., 2017). No characteristics or combinations of characteristics unique to *D. wilsoni* are observable on this element.

Quadratojugal

The quadratojugal is conservative morphologically across tyrannosaurids (Loewen et al., 2013). However, a single characteristic of the quadratojugal of *D. wilsoni* unites it with *D. horneri* and at least one specimen of the Dinosaur Park taxon (TMP 2001.36.1), and differs from the condition in *D. torosus* and less derived tyrannosaurids: a dorsal quadrate contact that is broadly visible in lateral view. In most tyrannosauroids, the dorsal quadrate contact of the quadratojugal is directed medially or rostromedially such that it is obscured by the body of the quadratojugal in lateral view. In *D. wilsoni*, *D. horneri* (MOR 590, Carr et al., 2017, Fig. 1), and TMP 2001.36.1 (Voris et al., 2019, Fig. 6), however, this process is directed caudomedially, exposing it laterally (Fig. 8).

The dorsal quadrate contact is marginally visible laterally in the holotype of *D. torosus*, CMN 8506 (Voris et al., 2019, Fig. 6; J. T. Voris, 2022, personal communication), but not nearly to the extent observable in the aforementioned taxa. The condition in *D. torosus* may therefore represent individual variation on the caudomedial orientation of most tyrannosaurids, or a structural antecedent to the condition present in other species of *Daspletosaurus*.

The caudomedial orientation of the dorsal quadrate contact is reversed in the paratype specimen of *D. horneri*, in which this process is hidden in lateral view (MOR 1130, Carr et al., 2017, Fig. S2K). Given that this specimen is younger stratigraphically than the holotype (MOR 590; Carr et al., 2017), this reversal may represent a phylogenetic signal (although it may instead represent intraspecific variation). *Tarbosaurus* and *Tyrannosaurus* share this feature with MOR 1130 (E. A. Warshaw, 2022, personal observations).

Quadrate

No discrete morphological characters distinguish the quadrate of *D. wilsoni* from those of its closest relatives. As in other derived tyrannosaurines, the quadrate is massive, with a shallow fossa on its medial surface and a pronounced pneumatic foramen (and surrounding fossa) at the rostral confluence of the mandibular condyles and the orbital process (Fig. 9; Carr et al., 2017). The paraquadrate foramen, bounded medially by the quadrate and laterally by the quadratojugal, is small and teardrop-shaped; only its lateral margin is made up by the quadratojugal, as the quadrate forms the dorsal and ventral borders of the foramen.

Although no palatal elements are known, the medial deflection of the quadrate’s pterygoid wing allows an approximation of the position of the pterygoids relative to the facial skeleton, and suggests a broad orbitotemporal region, as in other tyrannosaurines.
Dentary

The dentary of *D. wilsoni* is deep, with a relatively straight ventral margin and a dorsal (alveolar) margin that trends caudodorsally, increasing the depth of the mandible caudally (Fig. 10). As in other *Daspletosaurus*, the texturing of the dentary symphysis is more exaggerated in *D. wilsoni* than non-*Daspletosaurus* tyrannosaurines (e.g., *Tyrannosaurus, Thanatotheristes; Voris et al., 2020*), and is composed of several interlocking (presumably, as only the left dentary is known) ridges and cusps. There are 17 dental alveoli, as in *D. horneri* (*Carr et al., 2017*), and a sharp, narrow Meckelian groove with a rugose knob caudoventral to its rostral end. This knob is present in both other species of *Daspletosaurus*, as well as *Tyrannosaurus, Tarbosaurus*, and *Zhuchengtyrannus magnus*, but not *Thanatotheristes* or more basal tyrannosaurids (*Carr et al., 2017; Voris et al., 2020*).

The lateral surface of the dentary of BDM 107 bears two intersecting grooves caudoventral to the caudal termination of the alveolar margin (Fig. 10); the edges of these grooves are beveled and are likely pathological. They may represent bite marks, as have been described on the craniofacial bones of other tyrannosaurids (*Voris et al., 2020; Brown, Currie & Therrien, 2022*).

Splenial

The splenial of BDM 107 is typical of *Daspletosaurus* except in the size and form of the mylohyoid foramen (Fig. 11), an autapomorphy of this taxon. In most derived tyrannosaurines, including *D. torosus* and *D. horneri*, this foramen is extremely large, roughly the same dorsoventral depth as the rostral process of the splenial (*Carr et al., 2017*). In *D. wilsoni*, however, the foramen is dorsoventrally shallow, and rostrocaudally elongate, such that it is ellipsoid in form and roughly half the dorsoventral depth of the splenial’s rostral process. This is most similar to the condition in alioramins (*Brusatte, Carr & Norell, 2012*) and *Appalachiosaurus* (*Carr, Williamson & Schwimmer, 2005*).

Cervical vertebrae

Four cervical vertebrae are preserved in BDM 107 from the cranial-middle portion of the series. No atlas or axis were found. As in all tyrannosaurids, the spinous processes of the cervical vertebrae are subequal in dorsoventral height to their corresponding centra. Both the spinous processes and the centra are craniocaudally short, similar to and most exaggerated in the cervical vertebrae of *Tyrannosaurus* (see *Brochu, 2003*, and figures therein). As in *Tyrannosaurus* (and other large tyrannosaurids), the cranial and caudal faces of the cervical centra in BDM 107 are dorsoventrally displaced from one another in order to create the characteristic ‘S-curve’ of the neck, and the cranial cervical centra are extremely foreshortened craniocaudally (*i.e.*, much taller than long). This indicates a robustly built cranial portion of the neck, presumably in order to support the weight of the head.

Sacral vertebrae

The spinous processes of two sacral vertebrae are preserved. Both are sub-rectangular in form and bear rugose knobs near their apices, presumably the ossified bases of sacral ligaments.
Methods

The holotype specimen was collected under permit MTM 108829-e6 issued to DF by The US Bureau of Land Management.

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A cladistic phylogenetic analysis was conducted using the character matrix of Carr et al. (2017) (with modifications from Voris et al. (2020)), with additional modifications based on personal observation of specimens made by the lead author, including the addition to the character matrix of several proposed autapomorphies of D. horneri noted by Carr et al. (2017) to occur more broadly across Tyrannosauridae (see Supplemental Information for a comprehensive list of modifications). The analysis was run in TnT v1.5 (Goloboff, Farris & Nixon, 2008) using a “New Technology” search with settings identical to those of Voris et al. (2020) (ratchet, tree drift, tree fusing, and sectorial search set to default, and set to recover minimum length 10 times). Support for recovered clades was tested using bootstrapping with 1,000 replicates under a traditional search.

RESULTS

The cladistic analysis produced 12 Most Parsimonious Trees (MPTs; best score: 853). Within the strict consensus of these trees, the least inclusive clade containing Dynamoterror and Tyrannosaurus (i.e., all of Tyrannosaurinae more derived than Alioramus) was recovered as a large polytomy, with a sister relationship retained between Tyrannosaurus and Tarbosaurus, and Dynamoterror, Lythronax, and Teratophoneus recovered in a trichotomy (see Supplemental Information: Fig. S1).

Given the fragmentary nature of their respective holotypes (scored for <15% of characters), Nanuqsaurus hoglundi and Thanatotheristes were removed from the dataset (inclusion of either taxon collapsed the tree as above), and an additional analysis was conducted with the same settings. This analysis produced two MPTs (best score: 846), and recovered D. wilsoni as sister to a clade formed by D. horneri and more derived tyrannosaurs (Zhuchengtyrannus, Tarbosaurus, Tyrannosaurus). Alioramins were recovered within a polytomy, as were Dynamoterror, Teratophoneus, and Lythronax; all other topological relationships were as in Voris et al. (2020) (Fig. 12).

Bootstrapping of this result showed weak support (<70) for all clades within Tyrannosaurinae except for alioramini (90), derived tyrannosaurs (Daspletosaurus + (Zhuchengtyrannus (Tyrannosaurus + Tarbosaurus))) (82), tyrannosaurs more derived
than *Daspletosaurus* (85), and *Tyrannosaurus + Tarbosaurus* (85). Recovered support was particularly weak (≤9) for the interrelationships of *Daspletosaurus* (Fig. 12).

A single autapomorphy of *D. wilsoni* was recovered by the cladistic analysis: mylohyoid foramen of the splenial elongate and rostrocaudally ovoid (this foramen is much deeper in other *Daspletosaurus* species; see above).

The *D. wilsoni* + more derived tyrannosaurines clade was recovered with the following three synapomorphies: dorsoventrally tall orbit; mediolaterally oriented tooth row of the premaxilla; and short cornual process of the lacrimal. A further four synapomorphies united *D. horneri* and more derived tyrannosaurines to the exclusion of *D. wilsoni*: rostrilaterally directed orbits (resulting from the rostromedial bowing of the jugal); cornual process of the postorbital swollen and terminating far rostral to the laterotemporal fenestra; first interdental plate of the maxilla narrow, and second plate truncated (both plates are subsequently expanded in tyrannosaurines more derived than *D. horneri*); and mediolaterally oriented prefrontal. Additional autapomorphies of relevant taxa and synapomorphies of relevant clades are available in Supplemental Information.

**DISCUSSION**

Several aspects of the results presented here contrast with (or supplement) those of previous analyses, and therefore deserve mention. Noticeably, the results of the cladistic analysis place *Tyrannosaurus*—line tyrannosaurines (*Zhuchengtyrannus*, *Tarbosaurus*, and *Tyrannosaurus*) as successive sister taxa to *Daspletosaurus* (contra Carr et al., 2017, and Voris et al., 2020, both of which recovered these as sister lineages), and recovers a paraphyletic *Daspletosaurus*; these aspects of the results are the topic of a study by the lead author currently in review, and will not be discussed here (although it should be noted that similar results were recovered by Horner, Varricchio & Goodwin (1992), and the Bayesian analysis of Brusatte & Carr (2016), Loewen et al. (2013) also recovered a paraphyletic *Daspletosaurus*. Should this paraphyly be upheld by future studies, *D. wilsoni* and *D. horneri* may be assigned new genera in order to preserve monophyly; *D. wilsoni* is assigned to *Daspletosaurus* here to avoid the creation of a polyphyletic *Daspletosaurus* and for ease of discussion and comparison with its closest relatives). Instead, only the interrelationships and evolutionary history of *Daspletosaurus* are considered below.

Though not included in the cladistic analysis, the Dinosaur Park taxon agrees with *D. wilsoni* in several characters which differ in both of these taxa from the condition in *D. torosus* (see Description), including the orientation of the premaxillary tooth row, the height of the cornual process of the lacrimal, the inflation of the rostrodorsal ala of the lacrimal, and lateral exposure of the dorsal quadrate contact of the quadratojugal. All of these characters are also shared with *D. horneri*, although *D. wilsoni* and the Dinosaur Park taxon also share (to the exclusion of *D. horneri*) a cornual process of the postorbital that approaches the laterotemporal fenestra and is subdivided into a caudodorsal tuberosity and a supraorbital shelf, and a prefrontal that is oriented rostrocaudally rather than rostromedially or mediolaterally (both of these characters are also present in *D. torosus*). Similarity in all of these features suggests a close affinity between *D. wilsoni* and the Dinosaur Park taxon, although this could reflect either taxonomic synonymity or a
genuine sister relationship; this designation is reserved for future studies centered on the Dinosaur Park taxon (noted as forthcoming by Currie (2003) and Paulina Carabajal et al., 2021), which has yet to receive a formal description and may reveal autapomorphies (or synapomorphies with D. wilsoni) not considered here.

Should the Dinosaur Park taxon be demonstrated to represent a distinct species from D. wilsoni, it would potentially represent the first known instance of contemporaneity between more than one species of Daspletosaurus (Carr et al., 2017), given that the D. wilsoni holotype was preserved in strata likely corresponding in time to the deposition of the Dinosaur Park Formation (at least in part; see Geologic Context). However, this possibility rests both upon the taxonomic distinctiveness of the Dinosaur Park taxon and the absence of fine-scale stratigraphic separation between this species and D. wilsoni, both of which require additional study to confirm or deny (e.g., a formal description of the anatomy of the Dinosaur Park taxon and precise stratigraphic placement of individuals of this taxon and D. wilsoni). Discussion below will therefore exclude this possibility from consideration, although resulting hypotheses will be subject to revision should this exclusion prove to be erroneous.

Among described Daspletosaurus species, D. wilsoni fulfills the predictions made by Carr’s et al. (2017) hypothesis of anagenesis between D. torosus and D. horneri. Namely, D. wilsoni is stratigraphically, phylogenetically, and morphologically intermediate between these taxa (see Geologic Context, Results, and Diagnosis, respectively), and occurs within the same general geographic range (all three species of Daspletosaurus are found within Montana or Alberta; Carr et al., 2017). These points correspond to the criteria proposed by Carr et al. (2017) (and later Zietlow, 2020) for defensible hypotheses of anagenesis: (1) lack of stratigraphic overlap (but see above), (2) close phylogenetic relationships, (3) intermediate morphologies, and (4) similar geographic ranges. It should be noted that while the fulfillment of these criteria establishes anagenesis as a defensible hypothesis, it does not preclude cladogenesis in Daspletosaurus as the driving factor of the evolution of this genus (with successively more derived clades, e.g., D. wilsoni and more derived tyrannosaurines, representing cladogenetic events rather than portions of an anagenetic sequence).

However, several alternative lines of evidence are consistent with anagenesis and inconsistent with cladogenesis, and therefore strengthen the hypothesis of anagenesis as a predominant evolutionary mode in Daspletosaurus. Firstly, as noted by Wagner, Erwin & Anstey (1995), cladogenesis via punctuated equilibrium (with species diverging from an ancestral taxon in morphological stasis; Eldredge & Gould, 1972) can be identified by the presence of polytomies in a recovered cladogram, since descendant species of an ancestor in stasis will not form subclades. No polytomy was recovered within the clade formed by Daspletosaurus and more derived tyrannosaurines (Fig. 12); the origination of sampled Daspletosaurus species from a common ancestor in stasis can therefore be rejected based on the topology of the recovered cladogram alone. Secondly, D. wilsoni almost entirely lacks autapomorphies, displaying only a single feature not also present in either D. torosus or D. horneri (a rostrocaudally elongate mylohyoid foramen of the splenial, recovered by the cladistic analysis; see Results). Wagner, Erwin & Anstey (1995) and Szalay (1977) noted
that ancestors should lack apomorphies relative to descendants, such that a paucity of
autapomorphies suggests that an ancestral taxon has been sampled. Similarly, Wilson,
Ryan & Evans (2020) noted the absence of autapomorphies in several centrosaurine taxa
hypothesized therein to represent an anagenetically evolving lineage, with stratigraphically
successive taxa being defined by combinations of plesiomorphic and apomorphic
characters rather than species-level autapomorphies (forming “metaspecies;” Horner,
Varrichio & Goodwin, 1992; Wilson, Ryan & Evans, 2020). Indeed, a hypothesis of
cladogenesis at the root of the clade formed by D. wilsoni and more derived
tyrannosaurs would rest entirely upon the elongate mylohyoid foramen of this species
as evidence of divergence from other Daspletosaurus; in the absence of additional
characters supporting this hypothesis, the sole autapomorphy of D. wilsoni may
alternatively represent individual variation, or a character evolved within this species and
lost before the appearance of D. horneri (similar to the lateral exposure of the dorsal
quadrate contact of the quadratojugal, present in D. wilsoni and the holotype of D. horneri,
but not in the stratigraphically sequential paratype specimen or more derived
tyrannosaurs; see Description). The morphological evidence for a cladogenetic origin of
D. wilsoni is therefore weak; the blend of ancestral and derived characteristics in this
species and the near total absence of autapomorphies is more consistent with anagenesis
between stratigraphically antecedent (D. torosus) and subsequent (D. horneri) taxa.

In light of this evidence, we propose that the three species of Daspletosaurus represent
an anagenetically evolving lineage (Fig. 13); as noted above, this hypothesis will be subject
to revision following further study into the phylogenetic affinities of species within the
genus, additional discoveries of Daspletosaurus individuals from stratigraphically
intermediate horizons (which under an anagenetic model, should be intermediate in
morphology between species), and characterization of the range of individual variation
present in relevant characters proposed here to represent species-level autapomorphies or
morphological transitions between taxa.

Should branching events (i.e., cladogenesis) within Daspletosaurus be demonstrated by
future studies or discoveries (e.g., if the Dinosaur Park taxon is demonstrated to be both
distinct from and contemporaneous with D. wilsoni), this would not necessarily exclude
anagenesis from playing a role in the generation of morphological novelty within the
genus. Wagner, Erwin & Anstey (1995) noted the presence of anagenetic change between
branching events in plesiomorphic lineages (=ancestral lineages; the lineage from which
cladogenetically derived taxa branch) not in morphological stasis, which led these authors
to designate this pattern of speciation as “bifurcation,” reserving “cladogenesis” for
branching from morphologically static ancestral taxa. Although we do not adopt their
terminology, we agree that anagenesis can operate in concert with cladogenesis in order to
produce observed patterns of macroevolutionary change. In the case of Daspletosaurus,
while autapomorphies of individual species may represent the results of cladogenesis, the
synapomorphies of progressively more exclusive clades within the genus (e.g., coarse
symphyseal texture of the dentary in Daspletosaurus, inflated rostroventral ala of the
lacrimal in D. wilsoni + D. horneri, etc.) would remain anagenetically derived under a
typically cladogenetic model. Anagenesis therefore enjoys a predominant role in the
evolution of derived morphologies within derived tyrannosaurines, regardless of the presence of branching events within *Daspletosaurus* (in contrast to morphologically static genera, in which morphological change is concentrated at the base of cladogenetic events; *Eldredge & Gould, 1972*).

The low bootstrap support recovered for the results of the cladistic analysis may also be readily explained in the context of anagenesis. As noted by *Soltis & Soltis (2003)*, low bootstrap scores may be recovered for otherwise well-supported clades (e.g., clades recovered within all MPTs, as in all of the interrelationships of *Daspletosaurus* recovered here) if they are supported by few characters, given that the chance of supporting characters being included in a bootstrap resample is lower with fewer characters. This is a common occurrence among recently diverged clades which have not had much time to accrue synapomorphies (*Soltis & Soltis, 2003*), but the same would apply to an anagenetically evolving *Daspletosaurus*; should *D. wilsoni* represent a descendant of *D. torosus* as proposed here, then all of the synapomorphies of the *D. wilsoni* + more derived tyrannosaurines clade must have been evolved within the ~500 kyr window between *D. torosus* and *D. wilsoni* (it should be noted that this would also be true in the case of recent divergence of the *D. wilsoni* + more derived tyrannosaurine clade via cladogenesis; therefore, low bootstrap scores cannot be taken as evidence of anagenesis, but are at least consistent with it).

More generally, as sampling of a lineage increases, the temporal windows between sampled taxa must necessarily be reduced, and synapomorphies of progressively more derived clades will therefore be fewer (with the same number of character changes distributed among more clades as sampling increases), such that bootstrap scores can be expected to correlate negatively with sampling intensity. To this point, removal of *D. wilsoni* (in addition to *Nanuqsaurus* and *Thanatotheristes*, as described above; see Results) from the cladistic analysis recovers an identical tree topology, but increases bootstrap support for the *D. horneri* + more derived tyrannosaurines clade from 8 to 25 (still a low score, but over three times higher).

Bootstrap scores can also be affected by the inclusion of characters irrelevant to the node in question (*Soltis & Soltis, 2003*). The phylogenetic character matrix of *Carr et al. (2017)* used here contains characters informative across Tyrannosauroidea, including hundreds of characters that are not informative within *Daspletosaurus* or derived Tyrannosaurinae in general (i.e., characters not recovered as autapomorphies or synapomorphies for species or groups within this clade, respectively). We therefore regard the low bootstrap scores recovered for the phylogenetic placement of *D. wilsoni* not as evidence of an erroneous result, but as an expected consequence of higher taxonomic resolution among derived tyrannosaurines and the nature of the data matrix used.

**CONCLUSIONS**

*D. wilsoni* sp. nov., a stratigraphic and morphological intermediate between *D. torosus* and *D. horneri*, is hypothesized to represent a transitional form along an anagenetic lineage linking both previously named species of *Daspletosaurus*. This finding, in concert with previous identifications of anagenesis in contemporary dinosaur lineages, emphasizes the
explanatory power of anagenesis in the production of evolutionary trends among 
dinosaurs of the Late Cretaceous Western Interior (Scannella et al., 2014; Freedman Fowler & Horner, 2015; Fowler & Fowler, 2020; Wilson, Ryan & Evans, 2020). Indeed, as 
anagenesis continues to be identified among fossil lineages, the predominant relative 
frequency of strictly cladogenetic evolutionary models (e.g., punctuated equilibria; 
Eldredge & Gould, 1972) must eventually come under scrutiny. Future explorations of 
evolutionary mode in fossil taxa, including further tests of the hypotheses presented here, 
will be important in this regard, and have the potential to refine understanding of the 
pattern and process of dinosaur evolution.

ACKNOWLEDGEMENTS

Special thanks are given to John Wilson for his discovery of the holotype specimen. Thanks 
also to Elizabeth Freedman Fowler and Matthew Lavin for discussion and guidance that 
improved the quality of this manuscript, and to David Hone, Jared Voris, and Thomas 
Holtz for helpful reviews. Thanks to Joshua Chase, Greg Liggett, Pat Gunderson, and other 
employees of the Bureau of Land Management who assisted DF with land access, 
permitting, and excavation. We would also like to thank the members of BDM field crews 
that worked tirelessly to excavate “Jack’s B2,” including Meara Aberle, Jon Carr, Andrew 
Chappelle, Steven Clawson, Chalfant Conley, Brian Conway, Joshua deOlivera, Jordan 
Drost, Robert Ebelhar, Elizabeth Flint, Elizabeth Freedman Fowler, Joshua Fry, Sandi 
Guarino, Brad Hoole, Felipe Jannarone, Marianna Karagiannis, Ashley Lambert, Rachel 
Livengood, Kat Maguire, Marianna Rogers, Emily Waldman, Solin Wanders, Alyssa 
Wiegars, Jack Wilson, and many others, without whom study of the holotype specimen 
would not have been possible. Thanks to Steven Clawson, Amanda Hendrix, Destiny Wolf, 
and Darrah Steffen for their preparation and curation of the holotype specimen. Thanks 
also to the Bergtohl family for land access, and to Matt Poole and Montana State Office 
Glasgow for camping access.

INSTITUTIONAL ABBREVIATIONS

AMNH  American Museum of Natural History, New York, New York, USA
BDM    Badlands Dinosaur Museum, Dickinson Museum Center, Dickinson, North 
       Dakota, USA
CMN    Canadian Museum of Nature, Ottawa, Ontario, Canada
FMNH   Field Museum of Natural History, Chicago, Illinois, USA
GIN    Palaeontological Centre of the Mongolian Academy of Sciences, Ulaanbaatar, 
       Mongolia
GM     Ganzhou Museum, Ganzhou, China
MOR    Museum of the Rockies, Bozeman, Montana, USA
TMP    Royal Tyrrell Museum of Paleontology, Drumheller, Alberta, Canada
UALVP  University of Alberta Laboratory for Vertebrate Palaeontology, Edmonton, 
       Alberta, Canada
UMNH   Natural History Museum of Utah, Salt Lake City, Utah, USA
ZPAL   Institute of Palaeobiology of the Polish Academy of Sciences, Warsaw, Poland
ADDITIONAL INFORMATION AND DECLARATIONS

Funding
Funding for this project was provided by the City of Dickinson, ND; the US Bureau of Land Management, TC Energy, Conoco-Phillips, TC Energy employees, Destiny and Craig Wolf, Google employees. 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:
City of Dickinson, ND.
The US Bureau of Land Management.
TC Energy.
Conoco-Phillips.
TC Energy Employees.
Destiny and Craig Wolf.
Google Employees.

Competing Interests
The authors declare they have no competing interests.

Author Contributions
• Elias A. Warshaw conceived and designed the experiments, performed the experiments, analyzed the data, prepared figures and/or tables, authored or reviewed drafts of the article, and approved the final draft.
• Denver W. Fowler conceived and designed the experiments, prepared figures and/or tables, authored or reviewed drafts of the article, and approved the final draft.

Data Availability
The following information was supplied regarding data availability:
The holotype specimen (BDM 107) is stored in the collections of the Badlands Dinosaur Museum in Dickinson, North Dakota.
The matrix used in the cladistic analysis is available as a Supplemental File.

New Species Registration
The following information was supplied regarding the registration of a newly described species:
Daspletosaurus wilsoni sp. nov. species LSID: urn:lsid:zoobank.org:act:790497D0-3664-4F83-BB13-67CDB27B5428.
Publication LSID: urn:lsid:zoobank.org:pub:F7EE2619-89FC-4D72-93DA-EFE6BD549A77.
Supplemental Information
Supplemental information for this article can be found online at http://dx.doi.org/10.7717/peerj.14461#supplemental-information.

REFERENCES

Brochu CA. 2003. Osteology of *Tyrannosaurus rex*: insights from a nearly complete skeleton and high-resolution computed tomographic analysis of the skull. *Journal of Vertebrate Paleontology* 22(Sup4):1–138 DOI 10.1080/02724634.2003.10010947.

Brown CM, Currie PJ, Therrien F. 2022. Intraspecific facial bite marks in tyrannosaurs provide insight into sexual maturity and evolution of bird-like intersexual display. *Paleobiology* 48(1):12–43 DOI 10.1017/pab.2021.29.

Brusatte SL, Carr TD. 2016. The phylogeny and evolutionary history of tyrannosaurid dinosaurs. *Scientific Reports* 6(1):1–8 DOI 10.1038/srep20252.

Brusatte SL, Carr TD, Norell MA. 2012. The osteology of *Alioramus*, a gracile and long-snouted tyrannosaurid (Dinosauria: Theropoda) from the Late Cretaceous of Mongolia. *Bulletin of the American Museum of Natural History* 2012(366):1–197 DOI 10.1206/770.1.

Brusatte SL, Norell MA, Carr TD, Erickson GM, Hutchinson JR, Balanoff AM, Bever GS, Choiniere J, Makovicky PJ, Xu X. 2010. Tyrannosaur paleobiology: new research on ancient exemplar organisms. *Science* 329(5998):1481–1485 DOI 10.1126/science.1193304.

Carpenter K. 1990. Variation in *Tyrannosaurus rex*. In: Carpenter K, Currie PJ, eds. *Dinosaur Systematics: Perspectives and Approaches*. Cambridge: Cambridge University Press, 141–145.

Carr TD. 1999. Craniofacial ontogeny in tyrannosauridae (Dinosauria, Coelurosauria). *Journal of Vertebrate Paleontology* 19(3):497–520 DOI 10.1080/02724634.1999.10011161.

Carr TD. 2020. A high-resolution growth series of *Tyrannosaurus rex* obtained from multiple lines of evidence. *PeerJ* 8:e9192 DOI 10.7717/peerj.9192.

Carr TD, Varricchio DJ, Sedlmayr JC, Roberts EM, Moore JR. 2017. A new tyrannosaur with evidence for anagenesis and crocodile-like facial sensory system. *Scientific Reports* 7(1):1–11 DOI 10.1038/srep44942.

Carr TD, Williamson TE. 2004. Diversity of late Maastrichtian Tyrannosauridae (Dinosauria: Theropoda) from western North America. *Zoological Journal of the Linnean Society* 142(4):479–523 DOI 10.1111/j.1096-3642.2004.00130.x.

Carr TD, Williamson TE. 2010. *Bistahieversor sealeyi*, gen. et sp. nov., a new tyrannosaurid from New Mexico and the origin of deep snouts in Tyrannosauroidea. *Journal of Vertebrate Paleontology* 30(1):1–16 DOI 10.1080/02724630903413032.

Carr TD, Williamson TE, Britt BB, Stadtman K. 2011. Evidence for high taxonomic and morphologic tyrannosaurid diversity in the Late Cretaceous (Late Campanian) of the American Southwest and a new short-skulled tyrannosaurid from the Kaiparowits formation of Utah. *Naturwissenschaften* 98(3):241–246 DOI 10.1007/s00114-011-0762-7.

Carr TD, Williamson TE, Schwimmer DR. 2005. A new genus and species of tyrannosaurid from the Late Cretaceous (Middle Campanian) Demopolis Formation of Alabama. *Journal of Vertebrate Paleontology* 25(1):119–143 DOI 10.1671/0272-4634(2005)025[0119:ANGASO]2.0.CO;2.

Currie PJ. 2003. Cranial anatomy of tyrannosaurid dinosaurs from the Late Cretaceous of Alberta, Canada. *Acta Palaeontologica Polonica* 48(2):191–226 DOI 10.1017/CBO9780511608377.023.

Eberth DA, Currie PJ, Koppelhus EB. 2005. The geology. Vol. 3. Bloomington: Indiana University Press, 54–82.
Eldredge N, Gould SJ. 1972. Punctuated equilibria: an alternative to phyletic gradualism. In: Schopf TJM, ed. Models in Paleobiology. San Francisco: Freeman Cooper & Co., 82–115.

Fiorillo AR, Tykoski RS. 2014. A diminutive new tyrannosaur from the top of the world. PLOS ONE 9(3):e91287 DOI 10.1371/journal.pone.0091287.

Foster W, Brusatte SL, Carr TD, Williamson TE, Yi L, Lü J. 2021. The cranial anatomy of the long-snouted tyrannosaurid dinosaur *Qianzhousaurus sinensis* from the Upper Cretaceous of China. Journal of Vertebrate Paleontology 41(4):e1999251 DOI 10.1080/02724634.2021.1999251.

Fowler DW. 2017. Revised geochronology, correlation, and dinosaur stratigraphic ranges of the Santonian-Maastrichtian (Late Cretaceous) formations of the Western Interior of North America. PLOS ONE 12(11):e0188426 DOI 10.1371/journal.pone.0188426.

Fowler DW, Fowler EAF. 2020. Transitional evolutionary forms in chasmosaurine ceratopsid dinosaurs: evidence from the Campanian of New Mexico. PeerJ 8:e9251 DOI 10.7717/peerj.9251.

Freedman Fowler EA, Horner JR. 2015. A new brachylophosaurin hadrosaur (Dinosauria: Ornithischia) with an intermediate nasal crest from the Campanian Judith River Formation of northcentral Montana. PLOS ONE 10(11):e0141304 DOI 10.1371/journal.pone.0141304.

Goloboff PA, Farris JS, Nixon KC. 2008. TNT, a free program for phylogenetic analysis. Cladistics 24(5):774–786 DOI 10.1111/j.1096-0031.2008.00217.x.

Hone DW, Wang K, Sullivan C, Zhao X, Chen S, Li D, Ji S, Ji Q, Xu X. 2011. A new, large tyrannosaurine theropod from the Upper Cretaceous of China. Cretaceous Research 32(4):495–503 DOI 10.1016/j.cretres.2011.03.005.

Horner JR, Varricchio DJ, Goodwin MB. 1992. Marine transgressions and the evolution of Cretaceous dinosaurs. Nature 358(6381):59–61 DOI 10.1038/358059a0.

Hurum JH, Sabath K. 2003. Giant theropod dinosaurs from Asia and North America: skulls of *Tarbosaurus bataar* and *Tyrannosaurus rex* compared. Acta Palaeontologica Polonica 48(2):161–190.

Kauffman EG. 1977. Geological and biological overview: Western Interior Cretaceous basin: The Mountain Geologist. Vol. 14, 75–99.

Loewen MA, Irmis RB, Sertich JJ, Currie PJ, Sampson SD. 2013. Tyrant dinosaur evolution tracks the rise and fall of Late Cretaceous oceans. PLOS ONE 8(11):e79420 DOI 10.1371/journal.pone.0079420.

Lü J, Yi L, Brusatte SL, Yang L, Li H, Chen L. 2014. A new clade of Asian Late Cretaceous long-snouted tyrannosaurids. Nature Communications 5(1):1–10 DOI 10.1038/ncomms4788.

Matthew WD, Brown B. 1922. The family Deinodontidae, with notice of a new genus from the Cretaceous of Alberta. Bulletin of the American Museum of Natural History 46(6):367–385 DOI 10.5281/ZENODO.1048848.

Matthew WD, Brown B. 1923. Preliminary notices of skeletons and skulls of Deinodontidae from the Cretaceous of Alberta. American Museum Novitates 89:1–10 DOI 10.5281/zenodo.4715537.

McDonald AT, Wolfe DG, Dooley AC Jr. 2018. A new tyrannosaurid (Dinosauria: Theropoda) from the Upper Cretaceous Menefee Formation of New Mexico. PeerJ 6:e5749 DOI 10.7717/peerj.5749.

Molnar RE. 1991. The cranial morphology of *Tyrannosaurus rex*. Palaeontographica. Abteilung A, Paläozoologie, Stratigraphie 217:137–176.

Ogg JG, Hinnov LA. 2012. Cretaceous. In: Gradstein FM, Ogg JG, Schmitz MD, Ogg G, eds. The Geologic Time Scale. Vol. 2. Oxford, UK: Elsevier, 793–853.
Osborn HF. 1905. Article XIV. Tyrannosaurus and other Cretaceous dinosaurs. *Proceedings of the Academy of Natural Sciences of Philadelphia* 8:72.

Osborn HF. 1912. Crania of *Tyrannosaurus* and *Allosaurus*; integument of the iguanodont dinosaur *Trachodon*. Memoirs of the AMNH; new ser., v. 1, pt. 1-2. Available at [http://hdl.handle.net/2246/49](http://hdl.handle.net/2246/49).

Paul GS, Persons WS, Van Raalte J. 2022. The Tyrant Lizard King, Queen and Emperor: Multiple Lines of Morphological and Stratigraphic Evidence Support Subtle Evolution and Probable Speciation Within the North American Genus *Tyrannosaurus*. *Evolutionary Biology* 49(2):156–179 DOI 10.1007/s11692-022-09561-5.

Paulina Carabajal A, Currie PJ, Dudgeon TW, Larsson HC, Miyashita T. 2021. Two braincases of *Daspletosaurus* (Theropoda: Tyrannosauridae): anatomy and comparison¹. *Canadian Journal of Earth Sciences* 58(9):885–910 DOI 10.1139/cjes-2020-0185.

Rogers RR, Kidwell SM, Deino AL, Mitchell JP, Nelson K, Thole JT. 2016. Age, correlation, and lithostratigraphic revision of the Upper Cretaceous (Campanian) Judith River Formation in its type area (north-central Montana), with a comparison of low-and high-accommodation alluvial records. *The Journal of Geology* 124(1):99–135 DOI 10.1086/684289.

Russell DA. 1970. *Tyrannosaurs from the Late Cretaceous of western Canada*. Ottawa: National Museum of Natural Sciences, Publications in Palaeontology, No. 1.

Scannella JB, Fowler DW, Goodwin MB, Horner JR. 2014. Evolutionary trends in Triceratops from the Hell Creek formation. *Montana Proceedings of the National Academy of Sciences of the United States of America* 111(28):10245–10250 DOI 10.1073/pnas.1313334111.

Sereno PC, McAllister S, Brusatte SL. 2005. TaxonSearch: a relational database for suprageneric taxa and phylogenetic definitions. *PhyloInformatics* 8(56):1–25 DOI 10.5281/zenodo.59797.

Soltis PS, Soltis DE. 2003. Applying the bootstrap in phylogeny reconstruction. *Statistical Science* 18(2):256–267 DOI 10.1214/ss/1063994980.

Szalay FS. 1977. Ancestors, descendants, sister groups and testing of phylogenetic hypotheses. *Systematic Biology* 26(1):12–18 DOI 10.1093/sysbio/26.1.12.

Voris JT, Therrien F, Zelenitsky DK, Brown CM. 2020. A new tyrannosauarine (Theropoda: Tyrannosauridae) from the Campanian Foremost Formation of Alberta, Canada, provides insight into the evolution and biogeography of tyrannosaurids. *Cretaceous Research* 110(41):104388 DOI 10.1016/j.cretres.2020.104388.

Voris JT, Zelenitsky DK, Therrien F, Currie PJ. 2019. Reassessment of a juvenile *Daspletosaurus* from the Late Cretaceous of Alberta, Canada with implications for the identification of immature tyrannosaurids. *Scientific Reports* 9(1):1–10 DOI 10.1038/s41598-019-53591-7.

Voris JT, Zelenitsky DK, Therrien F, Ridgely RC, Currie PJ, Witmer LM. 2022. Two exceptionally preserved juvenile specimens of *Gorgosaurus* libratus (Tyrannosauridae, Albertosaurinae) provide new insight into the timing of ontogenetic changes in tyrannosaurids. *Journal of Vertebrate Paleontology* 41(6):e2041651 DOI 10.1080/02724634.2021.2041651.

Wagner PJ, Erwin DH, Anstey RL. 1995. Phylogenetic patterns as tests of speciation models. In: *New Approaches to Speciation in the Fossil Record*. New York: Columbia University Press, 87–122.

Wilson JP, Ryan MJ, Evans DC. 2020. A new, transitional centrosaurine ceratopsid from the Upper Cretaceous Two Medicine Formation of Montana and the evolution of the ‘Styracosaurus-line’ dinosaurs. *Royal Society Open Science* 7(4):200284 DOI 10.1098/rsos.200284.

Zietlow AR. 2020. Craniofacial ontogeny in Tylosaurinae. *PeerJ* 8(1):e10145 DOI 10.7717/peerj.10145.