

## **The fragile legacy of *Amphicoelias fragillimus* (Dinosauria: Sauropoda; Morrison Formation - Latest Jurassic)**

In the summer of 1878, American paleontologist Edward Drinker Cope published the discovery of a sauropod dinosaur that he named *Amphicoelias fragillimus*. What distinguishes *A. fragillimus* in the annals of paleontology is the immense magnitude of the skeletal material. The single incomplete dorsal vertebra as reported by Cope was a meter and a half in height, which when fully reconstructed, would make *A. fragillimus* the largest vertebrate ever. After this initial description Cope never mentioned *A. fragillimus* in any of his scientific works for the remainder of his life. More than four decades after its description, a scientific survey at the American Museum of Natural History dedicated to the sauropods collected by Cope failed to locate the remains or whereabouts of *A. fragillimus*. For nearly a century the remains have yet to resurface. The enormous size of the specimen has generally been accepted despite being well beyond the size of even the largest sauropods known from verifiable fossil material (e.g. *Argentinosaurus*). By deciphering the ontogenetic change of *Diplodocoidea* vertebrae, the science of gigantism, and Cope's own mannerisms, we conclude that the reported size of *A. fragillimus* is most likely an extreme over-estimation.

## Introductory Note:

The paper below, “The fragile legacy of *Amphicoelias fragillimus* (Dinosauria: Sauropoda; Morrison Formation - Latest Jurassic)”, was released on December 15<sup>th</sup>, 2014 in the second issue of the open access online journal *Volumina Jurassica*. Unfortunately, the version released was not our intended final version. This mistake was due to a miscommunication on our part during the manuscript revision stage. All of the information pertaining to our assessment that the values reported by E.D. Cope represent typographical errors remains the same in this released version. The striking difference between the two versions pertains to the taxonomic stability and proposed future treatment of the genus *Amphicoelias*. While the differences between these two versions may be minor, we feel that the version included below is the more proper of the two. Our intention with this version, released via PeerJ, is to serve as a supplemental or supporting file to the *Volumina Jurassica* version. Citations of this manuscript should still follow that of *Volumina Jurassica*. We greatly appreciate all of the help and assistance from the editors and staff of *Volumina Jurassica*, and we ask forgiveness from the journal and the readership for this discrepancy.

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- Since the release of the paper, a few additional points have been raised. Specifically, it was suggested to us that it is possible to crosscheck Cope's published measurements by comparing them with the original print size (1:10) of the figured *A. fragillimus* vertebra, which suggests a smaller vertebral size (yet still close to our calculated 1.83 M). We felt that due to the already speculative nature of the specimen, it was more conservative to propose that the numbers published in the text by Cope were legitimate, just that their specific order was not. However, we fully acknowledge that further typographical errors could exist. Regardless of the specifics, such questioning is wonderfully stimulating to the discussion of the plausibility of *A. fragillimus*.

## The fragile legacy of *Amphicoelias fragillimus* (Dinosauria: Sauropoda; Morrison Formation - Latest Jurassic)

D. Cary Woodruff<sup>1,2\*</sup>, John R. Foster<sup>3</sup>

<sup>1</sup> Museum of the Rockies, <sup>2</sup> Department of Earth Sciences, Montana State University, Bozeman, MT, USA <sup>3</sup> Museum of Moab, Moab, UT, USA

\*sauropod4@gmail.com

**Abstract.** In the summer of 1878, American paleontologist Edward Drinker Cope published the discovery of a sauropod dinosaur that he named *Amphicoelias fragillimus*. What distinguishes *A. fragillimus* in the annals of paleontology is the immense magnitude of the skeletal material. The single incomplete dorsal vertebra as reported by Cope was a meter and a half in height, which when fully reconstructed, would make *A. fragillimus* the largest vertebrate ever. After this initial description Cope never mentioned *A. fragillimus* in any of his scientific works for the remainder of his life. More than four decades after its description, a scientific survey at the American Museum of Natural History dedicated to the sauropods collected by Cope failed to locate the remains or whereabouts of *A. fragillimus*. For nearly a century the remains have yet to resurface. The enormous size of the specimen has generally been accepted despite being well beyond the size of even the largest sauropods known from verifiable fossil material (e.g. *Argentinosaurus*). By deciphering the ontogenetic change of Diplodocoidea vertebrae, the science of gigantism, and Cope's own mannerisms, we conclude that the reported size of *A. fragillimus* is most likely an extreme over-estimation.

**Key words:** *Amphicoelias fragillimus*; E. D. Cope; sauropod; gigantism

**Institutional Abbreviations:** AMNH – American Museum of Natural History, New York, NY, U.S.A.; CM – Carnegie Museum of Natural History, Pittsburgh, PA, U.S.A.; MOR – Museum of the Rockies, Bozeman, MT, U.S.A.; SMA – Sauriermuseum Aathal, Aathal, Switzerland; WDC – Wyoming Dinosaur Center, Thermopolis, WY, U.S.A.

### INTRODUCTION

Described by E. D. Cope in 1878, the holotype (and only) specimen of *A. fragillimus* consisted only of a distal end of a femur and a partial posterior dorsal neural arch and spine. What makes *A. fragillimus* truly unique in all of dinosaurian paleontology is the reported immense size of the material. As reported by Cope, the fragmentary posterior dorsal vertebra of *A. fragillimus* was 1.5 meters tall, which when reconstructed resulted in the complete dorsal

vertebra being at least 1.83 meters tall. Later studies based general proportions of *A. fragillimus* on those of *Diplodocus*, resulting in body length estimates of 58-60 meters (Paul, 1994; Carpenter, 2006). Since Cope's report of *A. fragillimus*, all other diplodocoids, sauropodomorphs, and all other dinosaurs pale in comparative size; thus *A. fragillimus* could be the largest dinosaur, and largest vertebrate in Earth's history (the Blue Whale being approximately 29 meters long [Reilly *et al.*, 2008], Fig. 1).

Surviving correspondence and journal entries between Cope and the collector O. Lucas address the presence and quarry location of *A. fragillimus* (McIntosh, 1998; Monaco, 1998; Carpenter, 2006) prior to Cope's 1878 formal description. However, at some point between Cope's 1878 description and Osborn and Mook's monograph on the sauropods of Cope in 1921, the *A. fragillimus* holotype material disappeared. This material was assigned an American Museum of Natural History collections number when the AMNH acquired the Cope collection in 1895 (AMNH 5777; McIntosh, 1998), so the specimen apparently survived at least until then. In the original description Cope noted the extreme fragile and delicate nature of the material (hence the species name *fragillimus*), and since fossil preservatives were not used at the time, it has been surmised that at some point prior to Osborn and Mook's survey that the deteriorating material was discarded, potentially even by Cope himself (Carpenter, 2006). Regardless of the whereabouts of the holotype material, several works have accepted Cope's measurements without question (Osborn and Mook, 1921; McIntosh, 1998; Carpenter, 2006).

Unfortunately in many respects the study of *A. fragillimus* represents a forever untestable scientific endeavor. Due to the disappearance of the holotype material, proportions cannot be re-measured, morphology and anatomy cannot be examined, and phylogeny and taxonomy are practically non-existent; in essence any further science is hindered. All we are left with is the scant information in Cope's 1878 description. Yet, based on what information survives, presumed body length and body mass have been calculated (Paul, 1994; Mazzetta *et al.*, 2004; Carpenter, 2006). This resulting body size has remained largely unquestioned, and dinosaur paleontology has been largely indoctrinated with the concept that *A. fragillimus* was possibly the largest dinosaur; although it is rarely mentioned in these discussions due to the

missing status of the type material. While any scientific assessment of *A. fragillimus* is conjectural, it is nonetheless important to question the biology of such a reported organism. If reconstructed elemental size and body length and mass can all be hypothesized, the same should hold true for other biological aspects. Through morphological comparisons this analysis shall examine more plausible elemental reconstructions while also considering biological and historical aspects in order to determine if a sauropod of the reported size of *A. fragillimus* could have existed.

## DISCUSSION

While the attention of *A. fragillimus* works are generally dedicated to length and weight of the animal (Paul, 1994; Mazzetta *et al.*, 2004), this analysis shall attempt to determine a more accurate reconstruction of the *A. fragillimus* material. Cope's original reconstructive comparisons were based on *Camarasaurus* and the second *Amphicoelias* species *A. altus* (AMNH 5764). However, based on vertebral morphological similarities, modern analyses prefer to use a diplodocid proxy, such as *Diplodocus* (Paul, 1994). Phylogenetically, both Rauhut *et al.* (2005) and Whitlock (2011) recover the second *Amphicoelias* species, *A. altus*, as a basal diplodocoid. Thus a diplodocid proxy is morphologically and phylogenetically supported. However, as Carpenter (2006) pointed out, if *A. fragillimus* was indeed a diplodocid, then either *Apatosaurus*, *Diplodocus*, or *Barosaurus* are suitable proxies. Indeed, it must be stated first and foremost that differing proxies would produce drastically different outcomes (i.e. the proportional differences between *Dicraeosaurus*, *Diplodocus*, and *Brachiosaurus*). In addition, all analyses presume that supergiant sauropods had proportions respective to their smaller relatives; for the purposes of this analysis, proportions and morphology shall be compared to *Diplodocus*, *Supersaurus*, and *A. altus*. The largest *Diplodocus* species, *D. hallorum*, has been temporarily excluded from this analysis since a large portion of the dorsal series remains articulated in a matrix block (Lucas *et al.*, 2006), and therefore comparable vertebral features and orientations cannot be examined.

### A. FRAGILLIMUS ELEMENTAL RECONSTRUCTIONS

## VERTEBRAL RECONSTRUCTION

In his 1878 description, Cope presumed that the neural arch of *A. fragillimus* represented the tenth dorsal, and he reconstructed the centrum similar to that of *A. altus*. Cope's *A. fragillimus* is reconstructed as having a very narrow and rectangular centrum (nearly twice as tall as wide) with a very small and rounded neural spine apex shortly above the preserved portion (Fig. 2). This reconstruction results in a posterior dorsal vertebra 1.83 meters tall. In 2006, based on presumed morphological similarities, K. Carpenter reconstructed the *A. fragillimus* vertebra as an extensively modified *A. altus* posterior dorsal vertebra. Carpenter's reconstruction resulted in a centrum that was slightly more circular (yet still taller than wide), with a very tall and narrow neural arch, slightly vertically oriented transverse processes, and a larger, more bulbous neural spine apex. This reconstruction, at least being based on a comparative specimen, results in the *A. fragillimus* posterior dorsal vertebra as being 2.7 meters tall (Fig. 2). Looking at a posterior dorsal from *Diplodocus carnegii* (D 10 from CM 84; Hatcher, 1901), the centrum is very circular, with the height and width being near equal. From the neural canal to the hyposphene, the neural arch is rather tall, but becomes quite narrow proximally to the neural canal. Proportionally, in the case of *D. carnegii* (D 10 from CM 84) the centrum is approximately 1/4 of the entire vertebral height. In a larger diplodocid, such as *Supersaurus vivianae* (WDC DMJ-021), the centrum is significantly larger than that of *D. carnegii*. In *S. vivianae* the centrum is much wider than it is tall. The neural arch from the neural canal to the hyposphene is much shorter than in *D. carnegii*, yet still retains the narrow portion proximal to the neural canal. Also, proportionally the centrum is a much larger component of the total vertebral height; in this case approximately 1/3 of the entire height. If these size and proportional differences between *D. carnegii* and *S. vivianae* are indicative of vertebral changes within large diplodocids, then it might be possible to predict the morphologies of even larger diplodocids. Since we do not have any material from a diplodocid between the size of *S. vivianae* and *A. fragillimus*, it is impossible to determine whether or not the centrum continued to widen and dominate more of the total vertebral height. However, assuming that the *A. fragillimus* posterior dorsal was

indeed roughly similar in relative proportions to that of *S. viviana*, this new reconstruction results in a posterior dorsal larger than Carpenter's (2006).

The relationship between the hyposphene to neural canal height versus the width and height of the centrum in *S. viviana* (WDC DMJ-021) results in the *A. fragillimus* centrum being rather circular, approximately 95 cm across by 80 cm tall. The neural arch was reconstructed as being rather short, as in the case of *S. viviana* (WDC DMJ-021), with a thinning, narrow portion proximal to the neural canal. In immature Diplodocidae the lateral sides of the neural arch are tall and straight, and throughout ontogeny the arch shortens in height and narrows lateral to the neural canal (Woodruff and Fowler, 2012). Based on the reported size, we presume that *A. fragillimus* was a fully mature animal, and thus would follow the same trend. The transverse processes were reconstructed as being slightly more horizontally oriented than in Carpenter's (2006) reconstruction, which is more akin to those observed in *D. carnegii* (CM 84; the transverse processes of WDC DMJ-021 are not preserved). The neural spine apex is nearly the same height, but much more bulbous (potentially the continuation of an ontogenetic trend observed by Woodruff and Fowler, 2012; Fig. 3). Using the ratio between the centrum height to the overall vertebral height of *S. viviana* (WDC DMJ-021), results in a total *A. fragillimus* vertebral height of 2.8 meters (Fig. 2).

#### FEMORAL RECONSTRUCTION

With regard to the femur of *A. fragillimus*, any attempted reconstruction is purely conjectural. Yet femoral dimensions can provide critical variables for life history information, such as body mass and growth rates. Since the femur can reveal such insightful information about the organism in question, we feel that an attempt to hypothesize the femoral morphology of *A. fragillimus* is a worthwhile endeavor.

In his 1878 description, Cope did mention that a distal end of a femur was located near the dorsal neural arch, and assumedly based on its large size from the same animal. However, Cope never figured or described this element in any detail. For comparative and speculative purposes, Cope noted that in the case of *A. altus* and *Camarasaurus supremus* the femora



were roughly twice as tall as the tallest dorsal vertebra (Cope, 1878a). Assuming that the preserved neural arch was indeed from the tallest of the series, this results in a femur over 3.6 meters tall (Cope, 1878a). In using *Diplodocus* proportions, the femur to dorsal 10 ratio is approximately 1.6-1.7. This produces an *A. fragillimus* femur anywhere from 3.1-4.76 meters tall (Paul, 1994; Carpenter, 2006; this analysis).

Assuming that *A. fragillimus* and *A. altus* were proportioned similar with respect to femur morphology, then the femur of *A. fragillimus* should be quite gracile and extremely long and narrow (*A. altus* displays the “stovepipe” diaphyseal morphology). It would seem rather peculiar for an organism with a potential mass of 122,400 kg (Carpenter, 2006) to be supported by such a proportionally narrow and gracile femur as opposed to a stocky and robust *Apatosaurus*-like femur. But, as the femur increases in length, so should the relative width. Following the development through *Diplodocus* femoral ontogeny, some general trends include the femoral head enlarging and inclining progressively horizontal, the greater trochanter protruding much more medially, the diaphysis thickening appositionally, and the tibial and fibular condyles becoming much larger and more robust (Fig. 4). Assuming the same developmental trends hold true (disregarding the questionable diaphyseal morphology mentioned above) in *A. fragillimus*, the femur of *A. altus* (AMNH 5764) has been modified in the same manner to reconstruct that of *A. fragillimus*. The resulting reconstructed femur is 4.76 meters tall (Fig. 5).

## THE SCIENTIFIC PLAUSIBILITY OF *A. FRAGILLIMUS*' MASSIVE SIZE

### ECOLOGY

The gargantuan size of *A. fragillimus* and the mystery surrounding the disappearance of the material makes this organism truly unique. That the purported largest terrestrial vertebrate of all time was found and lost before the advent of modern documentation technology is a travesty; and, alas, the memory of *A. fragillimus* may forever remain engrained in the annals of paleontology. While the evidence exists to verify that Cope did indeed extract the remains of some large sauropod from Colorado, the science of gigantism should dispel the potential myth of *A. fragillimus*' purported reconstructed length and body



mass. Intrinsic factors such as gravitational forces indicate the maximum weight limit for a terrestrial organism to be 75,000 kg (Günther *et al.*; 2002), while bone strength and muscles forces have been calculated to max out in terrestrial body sizes over 90,718 kg (Hokkanen, 1986). With regard to large body size, the most critical extrinsic property is resource availability. The larger an organism the more nutritional resource it needs (particularly if it has a higher basal metabolic rate) along with a larger home range for said resources (Burness *et al.*, 2001). If an organism is too large for the geography and biota to support it, that particular organism will be in direct conflict with the ecosystem and shall eventually go extinct. According to MacArthur and Wilson (1967), the larger a particular land area the more individuals per taxon will be present.

If a homogenized Morrison Formation sauropod biota were indeed correct, this means that at least ten genera and potentially twenty-one species with body masses ranging from 7-26 tons were all occupying the same landscape (Seebacher, 2001; Foster, 2003; Mazzetta *et al.*, 2004; Taylor, 2009). While there is evidence to suggest generic and ontogenetic niche partitioning (Whitlock *et al.*, 2010; Fowler and Sullivan, 2011), these sauropod genera would undoubtedly have had an impact on the plant biota; so, presumably a sauropod an entire order of magnitude greater in mass would have needed significantly more resources and would have potentially been in direct competition with other co-existing large herbivores, especially as juveniles and sub-adults. As in the case of the African savanna, the abundance of the largest herbivores is much lower than that of the smaller herbivores (Owen-Smith and Mills, 2008). Assuming the same would hold true for Morrison time, this could explain the over abundance of the relatively smaller *Camarasaurus*, compared to the rare giant dipodocids such as *Supersaurus* and *Diplodocus hallorum*.

## TAPHONOMY AND STRATIGRAPHY

Taphonomically the larger an organism, the more sediment that is needed to entomb the carcass. When one considers the immense size of the giant sauropods such as *Puertasaurus reuili* (Novas *et al.*, 2005), *Turiasaurus riodevensis* (Royo-Torres *et al.*, 2006), *Paralititan stromeri* (Smith *et al.*, 2001), *Diplodocus hallorum* (Lucas *et al.*, 2006),

*Futalognkosaurus dukei* (Calvo *et al.*, 2007), *Sauroposeidon proteles* (Wedel *et al.*, 2000), *Supersaurus viviana* (Jensen, 1985), *Alamosaurus sanjuanensis* (Gilmore, 1922), *Argentinosaurus huinculensis* (Bonaparte and Coria, 1993), and *Bruhathkayosaurus matleyi* (Yadagiri and Ayyasami, 1989), all of these sauropods are generally known from relatively sparse remains. So, one could make the argument that if *A. fragillimus* was considerably larger than any yet-described sauropod, this could support the taphonomic plausibility of a burial bias.

Stratigraphically the *A. fragillimus* (and *A. altus*) quarry sits within the Tithonian portion of the upper Brushy Basin Member of the Morrison Formation (Turner and Peterson, 1999). Likewise, the largest *Camarasaurus* (*C. supremus*), *Apatosaurus* (*A. louisae*), and *Diplodocus* (*D. hallorum*) species are all stratigraphically from the slightly lower Kimmeridgian portion of the upper Brushy Basin (Turner and Peterson, 1999). Perhaps increasing average body size is a stratigraphic trend in Morrison Formation sauropods; although the stratigraphically highest occurrence of *Apatosaurus*, in the upper meters of the Brushy Basin Member in Arches National Park, Utah (Foster, 2005), is of a size typical for the average adult lower in the formation. So, it is very likely that O. Lucas did indeed find the remains of a large upper Morrison Formation sauropod; and as Cope said in 1878, the remains were "...the largest saurian I have ever seen." Certainly this could be the case given that a posterior dorsal from *A. altus* is nearly the same size as that from *D. carnegii*. Given that most Morrison Formation sauropod quarries are located generally near the division between the Lower and Upper portions of the Brushy Basin Member (Turner and Peterson, 1999), the rarity of these larger species could also be attributed to collecting biases.

However, even though fragmentary, remains of the largest sauropods are regularly recorded (i.e. *Turiasaurus riodevensis* and *Alamosaurus sanjuanensis*; and the recent discovery of a bone bed of several large sauropods from Argentina with reported femoral lengths of nearly 2.4 meters). Considering the time and resources that have been spent exploring the vastness of the Morrison Formation, not a single substantiating piece of an *A. fragillimus* sized sauropod has been reported. Taphonomically at the very least, some partial

fragment should have been quite evident and found in the intervening years. As of this publication no such secondary material is scientifically known.

## TAXONOMY

In terms of the genus level taxonomy, this analysis agrees with Osborn and Mook (1921) and McIntosh (1998) that the second species, “*A. latus*”, is simply a synonym of *A. altus*. The distinct stout femur of “*A. latus*” could possibly be pathologic or individual variation. However, in terms of *A. altus*, this analysis agrees with Foster (2007) that there are not enough characters to separate *A. altus* from the genus *Diplodocus*. In the past, *A. altus* has been considered either likely a large cf. *Diplodocus* (Foster, 2001) or at least of uncertain status (Harris, 2006). Even the previously recognized “autapomorphies”, such as the circular femoral cross section (“stovepipe” diaphysis) in *A. altus* are now recognized to be present in many *Diplodocus* specimens (Carpenter, 2006; Woodruff and Fowler, 2012; R. Wilhite pers. comm.; C. Woodruff and J. Foster pers. obs.), while the proportionally longer ulna to femur (Carpenter, 2006), the taller and wider neural arch (Carpenter, 2006), a taller than wider centrum, and other minor vertebral differences are primarily indicative of immature characteristics (Woodruff and Fowler, 2012).

Yet regardless of its questionable status, *A. altus* has been included in several sauropodomorph phylogenies (initially by Wilson [2002]) before the holotype material has been clearly demonstrated to be distinguishable from *Diplodocus*. Several analyses list no autapomorphies for *A. altus*, despite the listing of such characters for other Morrison sauropod genera. The initial characters of *A. altus* used to support its position as a basal diplodcoid by Wilson (2002) include: (3) dentary with a sharply projecting chin, (7) anterior cervical neural spines bifid, (8) posterior cervical and anterior dorsal neural spines bifid; (10) anterior dorsal neural arches with divided centropostzygapophyseal lamina; (12) anterior caudal neural arches with spinoprezygapophyseal lamina on lateral aspect of the neural spine; (15) pubis with a prominent ambiens process. However, none of these characters are preserved in the holotype material (AMNH 5764), which consists of two dorsal vertebrae, a femur, and several other partial elements. We believe that at least some of these characters are based on a second

specimen referred by Wilson and Smith (1996) to *A. altus*, which was subsequently demonstrated by Woodruff and Fowler (2012) to in fact be an immature *Diplodocus*. The apparent taxonomic uncertainty of *A. altus* could arise from this “chimera” character assignment, and we would stress caution against further inclusion of the “genus” in future phylogenetic analyses until the type material has been more fully analyzed.

This analysis shall go one step further and suggest that if the differences between the comparable elements of *A. altus* and *A. fragillimus* are indeed correct, then this is indicative of an ontogenetic trajectory (comparable to the ontogenetic vertebral changes documented by Woodruff and Fowler [2012]). Thus, the *A. altus* material represents an immature animal, and *A. fragillimus* the more mature form. As *A. altus* was erected in 1887 and *A. fragillimus* in 1878, *A. fragillimus* should be synonymized into *A. altus* (likewise suggested by McIntosh, 1998). Stratigraphically and ontogenetically there is coinciding support for a very large uppermost Brushy Basin, Tithonian-aged diplodocine. Considering the lack of legitimate autapomorphies demonstrated for *A. altus* and given its uncertain distinction from the genus *Diplodocus*, this analysis preliminarily supports the referral of the collective *Amphicoelias* material to the genus and species *Diplodocus altus*. As the genus *Amphicoelias* (Cope, 1877) takes priority over *Diplodocus* (Marsh, 1878), this would mean that the genus level identification should be assigned as *Amphicoelias*, and *Diplodocus* would now be a junior subjective synonym. However, in consideration of its historical and cultural significance, coupled with the copious material and significant paleontological understanding of material referred to *Diplodocus*, this analysis calls for *Amphicoelias* being deemed the *nomen oblitum* and *Diplodocus* as a *nomen protectum*.

#### COULD E. D. COPE HAVE BEEN WRONG?

In the face of the historical evidence and Cope’s detailed anatomical training, it would seem hard to deny the existence of *A. fragillimus*. We will never know the answers as to why Cope never published more on *A. fragillimus*, why O. C. Marsh did not publicly question the material, and why the AMNH never publicly expressed vivid interest in displaying the world’s

largest dinosaur. All of these questions are intangible. However, with regard specifically to the described material, it is our belief that the measurements are a typographical error.

## TYPOGRAPHICAL ERROR

In the 1878 description of *A. fragillimus* Cope lists the measurements as follows, “total elevation of the neural arch preserved, 1500 m.; elevation of the posterior zygapophyses, 585; transverse expanse of posterior zygapophyses, 190; vertical diameter of base of diapophysis, 390.” (Cope, 1878a). This list of measurements is interesting because all subsequent works (such as Carpenter, 2006) refer to the units as mm (other 19<sup>th</sup> century works denote meters as “M.” and millimeters as “mm.”). Obviously, Cope did not mean to say that the neural arch was 1,500 meters tall, but immediately in the very first line of given measurements is a typographical error. With regard to the values themselves, some care should be taken as to the literal dimensions. In his 1877 description of *A. altus*, Cope described the femur as being exactly “six feet four inches” (Cope, 1877), while in his table the value given is “1.542 M” (1542 mm; approximately 5 feet; Cope, 1877). While this error does not result in a total re-evaluation of the relative size of the element, it is off by over 38 cm.

This analysis agrees with subsequent work in that the unit “m.” was a typographical error, which was intended to be mm. Yet a 1,500 mm neural arch still leads to the titanic vertebral reconstructions. It is historically unsupported and speculative, but if the value 1,500 mm is altered to 1,050 mm, then this results in a vertebral reconstruction significantly more comparable in size to the largest sauropods in the world and still represents the largest ever found in the Morrison Formation. Since Cope did not indicate where his precise measured reference points were, the newly calculated values are not identical, but they are surprisingly close. Based on Cope’s drawing (which assumes the drawing to be perfectly scaled and an exact 1:1 with the fossil material), with a 1,050 mm neural arch height, the elevation of the posterior zygapophyses is approximately 520 mm; the transverse expanse of posterior zygapophyses is approximately 160 mm; and the vertical diameter of the base of diapophysis is approximately 320 mm. While these values do differ from the originals (aside from the neural arch height the mean difference is 55 mm), proportionally these new values fit much

better with a 1,050 mm neural arch than the originals do with one 1,500 mm. This new arch value still produces a reconstructed total vertebral height nearly 1.83 m, which is close to the value estimated by Cope (Fig. 6).

#### DISAPPEARANCE OF THE *A. FRAGILLIMUS* MATERIAL

Given this new possible explanation, this analysis presents two new potential scenarios as to the disappearance of the *A. fragillimus* material: 1) O. Lucas did collect a large diplodocid partial vertebra, which Cope quickly described. However, the remains were of such poor preservation that Cope discarded the material soon after description (similar to Carpenter's [2006] explanation; and the assigned AMNH collection number probably was assigned after the acquisition of Cope's collection). While this scenario could only be substantiated by a confession, Cope's history is marked with typographical errors (Cope even referred to his own species on occasion as "*fragillimus*" and "*fragillisimus*"). Cope himself claimed credit for the theory that unitatheres possessed trunks and that secretaries and printers had left this information out through a series of errors (Wallace, 1999). And even Cope's longtime friend, H. F. Osborn noted that Cope's own furious rush to describe and publish on fossil material had led to numerous scientific errors (Osborn, 1931); 2) Considering the immense volume of scientific work Cope had compiled for himself, coupled with the fact that his publication finances were significantly limited compared with those of O. C. Marsh, it is possible that Cope deliberately chose not to pursue further in depth works on *A. fragillimus*. It is even possible that Cope himself was never aware of this potential typographical error. And, if O. C. Marsh's spies had reported on the specimen in Colorado, this could be why Marsh never disputed the description. If Cope did not personally dispose of *A. fragillimus*, then it is just as likely that at some point after the AMNH acquired the Cope collection in 1895, a fragmentary, poorly preserved portion of neural arch may have inadvertently slipped "under the radar" of the collections staff and was subsequently disposed of.

#### CONCLUSION

Whichever scenario holds true as to the disappearance of *A. fragillimus*, it is unfortunately most likely that the material was in one way or another destroyed. While this

analysis does not wish to claim that Cope had ulterior motives nor is it believed that Cope purposefully inflated measurement values, some strikingly absent historical events do raise questions. While it is a fact that Cope did not have the near limitless financial or supportive backing of O. C. Marsh, it is curious that Cope never mentioned *A. fragillimus* in any further detail (its discovery and taxonomy were extremely briefly addressed in a publication in 1881 [Cope, 1881]). Some speculations suggest that it was the limited resources that hindered any further works, yet merely mentioning or referencing *A. fragillimus* would not have been a grievous task. In December 1877, Cope described *A. altus* and “*A. latus*” in a review of dinosaurs from Colorado, and another on the same subject in February 1878 (Cope, 1877; 1878b). If *A. fragillimus* was indeed discovered in the summer of 1877, it is curious as to why was it not addressed in either of these sizeable works (the former was released within 6 months before the formal description). The entire formal description of *A. fragillimus* is only two paragraphs long, so certainly it was not omitted from either due to limited text space. Potentially Cope purposefully chose to do a separate description, but given the sparse size of the description this seems rather odd. In addition, it is perplexing as to why Cope did not even mention *A. fragillimus* within his theory of lineages increasing in body size through time (a theory now called Cope’s Rule, [Cope, 1896]). While Cope’s Rule is a loose general trend (and fraught with exceptions, i.e. Hone *et al.*, 2005; Carrano, 2006; Hone *et al.*, 2008; Sander *et al.*, 2011), considering he discovered the largest terrestrial vertebrate at the time, it is striking that Cope did not speculate on *A. fragillimus* and its evolutionary lineage contribution.

To confirm the story and size of *Amphicoelias fragillimus* may be nearly every sauropod paleontologist’s Holy Grail. When in the heart of Morrison Formation country one cannot help but think that the long lost *A. fragillimus* rests around the next bend. But, as in a mythical treasure hunt, the spoils may forever remain unattainable. While the mysteries surrounding *A. fragillimus* are alluring, when we stop to consider the validity of a gargantuan terrestrial organism through several aspects such as biology, ecology, and ontogeny (opposed to making these variables fit around it) the end result is that it is highly unlikely that a terrestrial quadruped of such a purported body size could have existed. Whatever the secrets of *A. fragillimus* are, Cope took them to his grave.



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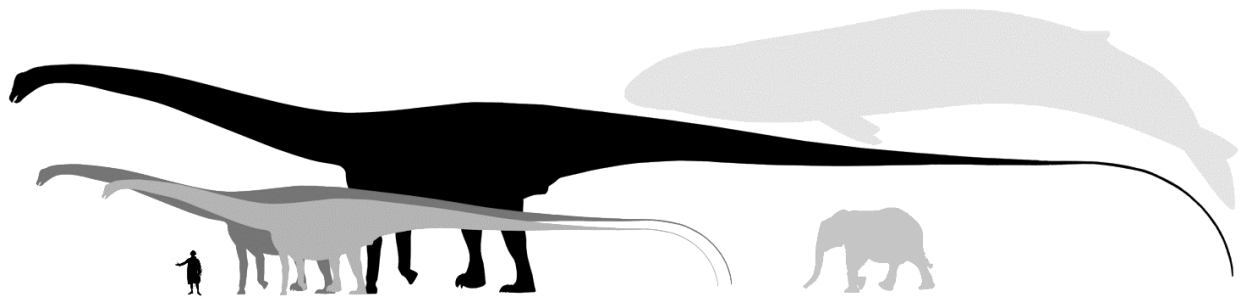


Fig. 1. Scaled life reconstruction of *Amphicoelias fragillimus*; light grey is *Diplodocus carnegii* (CM 84), dark grey is *Supersaurus vivianae* (WDC DMJ-021), black is *A. fragillimus*. Sauropod silhouettes used are *Diplodocus carnegii* available from S. Hartman and PhyloPic. The Blue Whale and African Elephant illustrate dwarfed comparative size (silhouettes from PhyloPic). All to scale. Human scale bar is Gilbert Stuart's "*George Washington*", depicting Washington as his true height of 1.88 M tall.

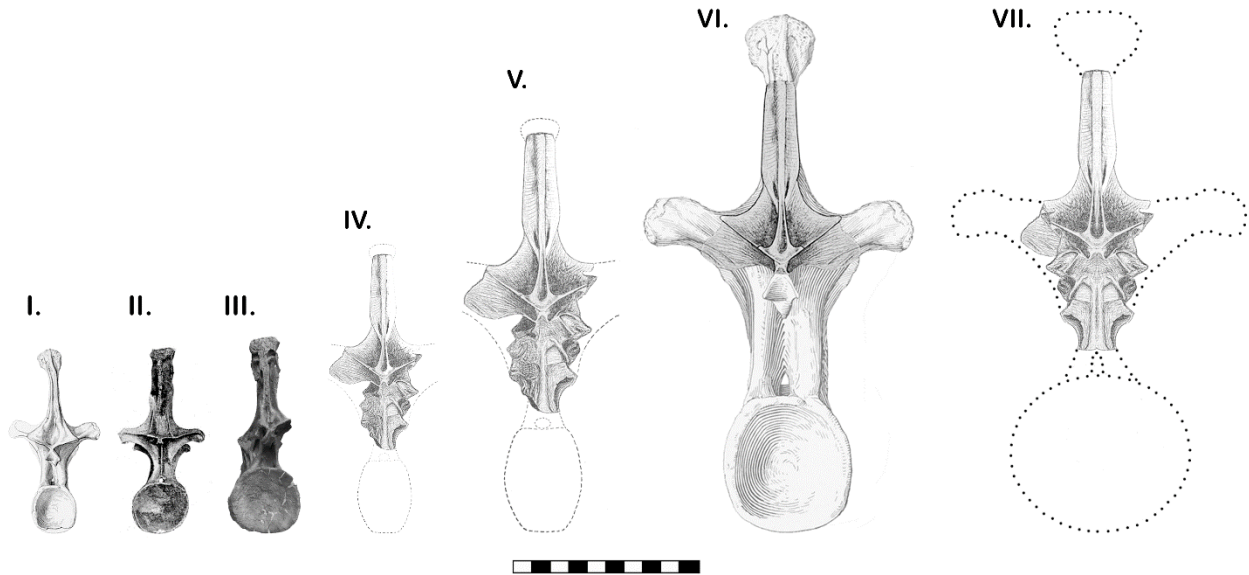


Fig. 2. Diplodocid vertebral comparisons and *Amphicoelias fragillimus* reconstructions. I. *Amphicoelias altus* (from Osborn and Mook, 1921), II. *Diplodocus carnegii* (CM 84, from Hatcher, 1901), III. *Supersaurus vivianae* (WDC DMJ-021), IV. dorsal neural arch of *A. fragillimus* with newly reconstructed neural arch height of 1050 mm (from Cope, 1878a), V. dorsal neural arch of *A. fragillimus* based on measurements described by Cope (from Cope, 1878b), VI. reconstructed dorsal vertebra of *A. fragillimus* modeled after *A. altus* (from Carpenter, 2006), VII. reconstruction of *A. fragillimus* using vertebral trends observed throughout Diplodocidae ontogeny (note the general proportions were modeled assuming *A. fragillimus* maintained similar proportions to that of *S. vivianae*). All vertebrae to scale. Scale bar = 1 M.

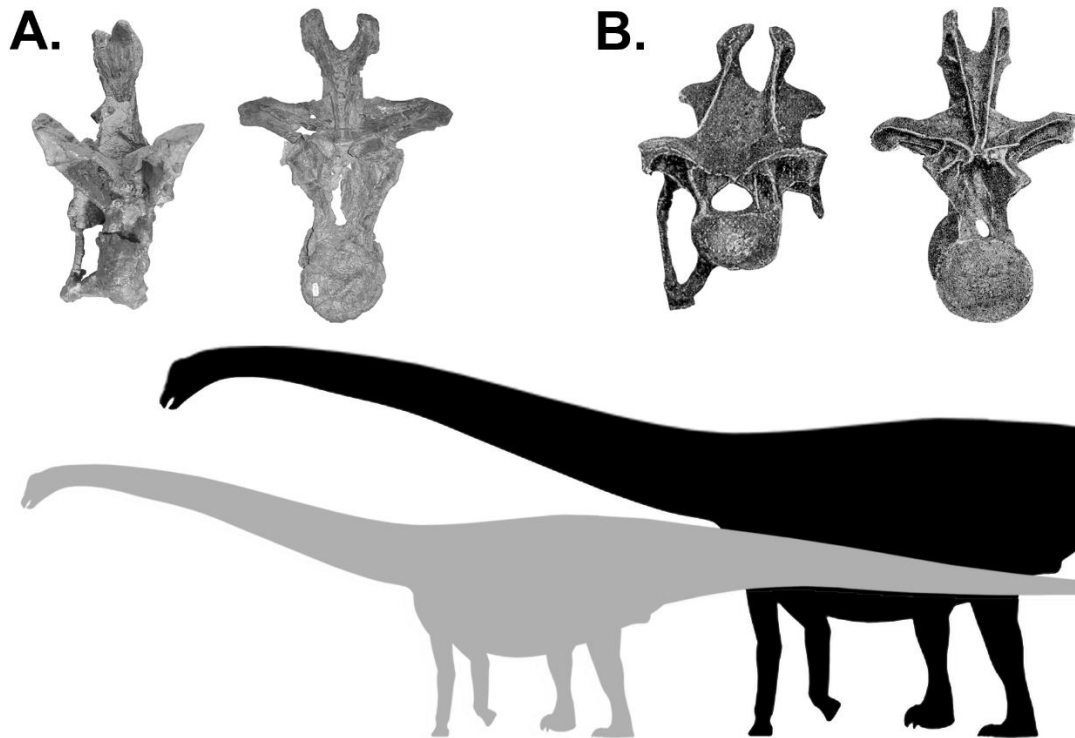


Fig. 3. Representative Diplodocidae (*Diplodocus*) anterior cervical and posterior dorsal vertebral ontogeny. A. and the grey silhouette represent the “sub-adult” *Diplodocus* condition, and B. and the black silhouette the “adult” condition. Silhouettes to relative scale, vertebrae not to scale. Modified from Woodruff and Fowler (2012). Silhouettes from S. Hartman and PhyloPic.



Fig. 4. Ontogenetic development of diplodocid (*Diplodocus*) femora. From left to right: SMA 0009, CM 33976, CM 30762, MOR 790 7-5-95-7, MOR 790 7-23-95-122, MOR 592-35, CM 84. *Diplodocus carnegii* silhouettes reflect the minimum and maximum body lengths of the femoral set (from S. Hartman and PhyloPic). Femora to scale. Scale bar equals 10 cm.

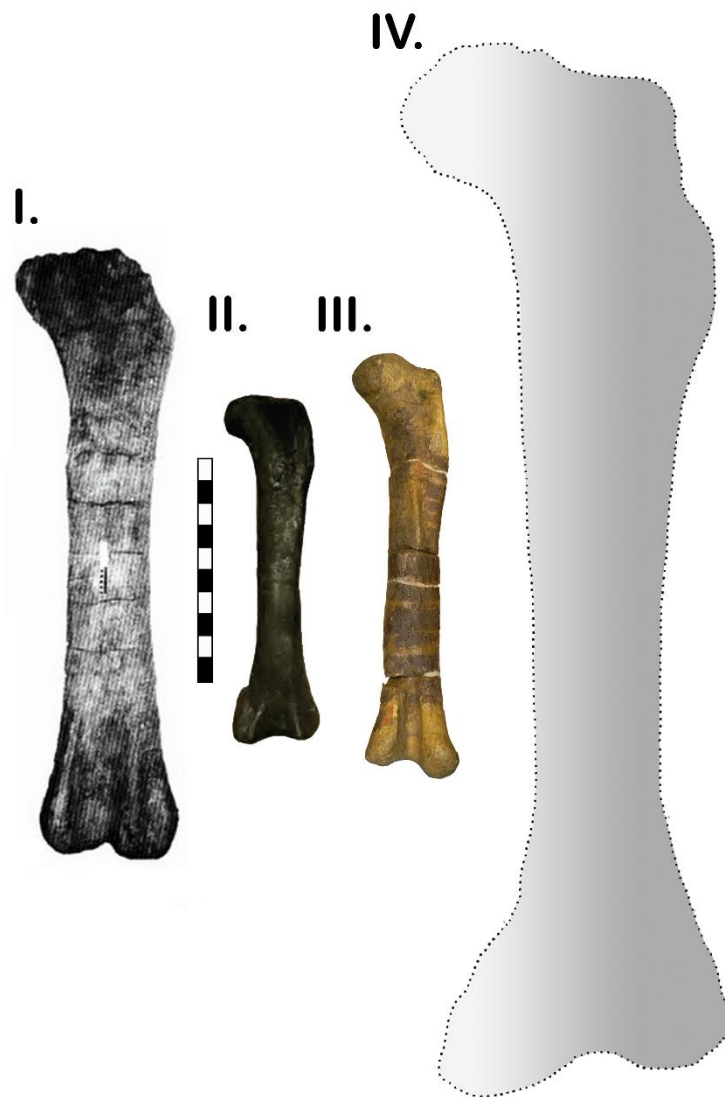
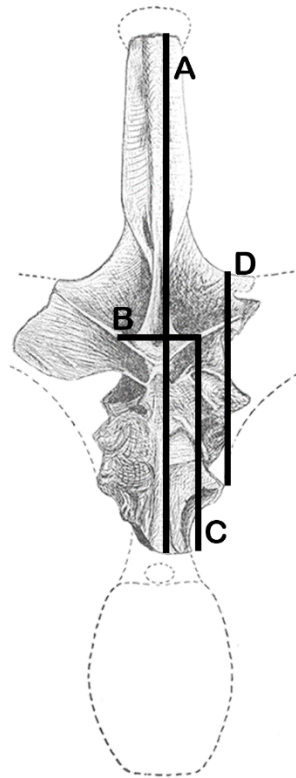


Fig. 5. Hypothetical femur of *Amphicoelias fragillimus* compared to known sauropods. I. *Ruyangosaurus giganteus* (from Lü *et al.*, 2009), II. *Diplodocus carnegii* (CM 84), III. *Amphicoelias altus* (AMNH 5764), IV. *Amphicoelias fragillimus* femur modeled after *A. altus* with implied features modified after *Diplodocus* femur ontogeny. Femora to scale. Scale bar = 1 M.



	New Arch Reconstruction (in mm)	Cope's Measurement (in mm)	Percent Difference
A	1050		
B	168.52	190	11%
C	437.5	585	25%
D	333	390	14%
	Cope's Arch Reconstruction (in mm)	Cope's Measurement (in mm)	Percent Difference
A	1500		
B	228.88	190	20%
C	624.57	585	7%
D	477	390	22%

Fig. 6. Neural arch measurement reference points. A. Neural arch height, B. Transverse expanse of posterior zygapophyses, C. Elevation of the posterior zygapophyses, D. Vertical diameter of the base of diapophysis. *Amphicoelias fragillimus* drawing from Cope, 1878a.