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A nearly complete skull of the sauropod dinosaur *Diamantinasaurus matildae* from the Upper Cretaceous Winton Formation of Australia and implications for the early evolution of titanosaurs

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Titanosaurian sauropod dinosaurs were diverse and abundant throughout the Cretaceous, with a global distribution. However, few titanosaurian taxa are represented by multiple skeletons, let alone skulls. *Diamantinasaurus matildae*, from the lower Upper Cretaceous Winton Formation of Queensland, Australia, was heretofore represented by three specimens, including one that preserves a braincase and several other cranial elements. Herein, we describe a fourth specimen of *Diamantinasaurus matildae* that preserves a more complete skull—including numerous cranial elements not previously known for this taxon—as well as a partial postcranial skeleton. The skull of *Diamantinasaurus matildae*

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shows many similarities to that of the coeval Sarmientosaurus musacchioi from Argentina (e.g. quadratojugal with posterior tongue-like process; braincase with more than one ossified exit for cranial nerve V; compressed-cone-chisel-like teeth), providing further support for the inclusion of both taxa within the clade Diamantinasauria. The replacement teeth within the premaxilla of the new specimen are morphologically congruent with teeth previously attributed to Diamantinasaurus matildae, and Diamantinasauria more broadly, corroborating those referrals. Plesiomorphic characters of the new specimen include a sacrum comprising five vertebrae (also newly demonstrated in the holotype of *Diamantinasaurus matildae*), rather than the six or more that typify other titanosaurs. However, we demonstrate that there have been a number of independent acquisitions of a six-vertebrae sacrum among Somphospondyli and/or that there have been numerous reversals to a five-vertebrae sacrum, suggesting that sacral count is relatively plastic. Other newly identified plesiomorphic features include: the overall skull shape, which is more similar to brachiosaurids than 'derived' titanosaurs; anterior caudal centra that are amphicoelous, rather than procoelous; and a pedal phalangeal formula estimated as 2-2-3-2-0. These features are consistent with either an early-branching position within Titanosauria, or a position just outside the titanosaurian radiation, for Diamantinasauria, as indicated by alternative character weighting approaches applied in our phylogenetic analyses, and help to shed light on the early assembly of titanosaurian anatomy that has until now been obscured by a poor fossil record.

1. Introduction

The early evolution of titanosaurian sauropod dinosaurs remains poorly understood [1,2], despite the fact that the fossil record of the clade spans much of the Cretaceous [3-7] and includes exemplars from every continent (e.g. [8-15]). One of the greatest impediments to resolving titanosaur phylogenetics has been the dearth of cranial remains. Several of the best known titanosaur skulls date to the latest Cretaceous, namely those of Rapetosaurus krausei from Madagascar [16,17], and Nemegtosaurus mongoliensis and Quaesitosaurus orientalis from Mongolia [18-20]. However, of critical importance to understanding the early evolution of Titanosauria has been the recent description of two titanosaur skulls from the mid-Cretaceous of South America: Tapuiasaurus macedoi from the Aptian of Brazil [21,22], and Sarmientosaurus musacchioi from the Cenomanian— Turonian of Argentina [23]. The skull of Tapuiasaurus is strikingly similar to those of the substantially geochronologically younger titanosaurs of Madagascar and Mongolia [21]. This implies either marked cranial morphological conservatism across a vast spatiotemporal interval, or convergent evolution. By contrast, the skull of Sarmientosaurus appears to be somewhat intermediate in morphology between earlier-deriving titanosauriforms such as brachiosaurids (e.g. 'Brachiosaurus' sp. [24]; Giraffatitan brancai [25]; Abydosaurus mcintoshi [26]), and most other titanosaurs known from substantial cranial remains [23]. Thus, Sarmientosaurus more fully embodies the cranial morphology perhaps expected of an early-branching titanosaur than does Tapuiasaurus, despite the latter taxon being geochronologically older. Unfortunately, very little is known of the postcranial anatomy of Sarmientosaurus, making it difficult to constrain its phylogenetic relationships [23] and leaving it susceptible to 'monophyly of the preserved' branch attraction effects (see [5]).

Although Cretaceous sedimentary rocks in northeast Australia have produced abundant evidence of sauropod dinosaurs [9,27–44], very few cranial elements have been discovered. Indeed, aside from teeth and a dentary fragment [33,35,43], only one partial skull has been described to date: that of a referred specimen of *Diamantinasaurus matildae* (AODF 0836), which preserves a braincase and several additional cranial remains [38,41]. Numerous similarities between the cranial remains and teeth of *Diamantinasaurus* and *Sarmientosaurus* have been identified, and a close relationship between these taxa, within the early-branching titanosaurian clade Diamantinasauria, has been supported by phylogenetic analyses [41]. Nevertheless, the overall morphology of the skull of *Diamantinasaurus* remains unclear, as does the position of the clade Diamantinasauria within Titanosauria [1,45].

In this paper, we describe a new specimen of *Diamantinasaurus matildae*, comprising a partial skull with associated postcrania from the lower Upper Cretaceous Winton Formation of Queensland, northeast Australia. We compare this specimen with other sauropods, assess its phylogenetic position, and discuss its evolutionary implications.

1.1. Institutional abbreviations

AODF, Australian Age of Dinosaurs Museum Fossil, Winton, Australia; AODL, Australian Age of Dinosaurs Museum Locality, Winton, Australia; ASDM, Arizona-Sonora Desert Museum, Tucson, USA;

CAMSM, Sedgwick Museum of Earth Sciences, University of Cambridge, England; CPT, Museo Aragonés de Paleontología, Fundación Conjunto Paleontológico de Teruel-Dinópolis, Teruel, Spain; DMNH, Denver Museum of Natural History, Denver, USA; DNHM, Dalian Natural History Museum, Liaoning, China; FC-DPV, Colección de Vertebrados Fósiles, Facultad de Ciencias, Universidad de la República, Uruguay; FMNH PR, Field Museum of Natural History, Chicago, USA; FPDM, Fukui Prefectural Dinosaur Museum, Fukui, Japan; FWMSH, Fort Worth Museum of Science and History, Fort Worth, USA; GSI, Geological Survey of India, Kolkata, India; IANIGLA-PV, Instituto Argentino de Nivología, Glaciología y Ciencias Ambientales, Colección de Paleovertebrados, Mendoza, Argentina; IVPP, Institute of Vertebrate Paleontology and Paleoanthropology, Chinese Academy of Sciences, Beijing, China; IZANUZ, Institute of Zoology, Uzbekistan Academy of Sciences, Tashkent, Uzbekistan; MACN PV, Museo Argentino de Ciencias Naturales 'Bernardino Rivadavia', Buenos Aires, Argentina; MAL, Malawi Department of Antiquities Collection, Lilongwe and Nguludi, Malawi; MAU, Museo Municipal 'Argentina Urquiza', Rincón de los Sauces, Argentina; MCF-PVPH, Museo Carmen Funes, Plaza Huincul, Neuquén, Argentina; MCNA, Museo de Ciencias Naturales de Álava/Arabako Natur Zientzien Museoa, Vitoria-Gasteiz, Spain; MCS, Museo de Cinco Saltos, Río Negro, Argentina; MDE, Le Musée des Dinosaures in Espéraza, France; MLP Av., Museo de La Plata, Rancho de Ávila Collection, La Plata, Argentina; MLP CS, Museo de La Plata, Cinco Saltos Collection, La Plata, Argentina; MMS/VBN, Musée Moulin Seigneurial/Velaux-La Bastide Neuve, Bouches-du-Rhône, France; MPCA-Pv, Museo Provincial 'Carlos Ameghino', Cipolletti, Río Negro, Argentina; MPEF-PV, Museo Paleontológico Egidio Feruglio, Trelew, Argentina; MPM, Museu de Paleontologia de Marília, Marília, São Paulo, Brazil; MPM PV, Museo Padre Molina, Río Gallegos, Argentina; MPZ, Museo Paleontológico de la Universidad de Zaragoza, Zaragoza, Spain; MRS Pv, Museo de Rincón de los Sauces, Neuquén, Argentina; MUCPv, Museo de Geología y Paleontología de la Universidad Nacional del Comahue, Argentina; MUVP, Mansoura University Vertebrate Paleontology Center, Mansoura University, Egypt; MZSP-PV, Museu de Zoologia da Universidade de São Paulo, Brazil; NMMNH, New Mexico Museum of Natural History and Science, Albuquerque, USA; PMU, Paleontological Museum of the University of Uppsala, Uppsala, Sweden; PVL, Fundacion Miguel Lillo, Universidad Nacional de Tucuman, San Miguel de Tucuman, Argentina; QM, Queensland Museum (Brisbane, Australia); RRBP, Rukwa Rift Basin Project, Tanzanian Antiquities Unit, Dar es Salaam, Tanzania; SM, Sirindhorn Museum, Changwat Kalasin, Thailand; TMM, Texas Memorial Museum, University of Texas, Austin, USA; TV, Musée des Dinosaures à Savannakhet, Laos; UA, Université d'Antananarivo, Antananarivo, Madagascar; UBB NVM1, Babes-Bolyai University, Cluj-Napoca, Romania; UNCUYO-LD, Universidad Nacional de Cuyo, Mendoza, Argentina; UNPSJB-PV, Universidad Nacional de la Patagonia 'San Juan Bosco'-Paleovertebrados, Comodoro Rivadavia, Argentina; USNM, National Museum of Natural History, Washington, DC, USA; Z. PAL, Instytut of Paleobiologii, Polish Academy of Sciences, Warsaw, Poland; ZIN PH, Paleoherpetological Collection, Zoological Institute, Russian Academy of Sciences, Saint Petersburg, Russia.

2. Geological setting

The Winton Formation (figure 1) is the stratigraphically uppermost Mesozoic-aged sedimentary unit in the Eromanga Basin of northeast Australia [48]. The mudstones, siltstones, sandstones and minor conglomerates that make up this unit were deposited on a low relief, forested floodplain [49], shortly after the recession of the epeiric Eromanga Sea [50,51]. The top 100 m of the Winton Formation (and the underlying Mackunda Formation) was subjected to substantial chemical weathering during the Cretaceous and early Cenozoic, following cessation of sedimentation in the Eromanga Basin [48]. Consequently, despite its great lateral extent, Winton Formation outcrop is limited to nodular sandstones, as well as erosion-resistant mesas comprising chemically weathered siltstones and sandstones [51]. The remaining Winton Formation is now blanketed by montmorillonite-rich soil [48].

Vertebrates are represented in the Winton Formation by both body fossils and ichnofossils. Actinopterygians are represented by the ichthyodectiform *Cladocyclus geddesi* from near Isisford [52], as well as a possible feeding trace from near Winton [42]. Isolated toothplates of two ceratodontoid dipnoan species (*Metaceratodus wollastoni* and *M. ellioti*) are known from several widely dispersed localities, [53–55], and possible lungfish feeding traces have also been identified near Winton [42]. Lissamphibians and synapsids are as yet unknown from the Winton Formation, whereas lepidosauromorphs are represented by a single vertebra originally interpreted as a dolichosaurid [56], but more recently regarded as an indeterminate varanoid [57]. Although fossil remains of testudinatans (possibly chelids) are seemingly

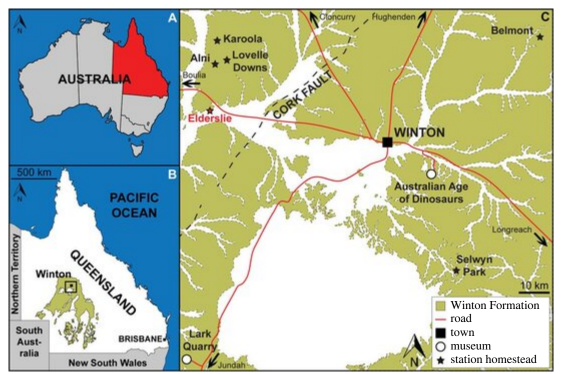


Figure 1. Location of the 'Ann' Site (AODL 0252). (*a*) Map of Australia showing the location of Queensland. (*b*) Map of Queensland showing the distribution of Winton Formation outcrop. (*c*) Map of the Winton area showing the extent of Winton Formation outcrop, the location of Elderslie Station and numerous other stock stations and sites in the area from which sauropod fossils have been collected. Map drafted by the senior author (S.F.P.) in Adobe Illustrator CC 2022 (modified from Poropat *et al.* [43], incorporating geological information from Vine [46] and Vine and Casey [47] (© Commonwealth of Australia (Geoscience Australia) 2022. This product is released under the Creative Commons Attribution 4.0 International Licence. http://creativecommons.org/licenses/by/4.0/legalcode).

fairly common near both Winton and Eromanga [9,34], the only described evidence for their presence constitutes possible ichnites from near Winton [42]. Crocodyliforms are well represented near both Isisford and Winton, with the former region producing Isisfordia duncani [58,59], and the latter producing Confractosuchus sauroktonos [60], as well as several trackways made by swimming or underwater-walking crocodyliforms [42]. Pterosaurs are rare, with the anhanguerid Ferrodraco lentoni and an isolated anhanguerian femur from near Winton being the only specimens described to date [61-63]. Ankylosaurian dinosaurs are also uncommon, the only published specimens constituting three teeth from near Winton [64]. The ornithopod body fossil record is restricted to a single tooth from near Winton [65,66], a partial skeleton found within the body cavity of Confractosuchus sauroktonos [60], and undescribed specimens from near Isisford [67]. Conversely, the ornithopod ichnofossil record is extensive, with both Lark Quarry and the Snake Creek Tracksite providing evidence of small- to medium-sized ornithopods [42,68-75]. Theropod teeth are fairly commonly found at vertebrate sites near Winton [76], but other body fossils are presently limited to the holotype specimen of the megaraptorid Australovenator wintonensis [9,76-78] and an indeterminate megaraptorid [79]. Tracks possibly made by theropods are also fairly common near Winton [42,68-70], although their attribution to theropod trackmakers has been the subject of some debate [42,71–75,80]. Sauropods are common as body fossils in the Winton Formation [28–31,33,42], with four taxa described to date: the early-branching somphospondylan (or perhaps diamantinasaurian titanosaur [34]) Wintonotitan wattsi [9,36] and the diamantinasaurian titanosaurs Diamantinasaurus matildae [9,37,38,41,44] and Savannasaurus elliottorum [38,40] from near Winton, and the diamantinasaurian titanosaur Australotitan cooperensis from near Eromanga [34]. Sauropod tracks have also been reported from both Winton [42] and Eromanga [34].

The new sauropod site, AODL 0252 (the 'Ann' Site), is approximately $10 \text{ m} \times 6 \text{ m}$ in area, but is divisible into two concentrated sections (one west and one east), between which few fragments of bone were found (figure 2). The western section of AODL 0252, which is about $3 \text{ m} \times 3 \text{ m}$, has yielded a partial sacrum, an anterior caudal vertebra, a chevron, the left ilium, the left pubis, both ischia, and much of the right hind limb including the femur, tibia, fibula, metatarsals I–V, and several pedal

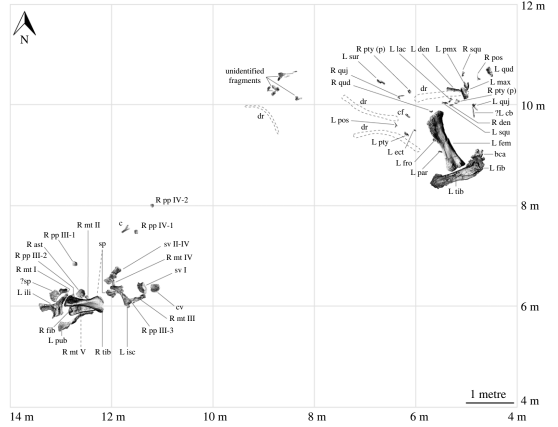


Figure 2. Diamantinasaurus matildae referred specimen (AODF 0906) site map. Site map showing the distribution of skeletal elements of the newly referred Diamantinasaurus matildae specimen (AODF 0906) within the 'Ann' Site (AODL 0252). ast, astragalus; bca, braincase; c, chevron; cb, ceratobranchial; cf, cervical fragment; cv, caudal vertebra; den, dentary; dr, dorsal rib; ect, ectopterygoid; fem, femur; fib, fibula; fro, frontal; ili, ilium; isc, ischium; L, left; lac, lacrimal; max, maxilla; mt ##, metatarsal and position; (p), part; par, parietal; pmx, premaxilla; pos, postorbital; pp ##-#, pedal phalanx and position; pty, pterygoid; pub, pubis; qud, quadrate; quj, quadratojugal; R, right; sp, sacral process; squ, squamosal; sur, surangular; sv #, sacral vertebra and position; tib, tibia.

phalanges including at least one ungual. Thus, the western section preserves elements restricted to the posterior half of the animal. The proximal end of the left pubis, and the right femur, tibia and fibula were found in the 'black soil' layer that represents the weathered cap of the Winton Formation. The remaining bones were found below the 'black soil' in the Winton Formation proper. The long axes of the left pubis, right tibia and right fibula were effectively parallel. Four of the five metatarsals and the probable astragalus fragment were found immediately beneath the tibia and fibula, and the fifth was found approximately 50 cm northeast, associated with fragmentary sacral vertebrae. The sole chevron and two other pedal phalanges were found 1-1.5 m northeast of this concentration of bones. Thus, all of these remains were found in close association.

The eastern section of AODL 0252, which is also about 3 m \times 3 m, yielded all of the sauropod cranial remains recovered from this site. The skull was almost entirely disarticulated, and its various elements were scattered around a partial left hind limb (comprising the femur, tibia and fibula) and several partial dorsal ribs. The left tibia and fibula were probably articulated prior to the incorporation of the former into the 'black soil', and both were found at the distal end of the left femur, with their long axes perpendicular to its long axis. The braincase was found near the proximal end of the left fibula, and several additional elements (including the left frontal, left parietal and a fragment of the right quadrate) were found scattered within the lowest sedimentary level occupied by the femur (an approx. 10 cm thick layer of siltstone, blue-grey when unweathered). Three partial dentigerous elements, all from the left side, were found near the proximal end of the femur. The left premaxilla was found wedged within the antorbital foramen of the maxilla, whereas the dentary was found in contact with, and with its long axis perpendicular to that of, the dentigerous portion of the maxilla. The remaining cranial elements were found in the same horizon, within 1.5 m of the proximal end of the femur.

All of the sauropod elements found within AODL 0252 are size congruent (notably the hind limb elements from the western and eastern sections), some articulate perfectly with one another, and there is no duplication of elements. Moreover, there is no evidence of other dinosaurs at the site, and little evidence of other animals (beyond a few possible turtle and fish fragments). Thus, we interpret all of the sauropod remains from AODL 0252 as deriving from a single individual. Based on measurements of overlapping elements, the sauropod at AODL 0252 was approximately the same size as, or perhaps slightly larger than, the *Diamantinasaurus matildae* holotype specimen (AODF 0603); however, it appears to have been more gracile based on the dimensions of the hind limb elements. Based on the size of the skull elements (e.g. quadrate dorsoventral height 216 mm), the AODF 0906 skull represents a slightly larger individual than AODF 0836 (quadrate dorsoventral height 184 mm).

3. Methods

Permission to dig on Elderslie Station was granted to the Australian Age of Dinosaurs Museum of Natural History by the landowners.

The cranial remains of AODF 0906 were CT scanned at St Vincent's Hospital (Melbourne, Victoria). One objective of this was to determine whether or not any replacement teeth were present within the dentigerous elements. As several were identified in the premaxilla, this element was subsequently scanned at the Australian Nuclear Science and Technology Organisation's (ANSTO) Australian Synchrotron (Melbourne, Australia) to provide higher-resolution imagery. Microtomographic measurements of the sauropod left premaxilla were performed using the Imaging and Medical Beamline (IMBL) at the Australian Synchrotron. For this investigation, acquisition parameters included a pixel size of 100× 100 µm, a pink beam of X-rays with a peak intensity of 220 keV generated using a superconducting multipole wiggler operating at 3T, a sample-to-detector distance of 500 mm, and use of the 'Quartz' Hamamatsu C9252-DK14 flat panel photodiode array imager (2432 x 100 pixels). As the height and width of the specimen exceeded the detector field-of-view, the specimen was aligned axially relative to the beam, its centre of rotation shifted toward one edge of the detector. Nineteen successive scans were required to cover the full specimen volume, each consisting of 1800 equally spaced angle shadowradiographs with an exposure length of 0.1 s, obtained every 0.10° as the sample was continuously rotated 180° about its vertical axis. Vertical translation of the specimen between tomographic scans was 80 mm. 100 dark (closed shutter) and beam profile (open shutter) images were obtained for calibration before and after shadow-radiograph acquisition. Total time for the scan was 70 min.

The raw 16-bit radiographic series were normalized relative to the beam calibration files and stitched with the in-house software 'IMBL-Stitch' to yield a 32-bit dataset which was reconstructed to yield a three-dimensional tomogram by the filtered-back projection method using the CSIRO's X-TRACT [81]. All tomographic scan data were rendered and visualized in Dragonfly v.2020.1.1.809 (Object Research Systems: www.theobjects.com/dragonfly).

All elements were surface scanned at Australian Age of Dinosaurs Museum of Natural History using an Artec Space Spider (www.artec3d.com/portable-3d-scanners/artec-spider-v2). All three-dimensional surface models were manipulated, and the skull reconstruction assembled, in Artec Studio 15 Professional (www.artec3d.com/3d-software/artec-studio). The screenshots of the elements used in the figures of three-dimensional models were taken in MeshLab (www.meshlab.net), owing to its customizable lighting. All figures were assembled in Adobe Photoshop 2022 and 2023 (www.adobe.com/au/products/photoshop.html) and outlined and annotated in Adobe Illustrator 2022 and 2023 (www.adobe.com/au/products/illustrator.html).

We scored AODF 0906 for the phylogenetic matrix of Poropat *et al.* [41]. We modified one existing character and added four new ones, while also altering several character scores for operational taxonomic units (OTUs) already included in the matrix based on new information on *Diamantinasaurus matildae* [41,43] and *Phuwiangosaurus sirindhornae* [82], as well as re-evaluation of the CT scan data of *Sarmientosaurus musacchioi* [23]. These additions and changes are documented in the appendix. The revised data matrix comprises 126 OTUs, scored for 556 characters, and is provided as a TNT file (electronic supplementary material).

A taxonomic aside: we have referred to USNM 5730 as simply *Brachiosaurus* in [2], and as '*Brachiosaurus*' sp. in this paper and a previous one [41], because the referral of this specimen to *Brachiosaurus altithorax* by D'Emic and Carrano [24] was not made based on direct anatomical overlap between that specimen and the holotype of the taxon (FMNH P25107), which comprises only postcrania [83–85]. The other specimen assigned to *Brachiosaurus altithorax* by D'Emic and Carrano

[24] (BYU 9754(4744)/USNM 21903) fails to bridge the anatomical gap: it comprises only postcrania, like the holotype. Thus, the referral of USNM 5730 to *Brachiosaurus* is circumstantial: the skull represents a brachiosaurid, it is from the Morrison Formation, and the only brachiosaurid heretofore named from the Morrison Formation is *Brachiosaurus altithorax*. In our view, this laissez faire taxonomic approach is not adequate grounds for referral, especially given that the Morrison Formation represents approximately 7 Myr of sedimentary deposition [86].

Phylogenetic analyses were run in TNT v.1.6 [87–89]. Eighteen of the 556 characters were treated as ordered (11, 14, 15, 27, 40, 51, 104, 122, 147, 148, 195, 205, 259, 297, 426, 435, 472 and 510). Eight taxa that have been determined in previous analyses to be unstable [6,41,90,91] were excluded *a priori*: *Astrophocaudia slaughteri, Australodocus bohetii, Brontomerus mcintoshi, Fukuititan nipponensis, Fusuisaurus zhaoi, Liubangosaurus hei, Malarguesaurus florenciae* and *Mongolosaurus haplodon*. The protocol employed for the equal weights and extended implied weights analyses was identical to that used by Poropat *et al.* [41]: under 'New Technology Search', we used the 'Stabilize Consensus' option with sectorial searches, drift and tree fusing. After five rounds of consensus stabilizing, the resultant trees were used as the starting topologies for a 'Traditional Search', which used tree bisection–reconnection (TBR). Two versions of the analysis were run: one with equal character weighting, the other with extended implied weighting and a *k*-value of 9 [92,93]. Following Poropat *et al.* [41], two further taxa were excluded *a priori* from analyses applying equal character weighting, namely the 'Cloverly titanosauriform' and *Ruyangosaurus giganteus*; these taxa were retained in the extended implied weighting analysis.

4. Systematic palaeontology

Sauropoda [94]
Macronaria [95]
Titanosauriformes [96,97]
Somphospondyli [95]
Diamantinasauria [41]
Diamantinasaurus matildae [9]

4.1. Holotype specimen

AODF 0603 ('Matilda'; AODL 0085): dentary fragment, tooth, three partial cervical ribs, three incomplete dorsal vertebrae, dorsal ribs, fragmentary gastralia, five coalesced sacral vertebrae, isolated sacral processes, right and left scapulae, right coracoid, partial right sternal plate, right and left humeri, right and left ulnae, right radius, right and left metacarpals I–V, eight manual phalanges (including right manual ungual I-2), right and left lilia, right and left pubes, right and left ischia, right femur, right tibia, right fibula, right astragalus, associated fragments [37,43].

4.2. Previously referred specimens

AODF 0836 ('Alex'; AODL 0127): left frontal, right and left parietals, left squamosal, right and left quadrates, braincase (comprising supraoccipital, right and left exoccipital–opisthotics, basioccipital, partial basisphenoid, right and left prootics, right and left laterosphenoids, right and left orbitosphenoids, and right and left possible sphenethmoids), left surangular, atlas intercentrum, axis, cervical vertebrae III–VI, middle/posterior cervical vertebral neural arch, three dorsal vertebrae, dorsal ribs, two co-ossified sacral vertebrae, right scapula, right and left iliac preacetabular processes, right and left pubes, right and left ischia, and abundant associated fragments, many representing ribs or partial vertebrae [38,41]; AODF 0663 ('Oliver'; AODL 0122): one cervical rib, two dorsal vertebral centra, three dorsal vertebral neural arches, several dorsal ribs, left scapula, right humerus, right manual ungual I-2, right femur, and associated fragments [44].

4.3. Newly referred specimen

AODF 0906 ('Ann'; AODL 0252): partial skull comprising left premaxilla, left maxilla, left lacrimal, left frontal, left parietal, right and left postorbitals, right and left squamosals, right and left quadratojugals, right and left quadrates, right and left pterygoids, left ectopterygoid, braincase (comprising supraoccipital, partial right and left exoccipital–opisthotics, fragmentary basioccipital, right and left

prootics, right and left laterosphenoids, right and left orbitosphenoids, and a possible right sphenethmoid), right and left dentaries, left surangular, ?left ceratobranchial, four dorsal ribs, five sacral centra, several sacral processes, one anterior caudal vertebra, one chevron, left ilium, left pubis, right and left ischia, right and left femora, right and left tibiae, right and left fibulae, a probable right astragalus fragment, right metatarsals I–V, right pedal phalanges III-1–3 and IV-1–2, and associated fragments.

4.4. Newly identified diagnostic characteristics

Quadratojugal and quadrate with horizontal ridge present across both elements anterior to their articulation point (lateral surface of quadrate, medial surface of quadratojugal).

4.5. Locality

AODL 0252 (The 'Ann' site), Elderslie Station, approximately 60 km NW of Winton, Queensland, Australia. Approximate coordinates 22° 15′ S, 142° 30′ E.

4.6. Horizon

Winton Formation; Cenomanian-lowermost Turonian, lower Upper Cretaceous [98,99].

5. Skull

5.1. General shape

The reconstructed skull of AODF 0906 is approximately 500 mm long anteroposteriorly. Based on the transverse width of the incomplete braincase, the mediolateral width of the left parietal (table 1), and the three-dimensional reconstruction of the skull presented herein (figure 3), the skull would have been approximately 250 mm wide transversely along its posterior margin. The incomplete left quadrate is approximately 210 mm tall dorsoventrally, indicating the minimum height of the posterior margin of the skull. Thus, the skull of AODF 0906 is slightly larger than the holotype skull of Sarmientosaurus musacchioi [23].

The premaxilla of Diamantinasaurus is not stepped, distinguishing the 'snout' from that of taxa such as Camarasaurus and Malawisaurus dixeyi ([100,101]; although see [102], and [103]). However, it is likely that the snout was 'boot'-shaped, as in brachiosaurids [104] and Sarmientosaurus [23]. As preserved, the premaxillary-maxillary index (PMI): [105] is approximately 45%. All other macronarians have PMI values greater than 60%, and those with anteriorly flattened snouts (like Antarctosaurus wichmannianus) have PMI values that exceed 75% [38,100,105]. However, the unusually low PMI value in Diamantinasaurus is almost certainly an underestimate, owing to the distortion to which the maxilla has been subjected. The nares appear to have been large and were presumably divided by the narial bar (comprising the narial processes of the premaxillae and the premaxillary processes of the nasals), although the direction in which they faced is unclear. By contrast, the antorbital fenestra on each side is small, ovate (dorsally tapered), faces laterally, and is bounded by the maxilla, lacrimal, and jugal; in all of these respects, it is akin to the antorbital fenestra of Sarmientosaurus [23]. The orbit of Diamantinasaurus was presumably ovate (tapering ventrally), although this cannot be stated with certainty, owing to the non-preservation of the jugal or prefrontal. The infratemporal fenestra is taller dorsoventrally than it is long anteroposteriorly, and is subtriangular, being expanded ventrally. This differentiates AODF 0906 from the titanosaur Rapetosaurus, which has slit-like infratemporal fenestrae [17]. The anterior margin of the infratemporal fenestra is situated posterior to the midpoint of the orbit, as in all macronarians [100,101,106,107]. The supratemporal fenestra is large, as in all neosauropods other than rebbachisaurids [104,108], and is bordered by the frontal, postorbital, squamosal and parietal. As in all somphospondylans, the greatest diameter of each supratemporal fenestra in Diamantinasaurus is less than the transverse distance between them [6,101,104]. The temporal bar precludes the supratemporal fenestra from being visible in lateral view, a feature that Diamantinasaurus shares with the titanosaurs Nemegtosaurus, Rapetosaurus and Sarmientosaurus, but that distinguishes it from non-titanosaurian macronarians and the titanosaur Tapuiasaurus [22,95,101]. Whether or not an external mandibular fenestra was present cannot be established with certainty in AODF 0906. However, it is probable that this fenestra was absent given that virtually all other

Table 1. Measurements of the skull and hyobranchial elements of AODF 0906 *Diamantinasaurus matildae*. An asterisk (*) indicates a tentative measurement based on an incomplete or distorted specimen.

element	dimension measured	measurement (mm)
premaxilla (left)	mesiodistal length along jawline	77
	dorsoventral height (perpendicular to jawline)	146
maxilla (left)	mesiodistal length along jawline	246*
	maximum dorsoventral height of base (anterior)	114
	maximum dorsoventral height of base (posterior)	74
	maximum dorsoventral height (total)	220
lacrimal (left)	maximum dorsoventral height	110
	maximum anteroposterior length (dorsal)	47
	maximum mediolateral width (dorsal)	24
	maximum anteroposterior length (ventral)	68
	maximum mediolateral width (ventral)	23
frontal (left)	maximum anteroposterior length	58*
	maximum mediolateral width	73*
parietal (left)	maximum dorsoventral height	51*
	maximum mediolateral width	108
postorbital (left)	maximum anteroposterior length	72*
	maximum dorsoventral height	117
postorbital (right)	maximum anteroposterior length	81
	maximum dorsoventral height	123
squamosal (left)	maximum anteroposterior length	50*
	maximum dorsoventral height	96*
squamosal (right)	maximum anteroposterior length	52*
1 (3)	maximum dorsoventral height	96*
quadratojugal (left)	maximum anteroposterior length	149
	maximum dorsoventral height	80*
quadratojugal (right)	maximum anteroposterior length	135*
	maximum dorsoventral height	76*
quadrate (left)	maximum dorsoventral height	216*
pterygoid (right)	maximum anteroposterior length	139*
ectopterygoid (left)	maximum anteroposterior length	68
braincase	maximum anteroposterior length	140*
	maximum transverse width	150*
dentary (left)	maximum length along jawline	310*
•	maximum dorsoventral height (anterior)	58*
	maximum dorsoventral height (posterior)	78
surangular (left)	maximum anteroposterior length	190
-	maximum dorsoventral height	60
ceratobranchial (?left)	maximum anteroposterior length	231*

macronarians lack external mandibular fenestrae [106,109]. *Tapuiasaurus* seems to be an exception: the presence of an external mandibular fenestra was tentatively inferred by Martínez *et al.* [23], although this opening was not described by either Zaher *et al.* [21] or Wilson *et al.* [22]. However, given the

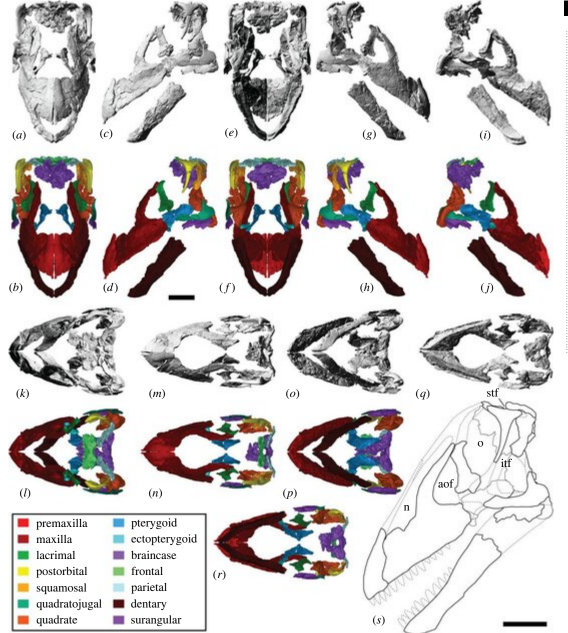


Figure 3. Diamantinasaurus matildae reconstructed skull (AODF 0906). (a-r) Diamantinasaurus matildae referred skull (AODF 0906) reconstruction in (a,b) anterior, (c,d) left lateral, (e,f) posterior, (g,h) right lateral, (i,j) left medial (right elements removed), (k,l) dorsal, (m,n) anterodorsal, (o,p) ventral and (q,r) posteroventral views. (a-r) are screenshots of three-dimensional models derived from surface scans. (s) Diamantinasaurus matildae referred skull (AODF 0906) reconstruction line drawing in left lateral view. aof, antorbital fenestra; itf, infratemporal fenestra; n, naris; o, orbit; stf, supratemporal fenestra. Scale bars = 100 mm.

existence of near-identical elliptical openings on both mandibles (MZSP-Pv 807: PDM, pers. obs. 2019), we contend that *Tapuiasaurus* is characterized by the presence of an external mandibular fenestra.

5.2. Premaxilla

The body of the left premaxilla is complete and generally well preserved, despite having suffered extensive fracturing (figure 4). The lateral surface of the premaxilla of AODF 0906 is smoothly convex (figure 4c,g), lacking the vascular grooves seen in *Nemegtosaurus* [104,110]. A slightly rough texture characterizes the lanceolate articular surface for the right premaxilla, which would have faced medially (figure 4i). The premaxilla articulates with the maxilla posteriorly via a similarly lanceolate articular surface (figure 4d,h). The medial surface of the premaxilla is overlapped by the premaxillary process of the maxilla. In lateral view, the premaxilla-maxilla contact is essentially straight, as in non-

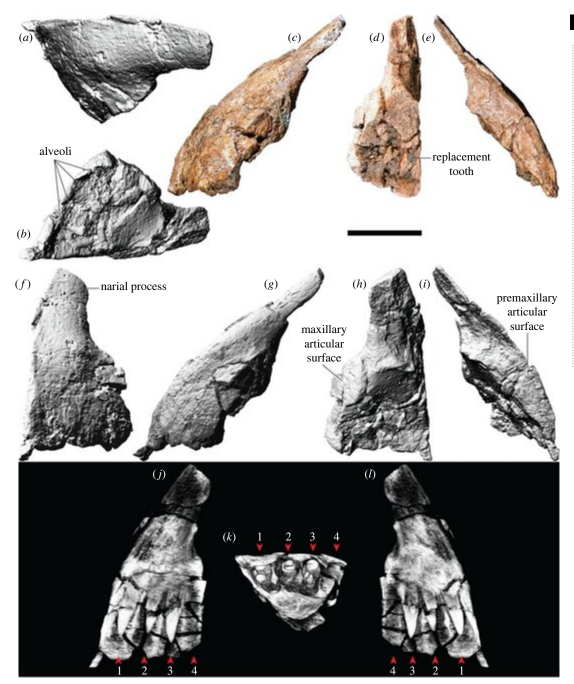


Figure 4. Diamantinasaurus matildae referred left premaxilla (AODF 0906). (a-i) Left premaxilla in (a) dorsal, (b) ventral, (c,g) lateral, (d,h) posterior, (e,i) medial and (f) anterior views. (a,b) and (f-i) are screenshots of three-dimensional models derived from surface scans, and (c-e) are photographs. (j-l) Three-dimensional digital models (derived from synchrotron data) of the left premaxilla in (j) labial, (k) occlusal and (l) lingual views. Numbers 1–4 indicate alveoli. Scale bar = 50 mm.

titanosauriform sauropods, *Euhelopus zdanskyi* [111–114], and some titanosaurs (e.g. *Malawisaurus* [115], *Muyelensaurus pecheni* [116], *Tapuiasaurus* [21,22]); this distinguishes *Diamantinasaurus* from brachiosaurids and *Nemegtosaurus*, in which this contact is sinuous [20,26]. Lateral to the tooth row, there is a prominent plate of bone, as in sauropods generally [106,117].

The premaxilla of AODF 0906 appears to lack maxillary processes. If this absence is genuine, this would serve to distinguish *Diamantinasaurus* from *Europasaurus holgeri* and *Camarasaurus*, in which both dorsomedial and ventromedial maxillary processes are present on the premaxilla [118,119], and from *Malawisaurus* and *Tapuiasaurus*, in which a single medial process is present [22,115,120]. However, given that maxillary processes are positioned near the base of the narial process of the premaxilla, that they tend to be relatively fragile, and that this portion of the premaxilla is incomplete in AODF 0906, it is possible that they were present in *Diamantinasaurus*, but have been lost.

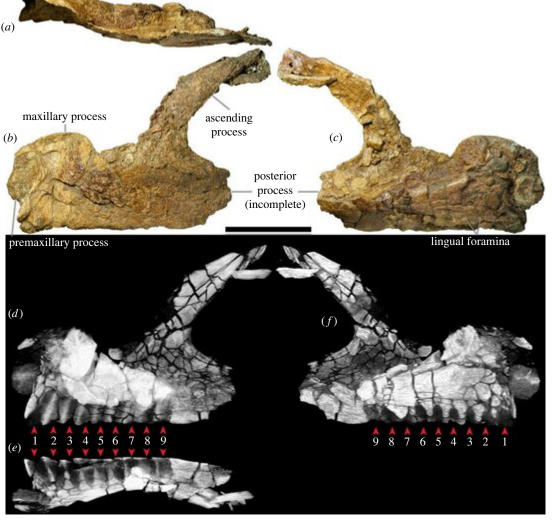


Figure 5. Diamantinasaurus matildae referred left maxilla (AODF 0906). (a-c) Photographs of the left maxilla in (a) dorsal, (b) lateral and (c) medial views. (d-f) three-dimensional digital models (derived from medical CT data) of the left maxilla in (d) lateral, (e) ventral and (f) medial views. Numbers 1–9 indicate alveoli. Scale bar = 50 mm.

The premaxilla of AODF 0906 hosts four tooth positions (figure 4b,j-l), as in sauropods generally [8,109]. The only reported exception—an embryonic titanosaurian from the Late Cretaceous of Argentina, described by Kundrát et al. [121] as having five premaxillary alveoli—might have been incorrectly interpreted: based on their size relative to each other and to the other alveoli, the mesialmost two alveoli in this specimen could represent a single alveolus. Although the active teeth have been dislodged from the premaxilla in AODF 0906, a total of five replacement teeth are present in the crypt (figure 4j-l): two in the first alveolus (both of which are partly exposed on the specimen in lingual view owing to loss of part of the wall of the crypt), one in the second alveolus, and two in the third one. The fourth alveolus is vacant.

5.3. Maxilla

The left maxilla (figures 5 and 6), which is lacking only the active dentition and the terminus of the posterior process, has suffered such extensive distortion that precise physical articulation with the premaxilla is not possible. Regardless, it is still morphologically informative.

The maxilla of AODF 0906 comprises a subrectangular dentigerous portion, a posterodorsally tapered ascending process, and an incomplete posterior process. Anteromedially, a prominent premaxillary process (anteromedial process sensu Wilson [20]) is present. Manifested as a semicircular flange, it is dorsoventrally taller, and consequently extends much further ventrally, than the same process in Europasaurus [119], Camarasaurus [118], 'Brachiosaurus' [24], 'Astrodon' / 'Pleurocoelus' [122,123] and Nemegtosaurus [20]. Although this potentially represents an autapomorphy of Diamantinasaurus, the

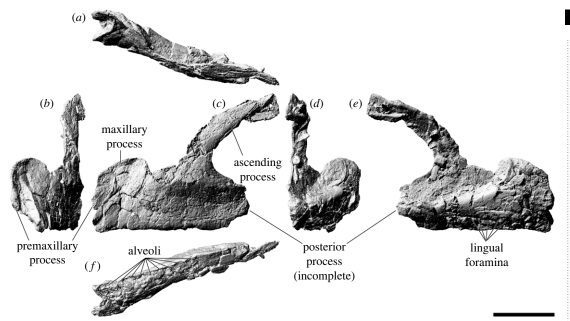


Figure 6. Diamantinasaurus matildae referred left maxilla (AODF 0906). (a-f) Three-dimensional digital models (derived from surface scans) of the left maxilla in (a) dorsal, (b) anterior, (c) lateral, (d) posterior, (e) medial and (f) ventral views. Scale bar = 50 mm.

morphology of this process cannot be assessed in most somphospondylans, either owing to genuine absence or non-preservation. Moreover, it is likely that the present orientation and position of this process in AODF 0906 has been impacted by distortion: it is not hard to conceive that this process was originally oriented more anteriorly (and medially) than preserved. At the posteroventral base of this process, a small foramen is present; following Madsen *et al.* [118], this is the subnarial foramen.

The premaxillary process unites dorsally with a flange of bone that presumably represents the contact for the contralateral maxilla. We suggest that this flange, which is now essentially vertical, would have been inclined dorsomedially in life, owing to the mediolateral compression to which the maxilla has been subjected, and based on the morphology of the premaxilla. Posterior to this flange, the dorsomedial margin of the maxilla is incomplete; it is quite likely that this surface was more extensive *in vivo*. A similar inference has been made about an isolated titanosaur maxilla from the mid-Cretaceous of Argentina (UNPSJB-PV 583 [23,124]).

The lateral surface of the dentigerous portion appears to have been completely smooth (figure 5b), lacking the vascular grooves seen in nemegtosaurids and, to a lesser extent, in *Giraffatitan* and *Sarmientosaurus* [6,20,23,25,104]. However, the poor preservation of this surface in AODF 0906 (figure 6c) might have destroyed any evidence of these, had they been present. If a preantorbital opening was present (which is unclear), it was only expressed as a shallow fossa, as in most macronarians [6,21,38,125]. Similarly, it is not clear whether or not an additional foramen anterior to the preantorbital fenestra, which is present in the titanosaurs *Nemegtosaurus* and *Tapuiasaurus* [20–22], was present in AODF 0906.

The preservation of the medial surface of the dentigerous portion of the maxilla evinces the degree of distortion to which this element was subjected (figures 5c,f and 6e). The dorsal surface of the crypt, which should be completely smooth, is broken into segments that have shifted slightly, despite remaining effectively in life position. Nevertheless, morphological information can be gleaned from the medial surface. As in all other sauropods, a rounded special foramen (sensu Edmund [126]; replacement foramen sensu Jin et al. [127]) is present dorsal to each tooth position. Although many of these special foramina are observable, none are complete—the ventral halves of all are missing. Dorsal to these special foramina, the medial surface of the crypt is smooth and essentially flat. Although the dentigerous portion of the maxilla articulated with the palatine and ectopterygoid posteromedially, neither of these contacts is evident, owing to breakage. The roof of the crypt is manifested as a posteroventrally slanted shelf, walled laterally by the base of the ascending process.

The posterior process of the maxilla of AODF 0906 is incompletely preserved but was clearly not as elongate as that of *Nemegtosaurus* [20], *Rapetosaurus* [17], or the embryonic titanosaurs from Auca Mahuevo [128–130]. Based on the preserved length of the dentary and the morphology of the quadratojugals in AODF 0906, we suggest that the posterior margin of the posterior process contacted

the jugal only, as in most macronarians [6,22,23,26,38,106,119]; by contrast, in *Camarasaurus* and *Nemegtosaurus*, the posterior process also has a substantial contact with the quadratojugal [20,118]. Whether or not the ventral margin of the posterior process of the maxilla possessed a distinct ventral emargination, as in other titanosauriforms [3,6,38], is not known.

The ascending process of the maxilla contacts the nasal and lacrimal dorsally. It forms the ventral and lateral margins of the naris and the anteroventral margin of the antorbital fenestra. In AODF 0906, the ascending process is almost complete, albeit broken. The grain of the bone, and the shapes of the fragments near the base, demonstrate that the ascending process should project more dorsally than preserved, relative to the dentigerous portion. However, the distortion to which both the dorsal half of the dentigerous portion and the ventral portion of the ascending process have been subjected implies that a simple dorsal orientation of the ascending process is not correct either: the present vertical orientation of the ascending process is probably only slightly exaggerated relative to its *in vivo* state. It seems unlikely that the ascending process projected posteriorly beyond the posterior process; as such, the ascending process in *Diamantinasaurus* appears to be characterized by the same morphology as most titanosauriforms, other than the titanosaurs *Nemegtosaurus* and *Rapetosaurus* [6,131].

As preserved, the tooth row of AODF 0906 terminates just anterior to the antorbital fenestra (figure 5d-f, 6f). However, given that the angle of the ascending process—and, consequently, the position of the anterior margin of the antorbital fenestra—has been altered by taphonomic distortion, we contend that it is more likely that the tooth row terminated somewhere below the antorbital fenestra. This would suggest that this aspect of the tooth row of Diamantinasaurus is more similar to titanosaurs such as Rapetosaurus and Sarmientosaurus, but differentiates it from others wherein the tooth row terminates anterior to the external naris, including Nemegtosaurus and Tapuiasaurus, as well as brachiosaurids [26,100,101,104,132]. Based on CT scan data of the left maxilla of AODF 0906, nine alveoli are present (figure 5d-f). This is fewer than in the early-diverging macronarian Europasaurus (12–13: [119]), brachiosaurids (10–14: [24– 26,133]), Euhelopus (10: [114]), and the titanosaurs Sarmientosaurus (11–12: [23]) and Tapuiasaurus (12: [21,22]). By contrast, it is greater than the number seen in the Auca Mahuevo titanosaur embryos (7-8: [129]) and (probably) Narambuenatitan (at least 8: [134]). Camarasaurus ranges from having 8 to 10 maxillary alveoli [118,135-140], and in nemegtosaurid titanosaurs either eight or nine maxillary alveoli are present [18-20]. Given that maxillary alveolus count in macronarian sauropods can vary between the two sides of a single individual (e.g. Sarmientosaurus [23]), within taxa known from multiple exemplars (e.g. Giraffatitan [25]), and probably changed ontogenetically (based on the low maxillary tooth count in the Auca Mahuevo titanosaur embryos [129]), attaching significance to maxillary alveolus count is tenuous. Thus, the relatively low maxillary alveolus count of Diamantinasaurus simply serves to distinguish it from most titanosauriforms other than nemegtosaurids.

5.4. Lacrimal

The lacrimal (figure 7) forms the posterior margin of the antorbital fenestra and the anterior margin of the orbit. The dorsal margin would have contacted the ascending process of the maxilla, the nasal and the prefrontal. If the lacrimal of AODF 0906 is complete dorsally, then the facet for the ascending process of the maxilla is a deep furrow (figure $7b_j$); however, it is far more likely that the apex of this element is incomplete, based on both its poor preservation and the height of the lacrimal relative to that of the ascending process of the maxilla (figures 5 and 6). Thus, it cannot be determined whether or not the lacrimal of Diamantinasaurus clasped the ascending process of the maxilla, as in Sarmientosaurus [23]. The dorsal four-fifths of the lacrimal (as preserved) is triangular in cross-section, with the element divided into lateral, anteromedial and posteromedial surfaces; a similar morphology was reported in Rapetosaurus [17] and Sarmientosaurus [23]. By contrast, at its ventral end, the lacrimal is anteroposteriorly flared and mediolaterally compressed. The lateral surface is flat to shallowly concave along its length and separated from the other two surfaces by pronounced flanges. The anteromedial and posteromedial surfaces are also separated by a pronounced medial flange (figure 7e,k). The anteromedial surface is shallowly concave, whereas the posteromedial surface is mostly flat, being concave only near the medial flange. The lacrimal foramen cannot be observed on the posteromedial surface, nor in CT scans of the specimen, owing to the distortion to which the lacrimal has been subjected (figure 7m,n). The lacrimal of AODF 0906 possesses a very weak anterior process (figure 7c, h), as in Giraffatitan, Nemegtosaurus and Sarmientosaurus [20,23,25]; this distinguishes Diamantinasaurus from Abydosaurus, Bonitasaura salgadoi, Rapetosaurus and Tapuiasaurus, wherein the anterior process is more pronounced [17,22,26,141]. The dorsal end of the lacrimal also possesses a posterior process

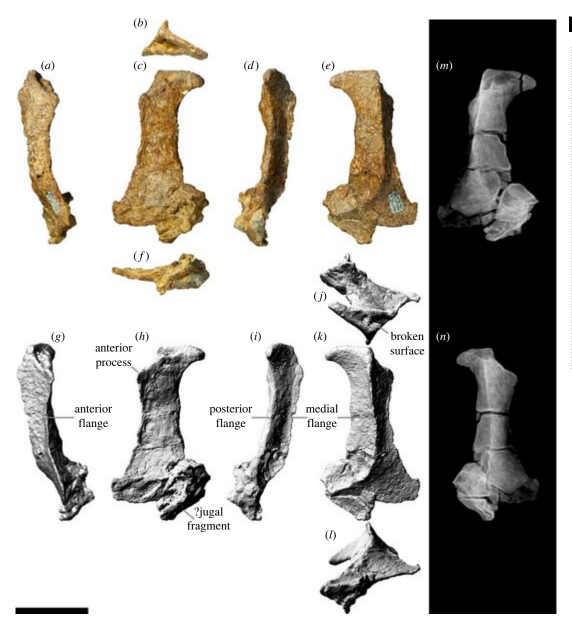


Figure 7. Diamantinasaurus matildae referred left lacrimal (AODF 0906). (a-f) Photographs of the left lacrimal in (a) anterior, (b) dorsal, (c) lateral, (d) posterior, (e) medial and (f) ventral views. (g-l) Three-dimensional digital models (derived from surface scans) of the left lacrimal in (g) anterior, (h) lateral, (i) posterior, (j) dorsal, (k) medial and (l) ventral views. (m,n) Three-dimensional digital models (derived from medical CT data) of the left lacrimal in (m) lateral and (n) medial views. Scale bar = 50 mm.

similar to that seen in *Nemegtosaurus* [20]; a similar process appears to be present in *Sarmientosaurus*, although in this taxon it is obscured laterally by the prefrontal [23].

The ventral margin of the lacrimal would have contacted the jugal ventrally. Indeed, the AODF 0906 lacrimal appears to be preserved in articulation with a fragment of the jugal (figure 7c,h). Based on the morphology of the ventral end of the lacrimal, and our reconstruction of the skull, we infer that the jugal made a small contribution to the antorbital fenestra, as in *Giraffatitan*, *Nemegtosaurus*, *Rapetosaurus* and *Sarmientosaurus* [6,104,142]; this distinguishes AODF 0906 from *Tapuiasaurus*, wherein the contribution of the jugal to the antorbital fenestra is extensive [21,22].

5.5. Frontal

Only the left frontal is preserved (figure 8), and it appears to be incomplete laterally and to have suffered significant *post mortem* damage. Despite its incompleteness, it is approximately 25% wider mediolaterally

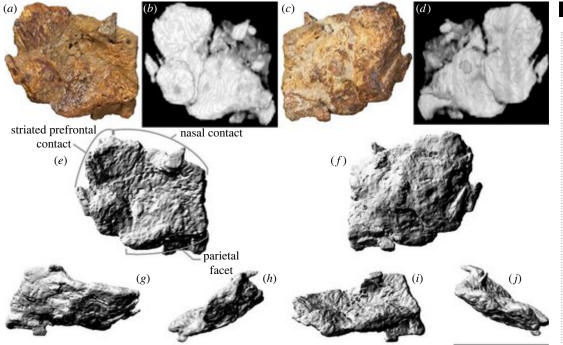


Figure 8. Diamantinasaurus matildae referred left frontal (AODF 0906). (a-d) Left frontal in (a,b) dorsal and (c,d) ventral views. (a) and (c) are photographs, (b) and (d) are three-dimensional models derived from CT data. (e-j) Three-dimensional models (derived from surface scans) of the left frontal in (e) dorsal, (f) ventral, (g) anterior, (h) medial, (i) posterior and (g) lateral views. Scale bar (g) and (g) anterior, (g) anterior, (g) anterior, (g) posterior and (g) lateral views.

than long anteroposteriorly. Thus, as in all non-euhelopodid titanosauriforms [6,38,131,143], including a previously referred specimen (AODF 0836) of *Diamantinasaurus* [38,41], the frontal of AODF 0906 is shorter anteroposteriorly than it is wide mediolaterally. The dorsal surface is broadly concave mediolaterally (figure 8a,e), although it becomes convex medially, as in the early-branching somphospondylan *Phuwiangosaurus sirindhornae* and the titanosaurs *Antarctosaurus*, AODF 0836, *Bonatitan*, *Bonitasaura*, *Narambuenatitan*, *Rapetosaurus* and *Saltasaurus loricatus* [3,6,17,38,41,134,141,144–147]. By contrast, the ventral surface of the AODF 0906 frontal is broadly convex but irregularly undulatory (figure 8c,f), which is presumably a consequence of deformation and/or under-preparation. The ventral surface would have been sutured to the dorsal margin of the orbitosphenoid and the anterodorsal portion of the laterosphenoid; unfortunately, neither point of contact is well preserved.

The anterior margin of the frontal is gently convex. The medial portion of this margin abutted the posterior margin of the nasal, whereas the lateral portion was overlapped by the prefrontal. The frontal–nasal contact is practically complete and, as in all macronarians, straight in dorsal view [6,131]. By contrast, the prefrontal articular surface is incomplete both anteriorly and laterally. Towards the lateral margin, the dorsal and ventral surfaces of the frontal host anteromedially–posterolaterally projecting striations, with those on the dorsal surface situated further posteriorly than the same on the ventral surface. The posterior margin of the frontal is convex posteriorly in dorsal view and sinusoidal in posterior view. The medial half of the posterior margin hosts a flattened facet for the parietal; thus, the parietal suture was situated near the anterior margin of the supratemporal fenestra, as in all macronarians [6,143]. The lateral half of the posterior margin either contacted the anterolateral wing of the parietal (if such a structure was present and in contact with the postorbital), or contributed to the supratemporal fenestra, as was inferred for AODF 0836 [38,41]. The medial margin of the frontal is straight and anteroposteriorly convex. No signs of interdigitation or breakage are evident on the medial margin (figure 8h); thus, it seems safe to presume that the frontals were not fused, as in most sauropods [106,148], including AODF 0836 [38,41].

5.6. Parietal

Only the left parietal is present in AODF 0906 (figure 9), and, despite being neither complete nor well-preserved, it is still anatomically informative. Moreover, it is practically identical to the parietals of AODF 0836 [41]. The parietal is broader mediolaterally than tall dorsoventrally, and is slightly tapered

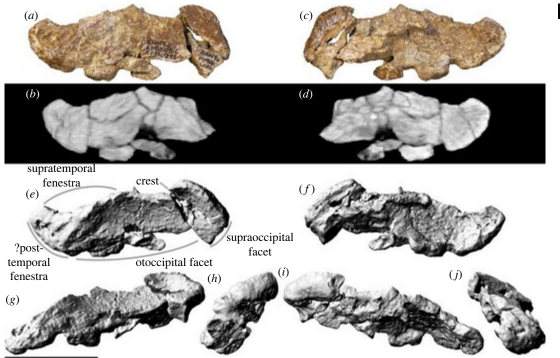


Figure 9. Diamantinasaurus matildae referred left parietal (AODF 0906). (a-d) Left parietal in (a,b) posterior and (c,d) anterior views. (a) and (c) are photographs, (b) and (d) are three-dimensional models derived from CT data. (e-j) Three-dimensional models (derived from surface scans) of the left parietal in (e) posterior, (f) anterior, (g) ventral, (h) medial, (i) dorsal and (g) lateral views. Scale bar = 50 mm.

laterally. The dorsoventral height of the parietal occipital process is greater than the maximum diameter of the foramen magnum, similar to Giraffatitan and Sarmientosaurus, but differentiating it from Abydosaurus, Europasaurus and titanosaurs such as Antarctosaurus and Tapuiasaurus [90,101,104]. As in other titanosaurs, the anterior margin is elevated such that it forms a crest [3,17,38]. The medial two-thirds of the posterior surface is shallowly concave (figure 9a,e), with a subtle ridge demarcating the medial and lateral margins (and presumably, when complete, the anterior margin). The lateral portion, by contrast, is shallowly convex dorsoventrally. Thus, AODF 0906 is similar to the parietals of AODF 0836, which were autapomorphically characterized as having their medial half concave and their lateral half convex [41]. The medial three-quarters of the dorsal margin of the parietal are dorsally convex in posterior view, whereas the lateral one-quarter (which would have formed the posterior margin of the supratemporal fenestra) is concave in posterior view. Unfortunately, the contact with the frontal has suffered extensive damage; thus, it is not clear if this contact occupied a deep trough, as in several other titanosaurs (e.g. Isisaurus colberti, Nemegtosaurus, Rapetosaurus, Saltasaurus and Tapuiasaurus [3,6,38]). The medial surface of the parietal, which appears to present finished bone, is still partially obscured by matrix. Thus, the nature of the inter-parietal contact is unclear. Whether or not a post-temporal fenestra was present in Diamantinasaurus is uncertain, although if there was one, the parietal would have contributed to its margin.

5.7. Postorbital

The right postorbital (figure 10a–l,y,z) is more complete than the left element (figure 10m–x, aa–ab); thus, the description below is based mainly on the former, with supplementary observations derived from the latter. In lateral view (figure 10e,f) the postorbital is triradiate, with a short anteromedial process, an even shorter posterior (squamosal) process, and an elongate anteroventral (jugal) process. The postorbital forms the posterodorsal margin of the orbit, the anterodorsal margin of the infratemporal fenestra, and the anterolateral margin of the supratemporal fenestra. The lateral surface of the postorbital bears a prominent nutrient foramen near the junction of the squamosal and jugal processes; the postorbitals of *Abydosaurus* and *Sarmientosaurus* possess a nutrient foramen in the same position [23,26]. Otherwise, the lateral surface of the postorbital of AODF 0906 is smoothly convex in all directions, albeit only shallowly so on the jugal process. By contrast, the anterior surface is mediolaterally convex and

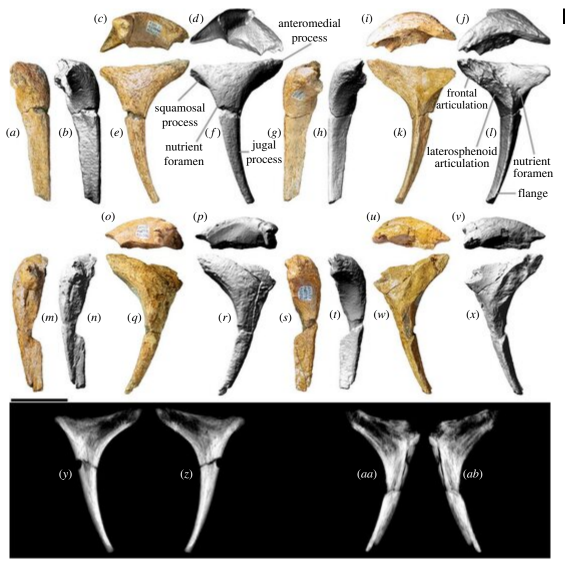


Figure 10. Diamantinasaurus matildae referred postorbitals (AODF 0906). (a-l) Right postorbital in (a,b) posterior, (c,d) ventral, (e,f) lateral, (g,h) anterior, (i,j) dorsal and (k,l) medial views. (a), (c), (e), (g), (i) and (k) are photographs, (b), (d), (f), (h), (j) and (l) are three-dimensional models derived from surface scans. (m-x) Left postorbital in (m,n) posterior, (o,p) ventral, (q,r) lateral, (s,t) anterior, (u,v) dorsal and (w,x) medial views. (m), (o), (q), (s), (u) and (w) are photographs, (n), (p), (r), (t), (v) and (x) are three-dimensional models derived from surface scans. (y,z) Three-dimensional digital models (derived from medical CT data) of the right postorbital in (y) lateral and (z) medial views. (aa-ab) Three-dimensional digital models (derived from medical CT data) of the left postorbital in (aa) lateral and (ab) medial views. Scale bar = 50 mm.

dorsoventrally concave (figure 10g,h), whereas the posterior surface (which would have articulated with the postorbital process of the jugal) is mediolaterally and (shallowly) dorsoventrally concave (figure 10a,b). The jugal process as a whole is anteroposteriorly compressed, as in macronarians generally [8,95]. In medial view (figure 10k,l), the anterior and posterior surfaces of the jugal process are divided by a flange that dissipates dorsally at the laterosphenoid articulation. This anteroposteriorly elongate, shallowly concave facet is more extensive anteroposteriorly in *Diamantinasaurus* than in *Giraffatitan* [25], although it is similar in extent to that of *Europasaurus* [119], *Euhelopus* [114] and *Sarmientosaurus* [23]. Anterior to the laterosphenoid facet lies the frontal facet, which is more strongly concave and faces anteromedially, rather than just medially. Dorsal to these facets, the dorsomedial surface (which would have formed the lateral margin of the supratemporal fenestra) is shallowly concave. The path of the supratemporal canal runs posterior to the laterosphenoid facet. On the medial surface, immediately posterior to the laterosphenoid facet, a small nutrient foramen is present. The squamosal process is broken in both postorbitals, but was clearly triangular and fairly short, albeit not reduced to the same extent as seen in *Antarctosaurus*, *Nemegtosaurus*, *Quaesitosaurus* and *Tapuiasaurus* [20–22,104].

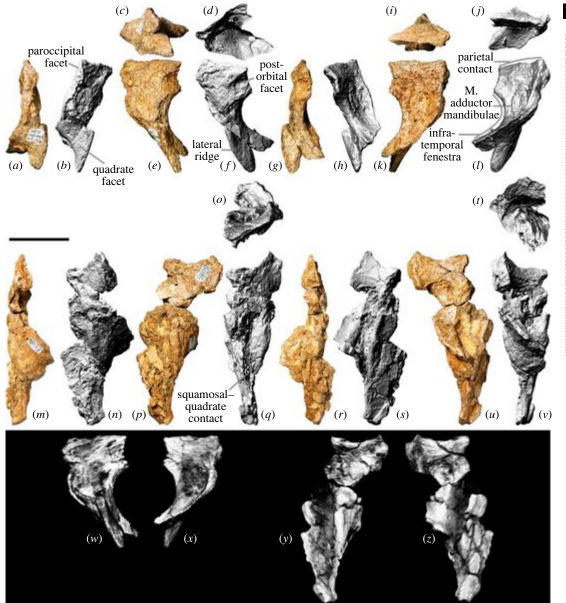


Figure 11. Diamantinasaurus matildae referred squamosals (AODF 0906). (a-l) Right squamosal in (a,b) posterior, (c,d) ventral, (e,f) lateral, (g,h) anterior, (i,j) dorsal and (k,l) medial views. (a), (c), (e), (g), (i) and (k) are photographs, (b), (d), (f), (h), (j) and (l) are three-dimensional models derived from surface scans. (m-v) Left squamosal (with attached left quadrate fragment) in (m,n) posterior, (o) ventral, (p,q) lateral, (r,s) anterior, (t) dorsal and (u,v) medial views. (m), (p), (r) and (u) are photographs, (n-o), (q), (s-t) and (v) are three-dimensional models derived from surface scans. (w,x) Three-dimensional digital models (derived from medical CT data) of the right squamosal in (w) lateral and (x) medial views. (y,z) Three-dimensional digital models (derived from medical CT data) of the left squamosal in (y) lateral and (z) medial views. Scale bar = 50 mm.

5.8. Squamosal

The right squamosal (figure 11a-l,w-x) is more complete and better-preserved than the left one (figure 11m-v,y-z). The latter is incompletely exposed, remaining in contact, albeit not in articulation, with a fragment of the dorsal section of the pterygoid process of the left quadrate. Consequently, the description below is based almost entirely on the right squamosal.

The squamosal was restricted to the postorbital region, distinguishing it from *Nemegtosaurus*, *Rapetosaurus* and *Tapuiasaurus*, wherein the squamosal extends beyond the posterior margin of the orbit [6]. Broadly speaking, the squamosal is comma-shaped in lateral view (figure 11e,f). The dorsal margin is manifested as a mediolaterally thin, shallowly anteroposteriorly concave ridge that terminates posteriorly at a small but pronounced point (figure 11i,j). The lateral surface of the short anterior process bears a

posteriorly tapered facet to receive the posterior process of the postorbital. Posterior to the postorbital facet, the lateral surface is smoothly convex. Ventral to this facet, a prominent lateral ridge is present, which divides the anteroventral process into anterior and posterior surfaces. The anterior surface of the anteroventral process (figure 11g,h)), which is shallowly concave, forms the posterodorsal margin of the infratemporal fenestra. The mostly flattened posterior surface of the anteroventral process hosts the articular facet for the quadrate (figure 11a,b), which is not well preserved in either squamosal of AODF 0906; nevertheless, it appears to be bilobate, as is also the case in AODF 0836 [38,41] and Sarmientosaurus, as observed in CT data [23]. Although the ventral processes of both squamosals are incomplete in AODF 0906, it is presumed that the ventral tip of the anteroventral process of the squamosal contacted the dorsal process of the quadratojugal, as in most macronarians [95,100,106,132].

The dorsomedial surface of the squamosal would have articulated with the posterolateral margin of the parietal. It seems likely that the squamosal of *Diamantinasaurus* contributed to the supratemporal fenestra, contrasting with the titanosaurs *Nemegtosaurus*, *Quaesitosaurus* and *Tapuiasaurus* [20,22,100,106]. A subtle horizontal ridge separates the relatively small parietal facet from the rest of the medial surface of the squamosal, which is strongly concave and would have accommodated the *M. adductor mandibulae*. The posterior margin of the squamosal is expressed as a thin, anteroventrally–posterodorsally oriented ridge for most of its length, although the dorsal one-third broadens into a flat (albeit still mediolaterally narrow) facet against which the paroccipital process abutted. The ventral prong situated ventrolateral to the paroccipital articulation in the squamosals of the non-titanosaurian somphospondylan *Phuwiangosaurus* and the titanosaurs *Nemegtosaurus* and *Tapuiasaurus* is not evident in AODF 0906 and seems to be genuinely absent [17,20,22,149].

5.9. Quadratojugal

The left quadratojugal of AODF 0906 (figure 12*a*–*l*) is more complete than the right (figure 12*m*–*x*), and it forms the basis for the entire description of this element herein. The quadratojugal is approximately L-shaped in lateral view (figure 12*f*–*h*), with an elongate jugal (anterior) process and an apparently shorter quadrate (dorsal) process. The latter process is not complete in either quadratojugal, but it was likely sufficiently elongate to contact the anteroventral process of the squamosal. In both quadratojugals, the jugal process also appears to be incomplete; nevertheless, it can safely be presumed that the jugal process was more than 1.3 times the length of the quadrate process, as in all macronarians [95,100]. It seems unlikely that the anterior extremity of the jugal process of the quadratojugal possessed a ventral triangular projection, differentiating *Diamantinasaurus* from brachiosaurids, *Europasaurus*, *Nemegtosaurus*, *Quaesitosaurus* and *Tapuiasaurus* [5,38]. The angle between the quadrate and jugal processes is less than 90° in AODF 0906. In the titanosaur *Sarmientosaurus*, and in non-titanosaurian macronarians generally, these processes meet at an angle of 90° or lower [6,23,106,132]; by contrast, the angle between these processes is greater than 90° in *Nemegtosaurus* and *Tapuiasaurus*, as well as the Auca Mahuevo titanosaur embryos [20,22,128].

The lateral surface of the jugal process of the quadratojugal is entirely convex, whereas that of the base of the squamosal process, which is distinctly twisted and thickened mediolaterally, is shallowly concave. The medial surface of the jugal process is anteroposteriorly convex but dorsoventrally concave, and it terminates posteriorly in a ridge that demarcates the anterior margin of the quadrate facet (figure 12j-l). At approximately two-thirds of the preserved height of the quadrate facet, a subtle horizontal ridge is present, which is aligned with a similar ridge on the quadrate when the elements are articulated. The quadrate facet of the quadratojugal is strongly concave, with two small pits present near the posterior margin. The quadratojugal lacks the posterioventral hook seen in *Tapuiasaurus* [22]. However, a posterior tongue-like process, situated posterior to the quadrate facet, is present. A similar process is present in *Sarmientosaurus* and was regarded as an autapomorphy of that taxon by Martínez *et al.* [23]. To test its distribution, we include its presence as a new character (C555).

5.10. Quadrate

The left quadrate (figure 13*a*–*m*) is almost complete but quite damaged, whereas the right quadrate (figure 13*n*–*s*) is only represented by a small, albeit far less distorted, ventral portion of the body. Consequently, the description below is based entirely on the left quadrate, which is virtually indistinguishable from the quadrates of AODF 0836 [41].

When complete, the quadrate would have been triangular in lateral and medial views, with a prominent anterior pterygoid flange. The quadrate was evidently oriented more or less vertically, as in

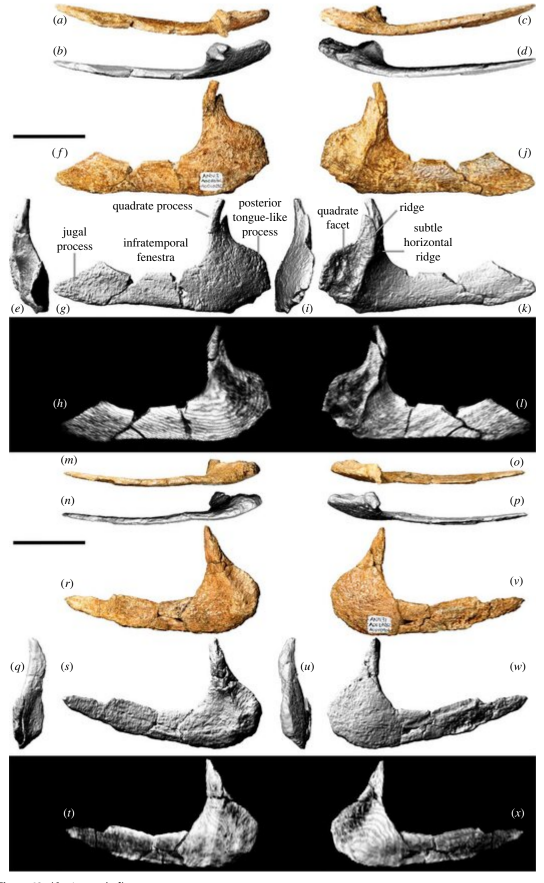


Figure 12. (Caption overleaf.)

Figure 12. (Overleaf.) Diamantinasaurus matildae referred quadratojugals (AODF 0906). (a–l) Left quadratojugal in (a,b) dorsal, (c,d) ventral, (e) anterior, (f–h) lateral, (i) posterior and (j–l) medial views. (a), (c), (f) and (j) are photographs, (b), (d–e), (g), (i) and (k) are three-dimensional models derived from Surface scans, and (h) and (h) are three-dimensional models derived from CT data. (h-h) Right quadratojugal in (h-h) ventral, (h-h) dorsal, (h-h) anterior, (h-h) medial, (h) anterior, and (h-h-h) lateral views. (h-h), (h-h) are photographs, (h), (h-h-h), (h-h) and (h) are three-dimensional models derived from CT data. Scale bar = 50 mm.

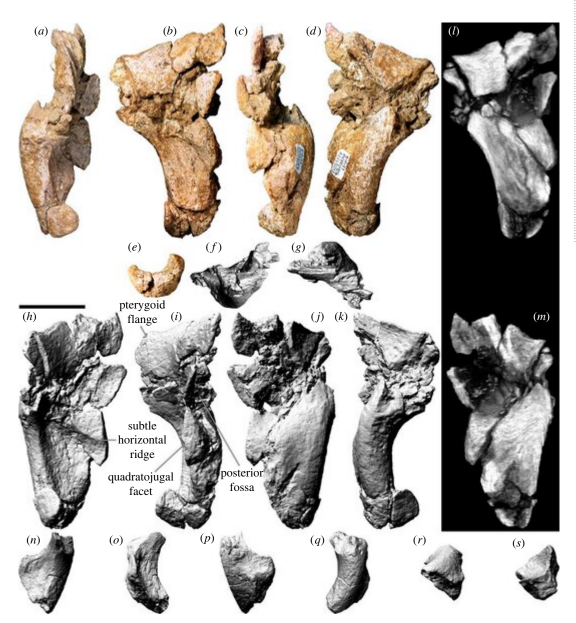


Figure 13. Diamantinasaurus matildae referred quadrates (AODF 0906). (a-e) Photographs of the left quadrate in (a) anterior, (b) lateral, (c) posterior, (d) medial and (e) ventral views. (f-k) Three-dimensional models (derived from surface scans) of the left quadrate in (f) ventral, (g) dorsal, (h) anterior, (i) lateral, (j) posterior and (k) medial views. (l,m) Three-dimensional models derived from CT scans of the left quadrate in (l) lateral and (m) medial views. (n-s) Three-dimensional models (derived from surface scans) of the right quadrate in (n) anterior, (o) lateral, (p) posterior, (q) medial, (r) dorsal and (s) ventral views. Scale bar = 50 mm.

some titanosaurs, such as *Sarmientosaurus*, but unlike taxa such as *Tapuiasaurus* [22,23]. Although the left quadrate of AODF 0906 is broken, the posterior surface is clearly characterized by a deep excavation (figure 13c,i,j), as in all macronarians [8,95]. In AODF 0906, this fossa faces posterolaterally, distinguishing it from most sauropods, with the exception of several titanosaurs, including AODF 0836, *Nemegtosaurus*, *Rapetosaurus*, *Sarmientosaurus* and *Tapuiasaurus* [17,20,22,23,41,104]. It seems

unlikely that the quadrate contacted the basal tuber, distinguishing AODF 0906 from *Nemegtosaurus*, *Quaesitosaurus*, *Rapetosaurus* and *Tapuiasaurus* [17,20,22,104].

The head of the quadrate, and therefore the squamosal contact, is largely missing. However, the articular surface for the quadratojugal is complete and well-preserved, such that a subtle horizontal ridge on the quadrate (figure 13h) is contiguous with a similar horizontal ridge on the quadratojugal when the two elements are articulated. A comparable ridge appears to be present on each quadrate of AODF 0836, although this feature was not identified in the original description of that specimen [41]. We consider this feature to be autapomorphic for *Diamantinasaurus matildae*. Ventral to the anterior flange, the lateral surface of the quadrate is strongly concave (figure 13b,i), whereas the medial surface is correspondingly strongly convex (figure 13d,k). The ventral process of the quadrate is expanded mediolaterally and laterally bevelled at a 45° angle. The ventral articular facet, in distal view, is crescentic. As in AODF 0836, the concave margin of the facet faces anterolaterally, whereas the convex one faces posteromedially [41].

5.11. Pterygoid

The left pterygoid (figure 14a-h,q) is represented by incomplete palatine and quadrate processes, whereas the right pterygoid (figure 14i-p,r,s) preserves nearly complete ectopterygoid and quadrate processes. Between the two elements, an almost complete pterygoid can be reconstructed, and morphologically it appears to be similar to those of Europasaurus [119], Giraffatitan [25] and Sarmientosaurus [23]. When complete, the pterygoid would have been triradiate, with the quadrate process directed posterolaterally, the palatine process anteriorly, and the ectopterygoid process ventrolaterally. Thus, the pterygoid of AODF 0906 was fairly robust, unlike those of Nemegtosaurus, Quaesitosaurus and Rapetosaurus, wherein the three processes are more or less coplanar [17,20,38]. The preserved portion of the palatine process (figure 14e,f) is sheet-like, being exceedingly thin mediolaterally and tall dorsoventrally. It is otherwise uninformative, although it appears to preserve bite marks (figure 14a,b). The distal end of the ectopterygoid process is mediolaterally thickened on its anterior surface (figure 141) and tapers to a thin ridge posteriorly (figure 14p). This ridge is confluent with the ventral margin of the quadrate process, which is similarly mediolaterally thin along its ventral surface (figure 14g,q). In anterior view, the distal end of the ectopterygoid process is mostly flat. The medial surface of the ectopterygoid process is similarly flattened (figure 14i.j), whereas its lateral surface is mostly convex (figure 14m.n). The exception to this is a small sulcus immediately proximal to the end of the process, situated at the ventrolateral margin. Whether or not this is a natural feature is unclear. As is the case in most sauropods for which this can be assessed, the articular surface for the ectopterygoid appears to extend along the dorsal half of the lateral surface of the ectopterygoid process; this distinguishes AODF 0906 from Rapetosaurus, Sarmientosaurus and Tapuiasaurus, wherein the articular surface for the ectopterygoid is restricted to the tip of this process [21,23,38]. The lateral surface of the quadrate process is mostly dorsoventrally convex, with a shallow, horizontal rise at one-quarter of the height of the process. This implies that the medial surface of the pterygoid flange of the quadrate, which is incompletely preserved, was dorsoventrally concave. The medial surface of the quadrate process of the pterygoid is more complex (figure 14ef, i,j): it is bisected by a prominent, medially projecting and anterodorsally-posteroventrally inclined ledge. The articular surface for the basipterygoid process is manifested as a socket in AODF 0906, distinguishing it from Nemegtosaurus, Quaesitosaurus, Rapetosaurus and Tapuiasaurus, wherein it forms a convex rocker-like surface [17,20,22,104]. Anteroventral to the ledge, the pterygoid is vaulted to form the posterior portion of the palate.

5.12. Ectopterygoid

The left ectopterygoid is virtually complete (figure 15), although the maxillary articular surface is incompletely exposed and appears to be broken. In anterior and posterior views, the ectopterygoid is ligulate (strap-shaped), being somewhat flared medially. The maxillary articular surface is circular and mostly concave, albeit partially bisected by a bulge. As in all neosauropods, the ectopterygoid did not articulate with the jugal [106]. Medial to the maxillary facet, the body of the ectopterygoid is tubular. Further medially, however, it flattens out to form a dorsoventrally expanded flange. This flange is biased posteriorly, such that the posterior surface of the ectopterygoid is essentially a mildly dorsoventrally convex sheet. The anterior surface is more strongly convex dorsoventrally than the posterior one, with a distinct dorsal shelf immediately medial to the maxillary articular facet. The medial flange is presumed to have articulated with the ectopterygoid process of the pterygoid posteriorly and the palatine anteriorly.

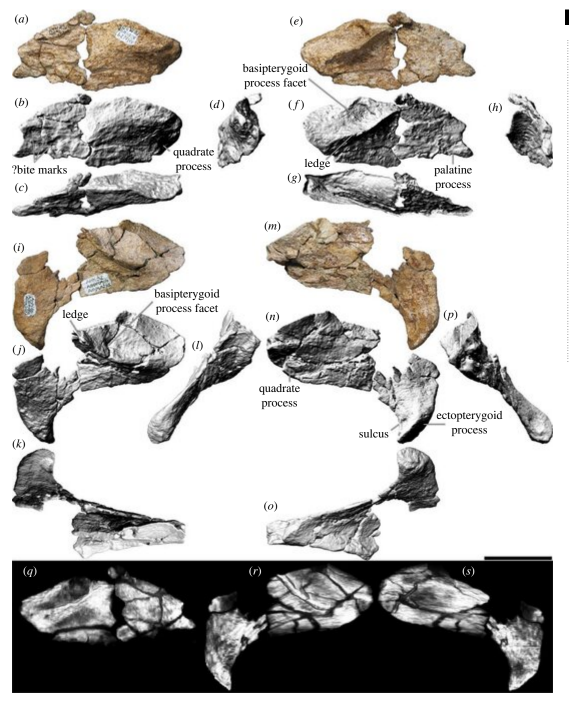


Figure 14. Diamantinasaurus matildae referred pterygoids (AODF 0906). (a-h) Left pterygoid in (a,b) medial, (c) dorsal, (d) posterior, (e,f) medial, (g) ventral and (h) anterior views. (i-p) Right pterygoid in (i), (j) medial, (k) dorsal, (l) anterior, (m,n) lateral, (o) ventral and (p) posterior views. (a), (e), (i) and (m) are photographs, and (b-d), (f-h), (j-l) and (n-p) are three-dimensional models derived from surface scans. (q) Three-dimensional model (derived from CT scans) of the left pterygoid in medial view. (r,s) Three-dimensional models (derived from CT scans) of the right pterygoid in (r) medial and (s) lateral views. Scale bar = 50 mm.

5.13. Braincase

The braincase is poorly preserved and in two sections (figure 16). The first comprises an incomplete supraoccipital, partial otoccipitals, fragmentary prootics, laterosphenoids and orbitosphenoids. The second comprises a partial basioccipital and basisphenoid. The photographs (figure $16a_ie_jh_ik_i,n$ and w), line drawings (figure $16g_ij_i,m$ and p), and CT slices (figure $16q_i-v$) of the braincase presented herein only illustrate the first fragment, whereas the screenshots of three-dimensional models derived from surface scans (figure $16b_i-d_i$, f_i , f_i , g_i and g_i) illustrate both fragments in approximate life position.

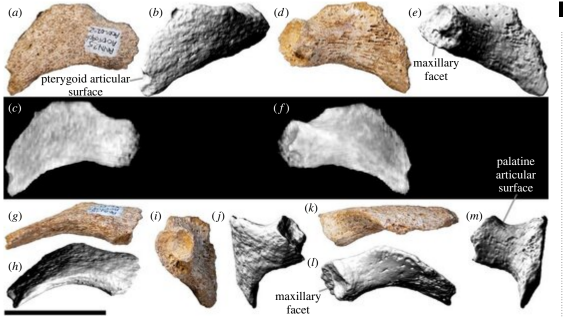


Figure 15. Diamantinasaurus matildae referred left ectopterygoid (AODF 0906). (a-m) Left ectopterygoid in (a-c) anterior, (d-f) posterior, (g,h) ventral, (i,j) lateral, (k,l) dorsal and (m) posterior views. (a), (d), (g), (i) and (k) are photographs; (b), (e), (h), (j) and (l-m) are three-dimensional models derived from surface scans; and (c) and (f) are three-dimensional models derived from CT scans. Scale bar = 50 mm.

5.13.1. Supraoccipital

Despite being incomplete, the supraoccipital is one of the better-preserved braincase elements. This subhexagonal element is slightly taller dorsoventrally (greater than 32 mm) than it is wide transversely (28 mm) and forms the dorsal margin of the foramen magnum (figure 16k-m). As is also the case in AODF 0836 and most other sauropods, the supraoccipital is slightly taller dorsoventrally (approx. 38 mm) than the foramen magnum, which contrasts with the titanosaurs Malawisaurus, Pitekunsaurus macayai and Rapetosaurus [101,104]. The nuchal crest, which runs dorsoventrally along the midline of the supraoccipital, is prominent anteriorly but fades out well before reaching the level of the foramen magnum; consequently, the 20 mm tall portion of the supraoccipital dorsal to the foramen magnum is almost entirely flat. Only the left half to two-thirds of the nuchal crest is preserved, and it is difficult to determine whether or not an anteroposterior groove was present on its dorsal surface. A hint of such a groove is preserved, although this might be an artefact of preservation and is situated too far to the left to be considered 'midline'. Although it is possible that there were two grooves on the nuchal crest, situated either side of the midline, this would be a highly unusual morphology for a sauropod, let alone a titanosauriform [150]. On both sides, lateral to the nuchal crest, the supraoccipital hosts a shallowly concave occipital fossa; however, both fossae might appear shallower than they were in vivo owing to their incompleteness.

5.13.2. Basioccipital

The basioccipital is incomplete: the occipital condyle has been lost, and the basal tubera are extremely fragmentary. Consequently, much of the morphology of the basioccipital cannot be determined, other than that it was sutured dorsolaterally with the otoccipital, dorsally with the prootic, and anteriorly with the basisphenoid.

5.13.3. Basisphenoid

The basisphenoid forms the floor of the endocranial cavity and is firmly sutured to the (missing) parasphenoid anteriorly, the orbitosphenoid anterodorsally, the laterosphenoid dorsally, the prootic posterodorsally and the basioccipital posteriorly. The basisphenoid forms the basipterygoid processes

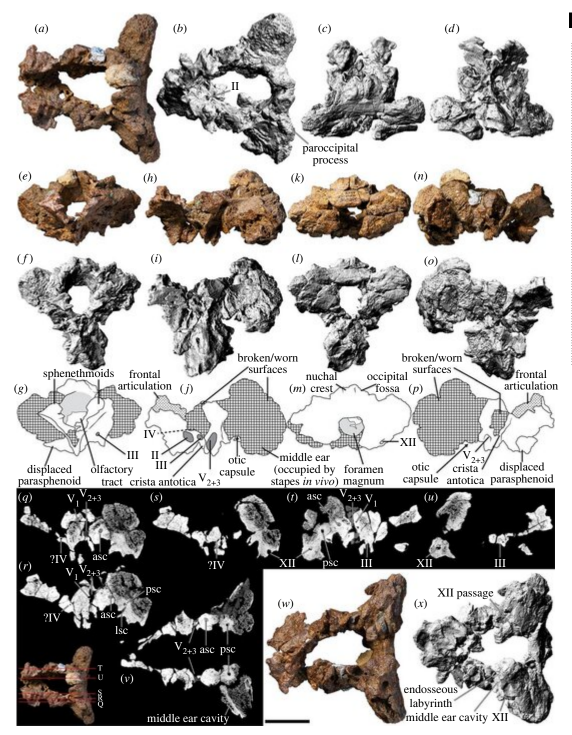


Figure 16. Diamantinasaurus matildae referred braincase (AODF 0906). (a-p) Braincase in (a,b) dorsal, (c) posterodorsal, (d) anteroventral, (e-g) anterior, (h-j) left lateral, (k-m) posterior and (n-p) right lateral views. (q-u) Braincase cross-sections in (q-s) left lateral, (t,u) right lateral and (v) dorsal views. (w-x) Braincase in ventral view. (a), (e), (h), (k), (n) and (w) are photographs; (b-d), (f), (i), (i), (o) and (x) are three-dimensional models derived from surface scans; and (q-v) are CT slices. Scale bar = 50 mm.

and a significant portion of the basal tubera. Unfortunately, both the basipterygoid processes and basal tubera of AODF 0906 are quite incomplete.

The hypophyseal extension is visible in dorsal view (figure 16b). The boundaries of this structure define an oval shape, lateral to which the paths of the carotid arteries can be traced. Although both are mostly open, this is almost certainly a consequence of breakage since finished bone is not evident throughout much of the length of the more complete left carotid artery canal. It seems likely that the carotid arteries opened lateral to the basipterygoid processes, as in most sauropods, including AODF 0836 [41] and *Sarmientosaurus* [23], but

contrasting with the condition in several titanosaurs (e.g. Antarctosaurus, Nemegtosaurus, Rapetosaurus, Saltasaurus), as well as the early-branching somphospondylan Tambatitanis amicitiae [151] and possibly Mongolosaurus haplodon ([152]: fig. 2), in which the carotid arteries exit medial to the basipterygoid processes [38,153].

5.13.4. Otoccipital

As in sauropods generally, the exoccipital and opisthotic are firmly sutured [8]; consequently, they are referred to herein as the otoccipital. The otoccipital is dorsomedially sutured to the supraoccipital, ventromedially to the basioccipital, anteroventrally to the prootic, and anteromedially to the laterosphenoid. The medial edge of the otoccipital forms the lateral margin of the foramen magnum. Immediately lateral to the foramen magnum lies the external opening for the hypoglossal nerve (CN XII). A single exit for CN XII is present on each side, as in most macronarians [38,153], including AODF 0836 [38,41]. This contrasts with Sarmientosaurus [23], as well as some specimens of Camarasaurus and Giraffatitan, which have two openings for CN XII on each side [153]. In AODF 0906, the left CN XII opening is broken, such that its passage from the endocast to the anterolateral surface of the occipital condyle is entirely exposed. By contrast, the right CN XII canal is intact, such that only its internal and external openings can be observed. Lateral to the left side of the foramen magnum, and dorsal to the external opening for CN XII, a node-like proatlantal facet appears to be present; a less well-preserved one appears to be present on the right side as well. Proatlantal facets have also been identified in AODF 0836 [41] and characterize most titanosaurs, although they are absent in Malawisaurus, Rapetosaurus and Sarmientosaurus [23,38,41]. When complete, the otoccipitals would have formed prominent paroccipital processes laterally; however, both are incomplete laterally, and the morphology of the paroccipital processes is unclear.

5.13.5. Prootic

Both prootics are incomplete in AODF 0906. Each prootic is firmly wedged between the laterosphenoid anteriorly, the parietal dorsally, the supraoccipital posteriorly, the opisthotic portion of the exoccipital-opisthotic complex posteroventrally, and the basisphenoid and basioccipital ventrally. The crista antotica marks the boundary between the laterosphenoid and prootic. Immediately posterior to this ridge is a foramen that would have hosted the maxillary (CN V_2) and mandibular (CN V_3) branches of the trigeminal nerve (CN V). This section of the prootic is poorly preserved on both sides of AODF 0906; nevertheless, it is clear that the morphology of this region is extremely similar to that of AODF 0836 [38,41]. Little evidence of the crista prootica is preserved on either prootic. Consequently, the precise position of the opening for the facial nerve (CN VII) is difficult to determine.

Neither prootic preserves its parietal or basisphenoid suture, meaning that the crista tuberalis is not preserved. By contrast, the crista interfenestralis is preserved on both sides. In AODF 0836, this feature was interpreted as a preservational artefact [38,41]; however, the fact that this structure in AODF 0906 occupies exactly the same position and has a near-identical morphology implies that it is a genuine anatomical feature. Posterior to the crista interfenestralis lies the metotic fissure. Anterior to the crista interfenestralis, the dorsal surface of a straight, narrow, posterolaterally–anteromedially oriented canal—the middle ear canal (or tympanic cavity)—can be observed: *in vivo*, this would have been occupied by the stapes (figure 16w-x). Medial to this canal, the matrix infilling the cochlear portions of both endosseous labyrinths has been removed, approximately to the level of the lateral semicircular canal; it is also possible to discern the approximate paths of the anterior and posterior semicircular canals. CT data reveal that the anterior semicircular canal is slightly larger in diameter than the posterior one (figure 16q,r, t and v). The anterior and posterior semicircular canals diverge from each other at approximately 100° , as in AODF 0836 [41].

5.13.6. Laterosphenoid

The anterolaterally concave laterosphenoid forms the posteromedial rim of the orbital fossa. Neither the dorsal suture with the frontal, nor the lateral suture with the postorbital, can be observed; by contrast, the completely preserved anterior suture with the orbitosphenoid has been largely obliterated. The orbitosphenoid suture line of the laterosphenoid is punctuated by the openings for the oculomotor (CN III), trochlear (CN IV) and abducens (CN VI) nerves, as in sauropods generally [154]. The laterosphenoid is sutured to the opisthotic portion of the otoccipital posteriorly, and to the prootic

posteroventrally. The contact between the laterosphenoid and otoccipital/prootic is manifested as the crista antotica. The ophthalmic branch (CN V_1) of the trigeminal nerve (CN V) lies entirely on the laterosphenoid, anterior to the crista antotica; the other two branches of the trigeminal nerve exited posterior to the crista antotica, as in AODF 0836 [38,41]. Among neosauropods, only *Phuwiangosaurus*, *Diamantinasaurus* and *Sarmientosaurus* are characterized by more than one ossified exit for CN V [23,38,41,82,149].

5.13.7. Orbitosphenoid

Both orbitosphenoids are incomplete. They are firmly sutured together along their ventral margins, and each is sutured posteriorly to its corresponding laterosphenoid. The frontal articulation, which would have been situated dorsally, is indistinct on both orbitosphenoids; however, based on the left side, it would seem that this suture was interdigitated, as in AODF 0836 [38,41]. The suture with the basisphenoid is poorly preserved on both.

The opening for the olfactory nerve (CN I) is situated anteriorly and is generally well preserved. A medially projecting prong is present within this opening on the right orbitosphenoid; a similar structure is present in the left orbitosphenoid of AODF 0836, and tentatively interpreted to represent the boundaries of the olfactory filaments [38,41]. One of few other titanosaurs in which the anteriormost portion of the orbitosphenoid is preserved, and in which the braincase was found disarticulated from the rest of the skull, is Bonatitan reigi: in this taxon, ethmoidal elements (specifically sphenethmoids) were inferred to be preserved anterior to, and in articulation with, the orbitosphenoids [155]. If this interpretation is correct, then at least one sphenethmoid appears to be preserved in AODF 0836, and both appear to be preserved in AODF 0906. The sphenethmoid would then be the element that hosts the medially projecting prong described above. The opening for the optic nerve (CN II) is hosted on the orbitosphenoid and is medially divided, as in nearly all neosauropods [156,157]. The opening for CN III is positioned immediately posterior to that for CN II, whereas the section that would have hosted CN IV (posterodorsal to CN II) is incomplete on both sides (although its passage can possibly be traced using the CT data, at least on the left side (figure 16q,r)). The passage for CN VI appears to be preserved on the left side (figure 16s). As in AODF 0836 [38,41], each CN VI canal projects lateral to the hypophyseal chamber and does not penetrate the pituitary fossa, meaning that Diamantinasaurus shows the derived condition that characterizes all titanosaurs [6,38,153]. In AODF 0906, as in many titanosaurs (including AODF 0836), the opening for CN VI is interpreted to have been situated anteroventral to, and quite distant from, the opening for CN III; by contrast, in non-titanosaurian macronarians, the external opening for CN VI lies ventral (and often quite close) to that for CN III [6,38,41,158].

5.14. Dentary

The left dentary is nearly complete and anteroposteriorly elongate (figure 17*a*–*l*); by contrast, the right dentary is represented only by a fragment (figure 14*m*–*t*). The description below is based on the left dentary unless otherwise indicated.

The dorsoventral height of the anterior end of the dentary (51 mm) is approximately equivalent to its height at mid-length (49 mm), as in most titanosauriforms, other than *Camarasaurus*, brachiosaurids and *Malawisaurus*, wherein the anterior end of the dentary is 120% taller than the mid-length [6,95,100,104]. As preserved, the anterior margin of the dentary is angled posteroventrally (figure 17f–h), as is the case in most sauropods [38,100], including *Camarasaurus* [118], *Europasaurus* [119], *Euhelopus* [114], and the titanosaurs *Malawisaurus* [115] and *Choconsaurus baileywillisi* [159]. This distinguishes AODF 0906 from brachiosaurids and most titanosaurs, including *Mansourasaurus shahinae*, *Nemegtosaurus*, *Rapetosaurus*, *Sarmientosaurus* and *Tapuiasaurus*, wherein the anterior margin is perpendicular (or nearly so) to the long axis of the dentary [17,20,22,23,26,160]. The 'chin' observed in *Mansourasaurus* [160] and many flagellicaudatans [100,161] is not present in AODF 0906, nor is there a labial tuberosity near the anterior end of the dentary, such as that which characterizes dicraeosaurids [131].

The paired dentaries would have formed a rounded lower jaw in occlusal view (figure 17a-c), as in non-titanosauriform macronarians, brachiosaurids, *Euhelopus*, and numerous titanosaurs, including *Ampelosaurus atacis*, *Choconsaurus*, *Karongasaurus gittelmani*, *Malawisaurus*, *Mansourasaurus*, *Nemegtosaurus*, *Quaesitosaurus*, *Rapetosaurus*, *Sarmientosaurus* and *Tapuiasaurus* [2]. This is in stark contrast to the condition in some other titanosaurs, such as *Antarctosaurus*, *Baalsaurus mansillai*, *Bonitasaura* and *Brasilotitan nemophagus*, wherein the dentaries are squared-off in occlusal view [141,145,162–165].

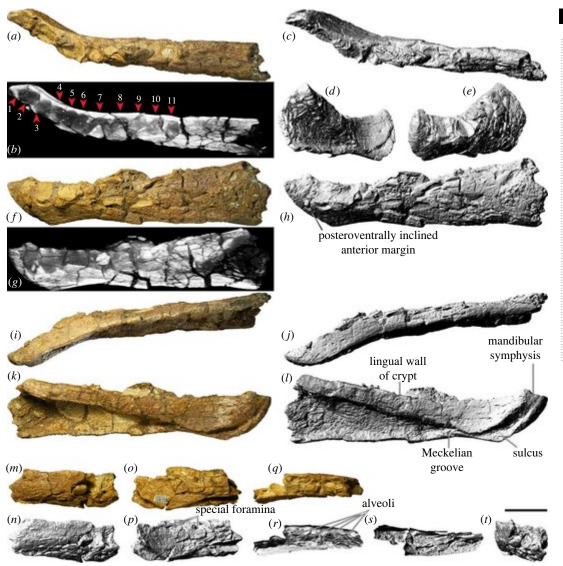


Figure 17. Diamantinasaurus matildae referred dentaries (AODF 0906). (a-l) Left dentary in (a-c) occlusal, (d) posterior, (e) anterior, (f-h) lateral, (i-j) ventral and (k-l) medial views. (a), (f), (i) and (k) are photographs; (b) and (g) are three-dimensional models derived from CT scans; and (c-e), (h), (j) and (l) are three-dimensional models derived from surface scans. Numbers 1–11 indicate alveoli. (m-t) Right dentary in (m-n) lateral, (o-p) medial, (q-r) occlusal, (s) ventral and (t) posterior views. (m), (o) and (q) are photographs, (n), (p) and (r-t) are three-dimensional models derived from surface scans. Scale bar = 50 mm.

The lateral surface of the left dentary (figure 17*a*–*c*), which is dorsoventrally convex along its length, has suffered extensive fragmentation. As a result, the tooth row appears to have been bowed outwards relative to the rest of the dentary, as in dicraeosaurids, *Nigersaurus taqueti* [166], as well as *Antarctosaurus* [90]. However, the preserved portion of the right dentary demonstrates that this was not the case *in vivo*—as in other macronarians, the lingual and labial margins of the alveoli were level, and the tooth row was not bowed either way (figure 17*q*,*r*). Presently, it is not possible to determine if the dentary of AODF 0906 bifurcated into two posterior processes (dorsal and ventral), nor if the posteroventral process was forked (as it is in brachiosaurids [26]).

The medial surface of the dentary (figure 17k,l) can be divided into four surfaces. The anteriormost and least extensive is the flattened articular surface for the opposite dentary. Immediately ventral to the mandibular symphysis, a distinct sulcus is present. Although it is separated from the anterior end of the Meckelian groove by a region wherein the lateral and medial surfaces meet directly (approx. 30 mm long anteroposteriorly), such that they are barely separated by a groove (in the left dentary) or entirely lack a groove (in the right one), it is possible that this feature represents an anterior continuation of the Meckelian groove proper. Among macronarians, this feature appears to be present only in ontogenetically immature individuals of the early-diverging macronarian *Europasaurus* [119]. Thus, this

feature might indicate that the AODF 0906 individual was not fully grown perimortem; alternatively, it might be locally autapomorphic for *Diamantinasaurus* within Titanosauria or even Titanosauriformes. The most extensive of the four medial surfaces of the dentary is the lingual wall of the crypt (within which the replacement teeth were generated and held). This surface is mostly flat, except where it is concave immediately posterior to the dentary articulation. This is also the position at which this surface is dorsoventrally tallest: it gradually decreases in dorsoventral height posteriorly (owing to the Meckelian groove) and anteriorly (owing to the anteroventral sulcus). Both dentaries preserve the ventral margins of several special foramina, although those on the less complete right dentary are more easily observable. Posteroventral to the crypt lies the Meckelian groove, which is dorsoventrally concave, anteriorly tapered and posteriorly expanded. The medial surface of the dentary within the Meckelian groove is concave and would have been overlain by the splenial in life.

CT scan data reveal that the incomplete right dentary of AODF 0906 hosts six alveoli, whereas the complete toothrow of the left dentary comprises 11 alveoli (figure 17a,g). The presence of 11 dentary alveoli separates AODF 0906 from non-titanosaurian titanosauriforms, which typically have 12 to 14 [24–26,114,118,119,133,135,137,167]. It also separates AODF 0906 from several titanosaurs with rounded snouts, including *Nemegtosaurus*, *Quaesitosaurus* and *Sarmientosaurus*, which have 13 [18–20,23]. However, alveolus count in titanosaurs with rounded snouts varies greatly. Some of the lowest counts are those seen in *Ampelosaurus* (nine) and *Mansourasaurus* (10) [160,168]. *Rapetosaurus* possesses at least 11 [17], and *Karongasaurus* at least 12 [115]. By contrast, other titanosaurs with rounded snouts have a greater number of alveoli, with 15 in *Tapuiasaurus* [22] and at least this many in *Malawisaurus* [115]. Similarly, alveolus counts in titanosaurs with squared-off snouts vary, from at least 13 in *Baalsaurus* [165], to 14 in *Brasilotitan* [164] and 16 in *Antarctosaurus* (MACN 6904, S.F.P. and P.D.M., pers. obs.). This places AODF 0906 at the lower end of the expected range of dentary alveoli for Titanosauria, particularly among early-branching titanosaurs.

5.15. Surangular

The left surangular (figure 18) is almost identical to, and more completely preserved than, the surangular of AODF 0836 [41]. It is also broadly similar to the surangulars of Giraffatitan [25] and Sarmientosaurus [23]. The surangular of AODF 0906 is mediolaterally thicker dorsally (figure 18j) than ventrally (figure 18i). The undulating dorsal margin (which is the only complete margin) is shallowly convex anteriorly and shallowly concave posteriorly. The preserved portion of the lateral surface of the surangular (figure 18b,c) is broadly convex, but shallowly concave posterodorsally. This part of the surangular in Tapuiasaurus accommodated the anteroventral prong of the quadratojugal [22]; however, the quadratojugal of AODF 0906 lacks this feature. As in AODF 0836, the dorsal portion of the medial surface of the surangular of AODF 0906 is manifested as a distinct double arch (figure 18f, g). The anterior arch is incomplete anteromedially, whereas the posterior arch is almost complete. Ventral to each arch, the medial surface of the surangular is shallowly concave. The anterior concavity (the adductor fossa) appears to preserve the posterior margin of a fairly large foramen. This appears to be homologous with, albeit smaller than, the anterior surangular foramen present in the surangulars of Nemegtosaurus [18,20] and Rapetosaurus [17]. The surangulars of Abydosaurus, Euhelopus, Europasaurus, Giraffatitan, Quaesitosaurus, Sarmientosaurus and Tapuiasaurus lack an enlarged surangular foramen, instead having a much smaller anterior surangular foramen, situated further anteriorly [19,22,23,25,114,119]. The morphology of this foramen varies in Camarasaurus: an enlarged foramen is present in some exemplars, whereas in others a smaller foramen is evident [118]. CT scan data of Sarmientosaurus reveal that the anterior surangular foramen connects to a posteromedially projecting canal, which expands greatly (diameter approx. 10 mm) within the body of the surangular, and exits on the medial surface near the posterior margin of the adductor fossa [23]—the precise position at which this foramen is located in AODF 0906 (and, evidently, in AODF 0836). It is likely, then, that these foramina are homologous. In living reptiles, the anterior and posterior surangular foramina accommodate cutaneous branches of the inferior alveolar nerve [169,170].

5.16. Teeth

No loose teeth were found at the AODL 0252 site, and no active teeth remained within the preserved dentigerous elements. Computed tomography (CT) scan data revealed that the premaxilla and maxilla preserve replacement teeth, but that the dentaries do not. In the maxilla, the few preserved replacement teeth are small, situated nearer the base of the crypt than the external margins of their

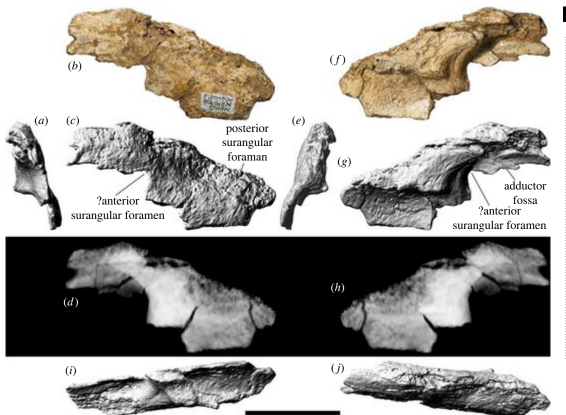


Figure 18. Diamantinasaurus matildae referred left surangular (AODF 0906). (a-j) Left surangular in (a) anterior, (b-d) lateral, (e) posterior, (f-h) medial, (i) ventral and (j) dorsal views. (a), (c), (e), (g) and (i-j) are three-dimensional models derived from surface scans; (b) and (f) are photographs; and (d) and (h) are three-dimensional models derived from CT scans. Scale bar = 50 mm.

alveoli, only present in the distalmost alveoli, and barely distinguishable from the bone of the maxilla. By contrast, the premaxillary replacement teeth are large, some are close to eruption, and all are clearly differentiable from the surrounding bone (figure 4j–l).

As outlined above, five replacement teeth are preserved within the left premaxilla of AODF 0906: two in the first alveolus, one in the second, and two in the third (figure 19). Thus, as in all sauropods other than diplodocoids, there were three or fewer replacement teeth per alveolus in AODF 0906 [104]. This description is based primarily on the largest replacement teeth present: the labialmost tooth in each of the first (figure 20) and third alveoli.

The teeth within the premaxilla of AODF 0906 are almost identical to those from the Diamantinasaurus matildae type (AODL 0085) and referred specimen (AODL 0127) sites, and those from the 'Mitchell' site (AODL 0270; [43]). They also show some similarity to some of the sauropod teeth described from Lightning Ridge [33,35]. The crown of each tooth of the AODF 0906 premaxilla is tapered apically, but otherwise has parallel sides, showing virtually no expansion above the root, as in neosauropods generally [100,171]. The cross-sectional shape of the crown of each tooth is generally D-shaped (figure 20d), although each becomes more rounded towards the root. The labial surface (figure 20a) of each tooth is generally smooth—as in titanosaurs, some early-deriving somphospondylans, and most diplodocoids—thereby contrasting with the longitudinally grooved surfaces seen in the teeth of most other sauropods [38,106]. The lingual surface (figure 20c) of each tooth is mostly convex, as is the case in titanosaurs generally [8,100,101]. The teeth lack an apicobasally oriented lingual ridge, as in most somphospondylans [101,172]. They also lack the bulges present towards the mesial and distal ends of the lingual surface (near the crown base), differentiating AODF 0906 from Euhelopus and Yongjinglong [113,114,173]. The teeth lack prominent mesial (figure 20d) and distal (figure 20b) carinae. Carinae characterize the teeth of most somphospondylans, although they are also absent in Ligabuesaurus [174], Sauroposeidon [175], and a small number of titanosaurs, including Rapetosaurus and Sarmientosaurus [41,101,152]. Based on the synchrotron scan data, the thickness of the tooth enamel is uniform labially and lingually. As in other somphospondylans, serrations or denticles are not evident on any of the teeth [8,101,104].

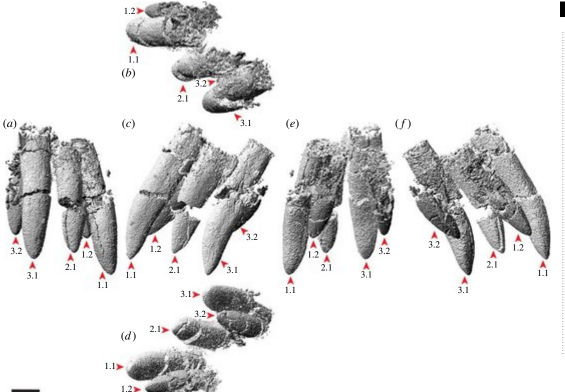


Figure 19. Diamantinasaurus matildae referred premaxillary replacement teeth (AODF 0906). (a-f) Three-dimensional models of the left premaxillary replacement teeth (derived from synchrotron scan data) in (a) anterior, (b) dorsal, (c) lateral, (d) ventral, (e) posterior and (f) medial views. The number preceding the period (1-3) indicates the alveolus position; the number succeeding the period (1 or 2) indicates the position of that tooth within the alveolus, with 1 being more labial than 2. Scale bar = 10 mm.

Four of the five premaxillary replacement teeth were measured (table 2), with the only exception being the severely damaged tooth in alveolus 3. The apicobasal heights of the preserved tooth crowns range from 25.40 to 28.07 mm (table 2). Thus, they are larger than all of the sauropod teeth described from the Griman Creek Formation [35], similar in size to the sauropod teeth from the sites that produced the *Diamantinasaurus matildae* holotype (AODL 0085) and referred (AODL 0127) specimens in the Winton Formation, and smaller than the sauropod teeth from AODL 0270, also in the Winton Formation [43]. The slenderness indices (SI = apicobasal length: mesiodistal width of crown ratio) of the premaxillary replacement teeth range from 2.33 to 2.74 (table 2). Similar tooth SI indices have been recorded in early-branching macronarians, brachiosaurids, most non-titanosaurian somphospondylans, and early-branching titanosaurs such as *Sarmientosaurus*, whereas much higher SI indices are evident in most titanosaurs [8,23,26,43,100,101]. The AODF 0906 teeth are generally more slender than those from the Griman Creek Formation: the SI indices of these range from 1.36 to 3.10 (mean SI = 2.09), although only four of the 26 teeth sampled have SI indices greater than 2.40 [35]. The AODF 0906 teeth are also generally slenderer than those from AODL 0270, the SI indices of which range from 2.00 to 2.88 (with six of the 10 below 2.10) [43].

The compression indices (CI = labiolingual breadth: mesiodistal width of crown ratio; [176]) of the AODF 0906 premaxillary replacement teeth range from 0.84 to 0.98. Thus, they are more compressed than the teeth from the Griman Creek Formation: CI indices range from 0.60 to 0.91, but only four of the 26 teeth sampled have CI values exceeding 0.85 [35]. The CI indices of the sauropod teeth from AODL 0270 range from 0.71 to 1.09, with five of the 10 having CI values below 0.80; thus, the AODF 0906 teeth are generally more compressed than those from AODL 0270.

6. Hyobranchial apparatus

An isolated, elongate and incomplete element (230 mm long), found associated with the left quadratojugal, appears to represent part of the hyobranchial apparatus (figure 21). It is described as if

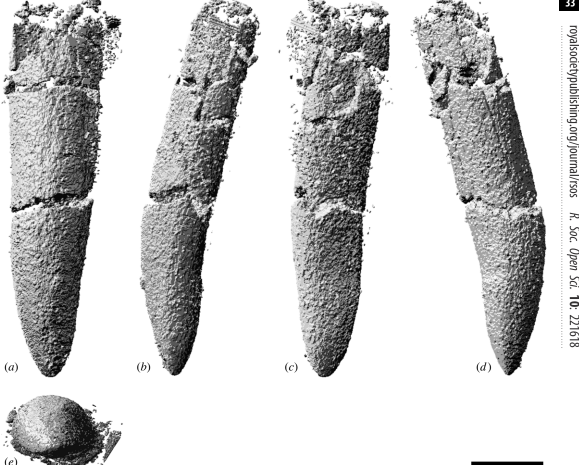


Figure 20. Diamantinasaurus matildae referred first premaxillary replacement tooth (AODF 0906). (a-e) Three-dimensional models of the left premaxillary replacement tooth 1.1 (derived from synchrotron scan data) in (a) labial, (b) distal, (c) lingual, (d) mesial and (e) apical views. Scale bar = 10 mm.

Table 2. Measurements of the premaxillary replacement teeth of AODF 0906 *Diamantinasaurus matildae*.

premaxillary tooth	apicobasal height	mesiodistal length	labiolingual breadth	SI	Cl
alveolus 1 (labialmost)	28.07	12.05	10.11	2.33	0.84
alveolus 1 (lingualmost)	25.89	9.44	8.71	2.74	0.92
alveolus 2	25.40	9.83	9.61	2.58	0.98
alveolus 3 (labialmost)	27.40	10.31	9.19	2.66	0.89

the long axis of the element runs anteroposteriorly, with the incomplete end interpreted as the anterior end and the complete end as the posterior tip, and the thinnest margin of the shaft being the dorsal one.

The incomplete anterior end is damaged (figure 21d), but is comma-shaped in cross-section. The dorsal margin of the shaft forms a narrow ridge along its length (figure 21a). The lateral surface is convex along its length (figure 21b,c). The medial surface (figure 21f,g) hosts a ridge along its midline, although this fades out towards the posterior end. Dorsal to this ridge on the anterior half, a prominent concavity is present. Ventral to this ridge, and dorsal to it on the posterior half, the medial surface is flat to slightly convex. The ventral margin is rounded except near the posterior end, where it becomes a mediolaterally narrow ridge (figure 21h). The complete posterior end is mediolaterally flattened (figure 21e).

Hyobranchial elements are not uncommonly preserved in early-branching sauropodomorphs but have relatively infrequently been identified in sauropods (table 3). Among the few sauropods for

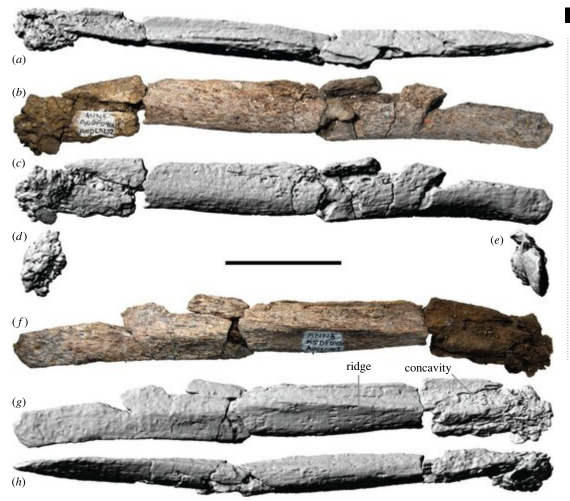


Figure 21. Diamantinasaurus matildae referred ?left ceratobranchial (AODF 0906). (a-h) ?Left ceratobranchial in (a) dorsal, (b,c) lateral, (d) anterior, (e) posterior, (f,g) medial and (h) ventral views. (a), (c-e) and (g-h) are three-dimensional models derived from surface scans; (b) and (f) are photographs. Scale bar = 50 mm.

which hyoid elements have been reported, those that have been described and/or illustrated in detail pertain to the early-diverging eusauropods *Shunosaurus lii* [207] and *Omeisaurus junghsiensis* [208], the diplodocid *Galeamopus pabsti* [211], the early-diverging macronarians *Europasaurus* [119,214], *Giraffatitan* [25] and *Abydosaurus* [26], and the somphospondylans *Phuwiangosaurus* [215] and *Tapuiasaurus* [21,22]. The AODF 0906 hyoid element bears little resemblance to the slender, straight element preserved in *Abydosaurus* [26], instead showing similarities to the flared posterior ends of the boomerang-like hyoid elements seen in the other sauropods listed above. The hyoid elements preserved in these sauropods, and specifically *Tapuiasaurus*, were interpreted by Wilson *et al.* [22] as second ceratobranchials, following the interpretation of an exquisitely preserved ankylosaurid hyobranchial apparatus by Hill *et al.* [177]. However, Yoshida *et al.* [178] revised the interpretation of this ankylosaurid, claiming that the element in question is more likely to be ceratobranchial 1 than ceratobranchial 2; according to those authors, ceratobranchial 2 is commonly lost in Archosauria. Thus, we interpret the probable hyoid element in AODF 0906 as a ceratobranchial, likely ceratobranchial 1.

7. Postcranial axial skeleton

7.1. Dorsal ribs

The four dorsal ribs that comprise part of AODF 0906 are poorly preserved and incomplete. Whereas all lack their proximal ends, some preserve enough of their distal ends to demonstrate that they were plank-like, as in all titanosauriforms [104], including all Australian Cretaceous sauropod taxa for which ribs are known: *Austrosaurus mckillopi* [39], *Wintonotitan* [9,36], *Diamantinasaurus* [9,39,44] and *Savannasaurus* [40].

Table 3. Sauropodomorphs for which hyoid elements have been reported. If the identification by Wilson *et al.* [22] of the hyoid elements of *Tapuiasaurus macedoi* as second ceratobranchials is correct (following the interpretation of the hyobranchial apparatus of the ankylosaurid *Pinacosaurus grangeri* presented by Hill *et al.* [177]), then the same identification should also be applied to the morphologically similar hyoid elements of (at least) *Shunosaurus lii, Omeisaurus junghsiensis, Diplodocus* sp., *Galeamopus pabsti, Europasaurus holgeri, Giraffatitan brancai, Phuwiangosaurus sirindhornae* and *Diamantinasaurus matildae* among sauropods. By contrast, if Yoshida *et al.* [178] are correct in their reinterpretation of the hyobranchial apparatus of *Pinacosaurus grangeri*, then the identification of these as first ceratobranchials is correct; this seems likely.

taxon	published interpreted identity	reference
non-sauropodan sauropodomorphs		
Buriolestes schultzei	hyoid	Cabreira <i>et al</i> . [179]
	hyoid	Müller <i>et al</i> . [180]
Pantydraco caducus	?hyoid	Kermack [181]
	?Corpus hyoideum + cornua	Galton & Kermack [182
Macrocollum itaqui	hyoid	Müller [183]
Unaysaurus tolentinoi	?hyoid	McPhee <i>et al</i> . [184]
Issi saaneq	ceratobranchial	Beccari <i>et al</i> . [185]
Plateosaurus spp.	illustrated but not identified	Jaekel [186]
	cornubranchial 1	Fürbringer [187]
	hyoid	Huene [188]
	hyoid	Huene [189]
	hypobranchial 1	Janensch [190]
	first ceratobranchial	Galton [191]
	ceratobranchial	Lallensack <i>et al</i> . [192]
Lufengosaurus huenei (='Fulengia youngi')	ceratobranchial 1	Carroll & Galton [193]
	ceratobranchial 1	Evans & Milner [194]
Xixiposaurus suni	hyoid	Sekiya [195]
Anchisaurus polyzelus	first ceratobranchial	Galton [196]
	ceratobranchial	Fabbri <i>et al</i> . [197]
Adeopapposaurus mognai	ceratobranchial	Martínez [198]
Leyesaurus marayensis	ceratobranchial	Apaldetti <i>et al.</i> [199]
Massospondylus carinatus	illustrated but not identified	Gow <i>et al.</i> [200]
	ceratobranchial 1 (cornu branchial 1)	Sues <i>et al</i> . [201]
Jingshanosaurus xinwaensis	hyoid	Zhang & Yang [202]
Melanorosaurus readi	ceratohyal	Yates [203]
Xingxiulong chengi	ceratobranchial	Wang <i>et al.</i> [204]
	ceratobranchial	Wang <i>et al.</i> [205]
Yunnanosaurus huangi	ceratobranchial	Barrett <i>et al.</i> [206]
auropods		
Shunosaurus lii	hyoid	Zhang [207]
Omeisaurus junghsiensis	hyoid	Dong <i>et al</i> . [208]
Mamenchisaurus jingyanensis	hyoid	Zhang <i>et al</i> . [209]
Diplodocus sp.	ceratobranchial	Woodruff <i>et al</i> . [210]
Galeamopus pabsti	ceratobranchial	Tschopp & Mateus [21
Brontosaurus excelsus	two sets of hyoid bones	Marsh [212]
Lavocatisaurus agrioensis	hyoid	Canudo <i>et al</i> . [213]

axon	published interpreted identity	reference
Europasaurus holgeri	ceratobranchial 1 (?)	Laven [214]
	ceratobranchial 1	Marpmann <i>et al.</i> [119]
Camarasaurus lentus	thyrohyals	Gilmore [135]
	three fragments pertaining to two bones	Wilson et al. [22]
Giraffatitan brancai	hypobranchial 1	Janensch [190]
	hypobranchial 1	Janensch [25]
Abydosaurus mcintoshi	hyoid	Chure <i>et al</i> . [26]
Phuwiangosaurus sirindhornae	hyoid	Martin <i>et al.</i> [215]
Diamantinasaurus matildae	ceratobranchial 1	this paper
Tapuiasaurus macedoi	hyoid	Zaher <i>et al</i> . [21]
	ceratobranchial 2	Wilson et al. [22]

7.2. Sacral vertebrae

In vertebrate palaeontology, a vertebra is interpreted as a sacral if it makes contact with the ilium [216,217]. However, sacral centra also coalesce through ontogeny, such that the vertebrae that contact the ilium also tend to become coossified in adults [95]. Thus, we contend that it is reasonable to interpret coalesced, non-pathological vertebrae in sauropods as sacral, particularly when they are morphologically consistent with other described sacral vertebrae.

The sacrum of AODF 0906 (figure 22*a–e*) was found in three pieces: one comprising the centrum and partial lower transverse processes of sacral vertebra I, a second comprising sacral centra II–IV (and the bases of the lower transverse processes of sacral vertebra IV), and a third comprising the centrum and partial lower transverse processes of sacral vertebra V.

The sacrum of AODF 0906 comprises five sacral vertebrae, albeit with the centrum of sacral vertebra V unsutured to sacral vertebra IV. Although some non-titanosaurian somphospondylans, namely Huanghetitan liujiaxiaensis [218], Sauroposeidon [175], Sibirotitan [219] and Tastavinsaurus [220,221], also have five sacral vertebrae, most somphospondylans-including all known titanosaurs that preserve complete sacra, as well as several taxa that probably lie outside of Titanosauria (e.g. Dongyangosaurus, Huabeisaurus, Phuwiangosaurus and Ruyangosaurus [149,222–224])—have six or more [95,100,101,106,109]. Thus, AODF 0906 appears to be plesiomorphic with respect to this character. The sacrum of the holotype specimen of Diamantinasaurus matildae (AODF 0603), which has been more completely prepared since its initial description [37], is now known to also comprise only five coalesced vertebrae (figure 22f). The previously figured portion of the sacrum of AODF 0603 comprises sacral vertebrae II-V, not III-VI as labelled in Poropat et al. [37]). An as yet unpublished diamantinasaurian specimen from the Winton Formation (AODF 0888), which is also likely referable to Diamantinasaurus matildae, preserves a sacrum comprising five coossified vertebrae: the transverse processes of the posterior four sacral vertebrae unequivocally form a sacricostal yoke that contacts the ilium (figure $22g_{,h}$). We infer that the vertebra fused to the anteriormost of these four also contacted the ilium, giving a sacral count of five. Whether or not an additional sacral vertebra was present anterior to this is unclear, but we consider it unlikely.

In AODF 0906, the anterior articular surface of the centrum of sacral vertebra I is convex and dorsoventrally compressed. Each lateral surface of sacral vertebra I preserves a pneumatic foramen, with that on the right side (figure 22b) better preserved than that on the left (figure 22e). The holotype sacrum of *Diamantinasaurus matildae* (AODF 0603) also has a prominent pneumatic foramen on sacral vertebra I (figure 22f). However, in both AODF 0603 and AODF 0906, pneumatic foramina are not evident on any of the other sacral centra. The presence of a pneumatic foramen on at least sacral vertebra I in both AODF 0603 and AODF 0906 distinguishes *Diamantinasaurus* from some titanosaurs, including *Opisthocoelicaudia skarzynskii* and *Saltasaurus*, wherein all sacral vertebrae lack pneumatic foramina [100,225]. Sacral vertebrae II and III in AODF 0906 have suffered extensive taphonomic dorsoventral compression (possibly a result of *post mortem* trampling by another dinosaur) and are too poorly preserved to elicit much comment. Their internal texture is camellate (possibly semicamellate), as in somphospondylans generally [101], and the transverse widths of the

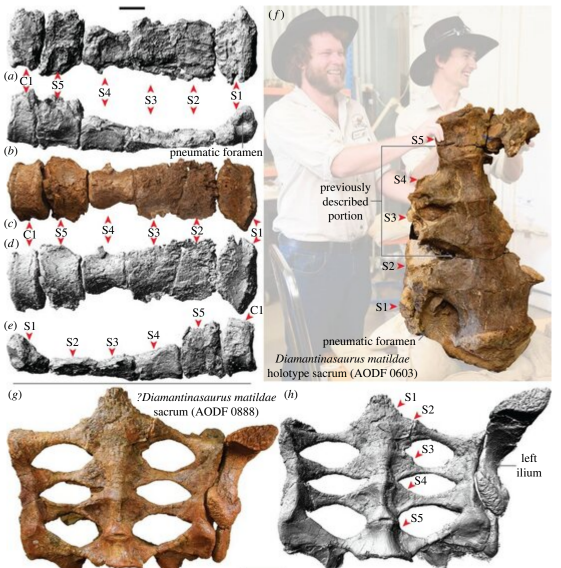


Figure 22. Diamantinasaurus matildae referred sacrum and caudal vertebra 1 (AODF 0906), Diamantinasaurus matildae holotype sacrum (AODF 0603); and ?Diamantinasaurus matildae referred sacrum and left ilium (AODF 0888). (a-e) Diamantinasaurus matildae referred (AODF 0906) sacral vertebrae 1–5 and caudal vertebra 1 of AODF 0906 in (a) dorsal, (b) right lateral, (c,d) ventral and (e) left lateral views. (a-b) and (d-e) are three-dimensional models derived from surface scans, (c) is a photograph. Scale bar = 100 mm. (f) Diamantinasaurus matildae holotype (AODF 0603) sacrum in ventrolateral view, with anterior towards bottom of page, showing the position of the previously described portion indicated. (g,h) ?Diamantinasaurus matildae referred (AODF 0888) sacrum and left ilium in ventral view. (g) is a photograph, (h) is a three-dimensional model derived from a surface scan. Scale bar = 200 mm.

middle sacral centra appear to be little different from those of the first and last sacral centra, distinguishing AODF 0906 from saltasaurids and aeolosaurines, in which the middle centra are 'waisted' relative to the first and last sacral centra [38,226–228]. The ventral surfaces of sacral vertebrae IV and V are anteroposteriorly concave and transversely convex. The acuteness of the transverse convexity is greater in sacral IV than in sacral V, such that it almost forms a weak median ridge in the former (figure 22d). Sacral vertebrae IV and V each preserve the bases of their transverse processes, and these are situated near the mid-length of the centrum in both vertebrae. The posterior articular surface of sacral vertebra IV and the anterior articular surface of the centrum of sacral vertebra V are both essentially flat, and these vertebrae were clearly unsutured in life; this implies that AODF 0906 was osteologically immature at death. The posterior articular surface of sacral vertebra V is shallowly concave and slightly wider transversely than tall dorsoventrally.

Table 4. Measurements of the sacral vertebrae and caudal vertebra of AODF 0906 *Diamantinasaurus matildae*. An asterisk (*) indicates a tentative measurement based on an incomplete or distorted specimen.

			sacral	vertebrae				caudal
measureme	nts (mm)		T	II	III	IV	٧	vertebra
centrum	anteroposte	rior length	118	138	139	147	134	116
	anterior	maximum dorsoventral height	124	—	—	—	155	165
		maximum transverse width	247	194*	145*	126	144	201
	midline	minimum transverse width	192	137*	127*	92	149	—
	posterior	maximum dorsoventral height		—	—	112	168	162
		maximum transverse width	—	218*	154	135	214	205

7.3. Caudal vertebra

The only caudal vertebra preserved is represented by an incomplete centrum (figure 22a-e). It is presumably caudal vertebra I, based on the size congruence of its anterior articular surface with that of the posterior articular surface of sacral vertebra V. Breakages on the dorsal surface of the centrum either side of the ventral margin of the neural canal reveal spongiose, but not camellate, internal texture. In this regard, it is similar to most sauropods, in which camellae do not extend into the tail [104,229]; however, it is possible that the AODF 0906 caudal neural arch (which is not preserved) was characterized by camellae, given that this tissue structure is present in that region, but not the centrum, in anterior caudal vertebrae of Savannasaurus [40]. The articular surfaces of the centrum are wider transversely than they are tall dorsoventrally (table 4), as in many titanosauriforms [8,101]. Both articular surfaces are shallowly concave, as is the case in the anteriormost caudal centra of some brachiosaurids and non-titanosaurian somphospondylans (wherein they are flat or concave), and differs from nearly all other titanosaurs, wherein one or both articular surfaces are convex [6,38,96,100,101,104,106,109,131]. However, Savannasaurus is also characterized by amphicoelous anterior caudal centra [38,40]. The average Elongation Index of the centrum of AODF 0906, calculated by dividing the anteroposterior length of the centrum by the mean of the mediolateral width and dorsoventral height of the anterior articular surface of the centrum [6,8,100,101,106,132], is 0.63. The ventral surface is anteroposteriorly concave and transversely convex (figure 22e). A tiny foramen (possibly pneumatic) is evident on the right lateral surface (figure 22b), and several probable nutrient foramina are present on the less complete left lateral surface (figure 22e). Despite the nearly complete state of the centrum, there is no evidence of a transverse process or a pneumatic foramen; this distinguishes AODF 0906 from the proximal-middle anterior caudal vertebrae of Savannasaurus [40]. However, a pneumatic opening is not always present in the anteriormost caudal centra, with substantial serial variation observed in some taxa [230,231], and thus its absence in AODF 0906 might merely reflect its proximal placement in the tail.

7.4. Chevron

The sole chevron preserved is almost complete (figure 23; table 5), missing only the distal end, the proximal left ramus, and the medial half of the right ramus. The chevron has also been slightly impacted by *post mortem* distortion. As is typical of titanosauriform chevrons, it is Y-shaped in anterior (figure 23a,b) and posterior (figure 23g,h) aspects. Similarly, as in most titanosauriforms (other than, for example, *Dongbeititan dongi*, *Daxiatitan binglingi* and *Xianshanosaurus shijiagouensis*), the haemal canal is open proximally rather than bridged [100,101,106,144,232]. The depth of the haemal canal is slightly less than one third the total proximodistal length of the chevron; the chevrons of the contemporaneous titanosauriform *Wintonotitan* show a similarly shallow haemal canal [36], as do those of some other titanosauriforms, including the non-titanosaurian somphospondylans *Dongbeititan*, *Daxiatitan*, *Xianshanosaurus* and *Sauroposeidon*, and the saltasaurine titanosaur *Saltasaurus* [101]. As preserved, the proximal articular surface of the right ramus is rounded and undivided (figure 23e,f); if this reflects the true morphology of the proximal articular surface, then AODF 0906 lacks the grooves on this surface seen in some somphospondylans, including the early-branching members *Tangvayosaurus hoffeti* and *Phuwiangosaurus* [5], and several titanosaurs, such as *Epachthosaurus sciuttoi*, *Lohuecotitan*

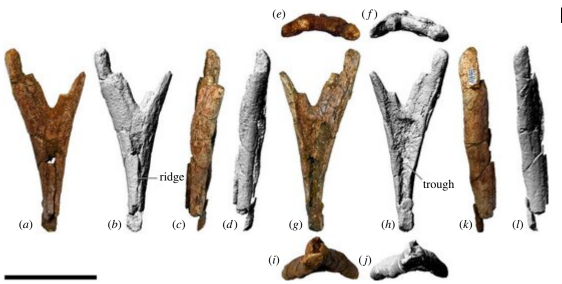


Figure 23. Diamantinasaurus matildae referred chevron (AODF 0906). (a-l) Chevron in (a,b) anterior, (c,d) left lateral, (e,f) dorsal, (g,h) posterior, (i,j) ventral and (k,l) right lateral views. (a), (c), (e), (g), (i) and (k) are photographs, and (b), (d), (f), (h), (j) and (l) are three-dimensional models derived from surface scans. Scale bar = 100 mm.

Table 5. Measurements of the chevron of AODF 0906 *Diamantinasaurus matildae*. An asterisk (*) indicates a tentative measurement based on an incomplete or distorted specimen.

measurements (mm)		chevron
dorsoventral height		201*
haemal canal depth		62*
haemal canal depth: chevron dorsoventral height		~0.31
distal shaft	anteroposterior	23
	transverse	21.5

pandafilandi, Mendozasaurus neguyelap, Notocolossus gonzalezparejasi and Aeolosaurus [5,38,145,233–236]. The anterior surface of each ramus, and the anterior surface immediately distal to their junction, is smoothly transversely convex. The lateral surface of each ramus lacks ridges and bulges (figure 23c,d,k, l), thereby contrasting with the chevrons of *Epachthosaurus* and some saltasaurids, in which such structures are present [38,234], although these are sometimes absent from the anteriormost chevrons [2]. The distal blade forms a prominent ridge anteriorly and hosts a deep trough posteriorly. This contrasts with *Wintonotitan*, wherein both the anterior and posterior margins of the distal blade form ridges [36].

8. Pelvic girdle

8.1. Ilium

The incomplete left ilium preserves the pubic and ischiadic processes—and therefore most of the acetabular margin—but lacks much of the blade, including the pre- and postacetabular processes (figure 24*a*,*c* and table 6). In all respects, it is practically identical to the ilium of the holotype of *Diamantinasaurus matildae* (figure 24*b*; [37]); it is also somewhat akin to the similarly incomplete ilium of *Wintonotitan wattsi* [36]. Despite being incompletely preserved, it is clear that, when viewed dorsally, the preacetabular lobe would have flared anterolaterally, as in neosauropods generally [100,106,109,144]. The pubic peduncle is perpendicular to the long axis of the AODF 0906 ilium, as in virtually all titanosauriforms [96]. The anteroposterior length (61 mm) of the incomplete pubic peduncle of the ilium is barely more than 25%

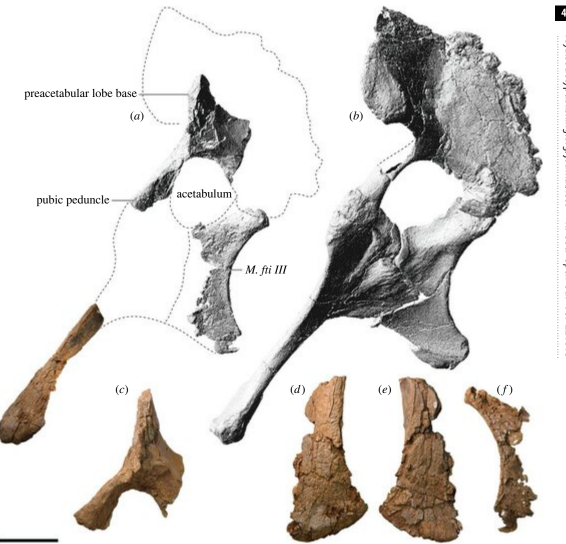


Figure 24. Diamantinasaurus matildae referred pelvis (AODF 0906), and Diamantinasaurus matildae holotype pelvis (AODF 0603).

(a) Diamantinasaurus matildae referred (AODF 0906) left ilium, pubis and ischium in lateral view. (b) Diamantinasaurus matildae holotype (AODF 0603) articulated left ilium, pubis and ischium. (c) Diamantinasaurus matildae referred (AODF 0906) left ilium in lateral view. (d,e) Diamantinasaurus matildae referred (AODF 0906) left pubis in (d) anterior and (e) posterior views. (f) Diamantinasaurus matildae referred (AODF 0906) left ischium in medial view. (a) and (b) are three-dimensional models derived from surface scans (excluding the pubis in (a)), and (c-f) are photographs. M. fti III, M. flexor tibialis internus III. Scale bar = 200 mm.

Table 6. Measurements of the left ilium and ischium of AODF 0906 *Diamantinasaurus matildae*. An asterisk (*) indicates a tentative measurement based on an incomplete or distorted specimen.

measurement (mm)	left ilium	left ischium
maximum proximodistal length	_	646
maximum anteroposterior length	422*	242
maximum mediolateral diameter of pubic peduncle	228*	
maximum anteroposterior diameter of pubic peduncle	61*	
maximum mediolateral diameter of iliac peduncle		63*
maximum anteroposterior diameter of iliac peduncle		77*

its mediolateral width (220 mm), compared with just under 50% in the holotype of *Diamantinasaurus matildae* [37]. In saltasaurids, the anteroposterior length of the pubic peduncle of the ilium is at least 50% its mediolateral width [6].

8.2. Pubis

The left pubis (figure 25a and d,e) is represented by its distal one-half and is practically identical to the pubes of the holotype of *Diamantinasaurus matildae* [37]. It is also very similar to the pubes of *Australotitan cooperensis* [34]. The distal end is laminar and expanded to form an anterior boot, as in some other titanosaurs [3,38,101], albeit not to the extent seen in *Savannasaurus elliottorum* [40].

8.3. Ischium

The left ischium is more complete than the right (figure 24a and f; table 6), although it is missing almost the entire pubic articular surface, a substantial portion of its iliac peduncle, and most (if not all) of the ischiadic articular surface. In all respects it is very similar to the holotype ischia of *Diamantinasaurus matildae* (figure 24b; [37]), *Savannasaurus elliottorum* [38,40] and *Australotitan cooperensis* [34]. It is also very similar to the holotype left ischium (QM F7292) of *Wintonotitan wattsi*, although the lateral ridge for *M. flexor tibialis internus III* is not as pronounced in AODF 0906 [36]. The preserved portion of the iliac peduncle demonstrates that it does not narrow anteroposteriorly in lateral view, and the mediolateral width of the acetabular margin is effectively uniform along its length. The only measurable portion of the proximal plate has an anteroposterior length of 177 mm, slightly greater than 25% the total proximodistal length of the ischium. This distinguishes AODF 0906 from several non-titanosaurian titanosauriforms, wherein the anteroposterior length of the proximal plate comprises less than 25% of the proximodistal length of the ischium [6,101]. The long axis of the shaft of the ischium projects towards the lower acetabular margin, as in all somphospondylans [95,100,101,106]. The attachment point for *M. flexor tibialis internus III* is a low ridge. No groove accompanies this ridge, with this also absent in most other titanosauriforms [5,6,38].

9. Hind limb

9.1. Femur

The right femur is incomplete and poorly preserved in two sections. By contrast, the left femur (figure 25*b*–*h*; table 7) is well preserved and almost complete, missing only the greater trochanter, part of the fibular condyle, and the lateral epicondyle (assuming it was present). It has, however, suffered some distortion, especially on the lateral margin (the curvature seen in figure 25*d* is not considered to be natural). The subsequent description is based entirely on the left femur.

In all measured dimensions, the left femur of AODF 0906 (table 7) is very slightly larger than the holotype right femur of *Diamantinasaurus matildae* (AODF 0603; [9,37,44,237]). However, the measured proximodistal length is likely exaggerated (possibly by as much as 50 mm) by infilled fractures in the element. Morphologically, the femur is also closely comparable with the holotype femur of *Diamantinasaurus matildae* [9,37]. The only notable difference between the femora of AODF 0906 and the *Diamantinasaurus matildae* holotype specimen (AODF 0603: figure 25a,i) is the morphology of the lateral bulge: in the latter, this structure is separated from the rest of the femur by a deep, longitudinal groove [37]. However, in AODF 0906 this section of the femur is flat. The AODF 0906 left femur shows evidence of substantial distortion, thereby accounting for the slight morphological disparity between it and the *Diamantinasaurus matildae* holotype femur observed here.

The femoral head is directed medially, rather than dorsomedially (figure 25*b,c,g*); among titanosaurs, medially deflected femoral heads are present in relatively few taxa, including *Alamosaurus*, *Diamantinasaurus*, *Jainosaurus*, *Malawisaurus*, *Mendozasaurus* and *Pitekunsaurus* [3,8,38,238]. The proximal lateral margin forms a flange-like trochanteric shelf, as in nearly all titanosaurs [101]. The proximolateral margin (dorsal to the lateral bulge) lies medial to the lateral margin of the distal half of the femoral shaft, differentiating AODF 0906 from some early-branching somphospondylans (e.g. *Tastavinsaurus*), *Patagotitan mayorum*, and saltasaurid titanosaurs, wherein this margin lies lateral to the lateral margin of the distal half of the femoral shaft [96,101,109,221,232,238,239]. The anterior surface (figure 25*b,c*) preserves a longitudinal midline ridge (*linea intermuscularis cranialis*) and a concavity on its proximal two-thirds, as is also the case in the *Diamantinasaurus matildae* holotype (figure 25*a*; [37]), as well as several other titanosaurs, including *Alamosaurus*, *Australotitan*, *Pellegrinisaurus powelli* and saltasaurines [5,34,240,241]. The fourth trochanter, which is situated on the posterior surface, slightly proximal to the mid-length (figure 25*g*), is manifested as a low, rounded ridge, contrasting with saltasaurines, wherein it is reduced to near absence [2]. The fourth trochanter is not visible in anterior view, distinguishing AODF 0906 from most non-somphospondylan macronarians, as well as some non-titanosaurian (e.g. *Phuwiangosaurus*, *Sauroposeidon*) and titanosaurian (e.g.

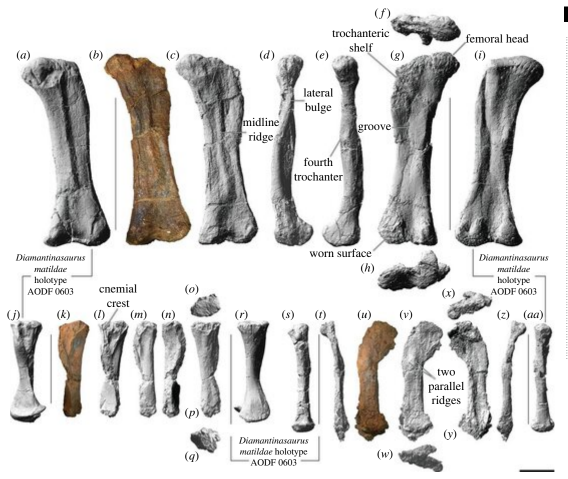


Figure 25. Diamantinasaurus matildae holotype hind limb elements (AODF 0603), and Diamantinasaurus matildae referred hind limb elements (AODF 0906). (a and i) Diamantinasaurus matildae holotype (AODF 0603) left femur (right femur mirrored) in (a) anterior and (i) posterior views. (b–h) Diamantinasaurus matildae referred (AODF 0906) left femur in (b–c) anterior, (d) lateral, (e) medial, (f) proximal, (g) posterior and (f) distal views. (f) Diamantinasaurus matildae holotype (AODF 0603) left tibia (right tibia mirrored) in (f) anterior and (f) posterior views. (f–g) Diamantinasaurus matildae referred (AODF 0906) left tibia in (f-g) anterior, (f) medial, (f) posterior and (f) posterior and (f) distal views. (f-g) Diamantinasaurus matildae holotype (AODF 0603) left fibula (right fibula mirrored) in (f) anterior and (f) posterior views. (f–g) Diamantinasaurus matildae referred (AODF 0906) left fibula in (f) anterior, (f-g) lateral, (f) distal, (f) proximal, (f) medial and (f) posterior views. (f-g) Diamantinasaurus matildae referred (AODF 0906) left fibula in (f) anterior, (f-g) lateral, (f) distal, (f) proximal, (f) medial and (f) posterior views. (f-g), (f-g) are three-dimensional models derived from surface scans, and (f), (f) and (f) are photographs. Scale bar = 250 mm.

Table 7. Measurements of the left femur of AODF 0906 *Diamantinasaurus matildae*.

measurements (mm)	left femur
proximodistal length	1366
proximal end mediolateral width	430
mid-shaft mediolateral width	261
mid-shaft anteroposterior length	126
mid-shaft minimum circumference	686
distal end maximum mediolateral width	435

Malarguesaurus, *Tapuiasaurus*) somphospondylans [101]. The femoral shaft is anteroposteriorly compressed (ratio of mediolateral width: anteroposterior length at midshaft = 2.07 versus 2.18 in the holotype; figure 25*d*, *e*) and the greatest anteroposterior thickness of the shaft (93.5 mm) is slightly more than one-third that of the anteroposterior length of the tibial condyle (272.5 mm); by contrast, in some non-titanosaurian macronarians,

Table 8. Measurements of the left tibia of AODF 0906 *Diamantinasaurus matildae*. An asterisk (*) indicates a tentative measurement based on an incomplete or distorted specimen.

measurements (mm)	left tibia	right tibia
proximodistal length	786*	532*
proximal end mediolateral width	152*	
proximal end anteroposterior width	247*	
mid-shaft long axis diameter	135*	
mid-shaft dimension perpendicular to long axis	98*	
mid-shaft minimum circumference	457	
distal end maximum mediolateral width	>155*	241
distal end maximum anteroposterior width	132*	170

the anteroposterior thickness of the shaft is at least half the anteroposterior length of the tibial condyle [131]. A longitudinal groove extends along the mid-shaft of the posterior surface (figure 25g). The distal condyles are similar to one another in mediolateral width (figure 25h) and are bevelled dorsomedially such that the fibular condyle extends further distally than the tibial one, as in many titanosaurs [101,104,242], including the *Diamantinasaurus matildae* holotype [37]. The distal condyles do not extend on to the anterior surface of the shaft, contrasting with the condition in titanosaurs such as *Rapetosaurus* and saltasaurines [104,242].

9.2. Tibia

The right tibia is incomplete, missing much of its proximal end. However, its distal end is reasonably well preserved and undistorted. The left tibia is similarly fragmentary (figure 25*k*–*q*; table 8), but does preserve the proximal end (which is incomplete on all margins except the anterior one), despite lacking the distal one. The following description is largely based on the left tibia, with supplementary information derived from the right one.

The tibia of AODF 0906 is very similar to that of the *Diamantinasaurus matildae* holotype specimen (figure 25*j* and *r*; [37]). The proximodistal length of the composite tibia is just under 60% the length of the femur. This is comparable to the condition in the holotype of *Diamantinasaurus matildae* [37], as well as in several non-titanosaurian somphospondylans [38], including *Chubutisaurus*, *Ligabuesaurus* and *Tastavinsaurus* [174,220,243]. The lateral edge of the proximal end (posterior to the cnemial crest) lacks the pinched-out projection, or 'second cnemial crest', seen in some somphospondylans [101]. The deep anterolateral fossa hosts a small proximal projection and is bordered posteriorly by a sharp longitudinal ridge (the presence of a double ridge cannot be determined owing to incompleteness). Several tibial autapomorphies of *Diamantinasaurus matildae* [37] cannot be assessed in AODF 0906, including whether or not a fossa was present posterior to the lateral longitudinal ridge(s). No tubercle is evident on the posterior/medial surface of the cnemial crest, contrasting with some early-diverging macronarians [157]. The complete and undistorted distal end of the right tibia is, in all respects, effectively identical to the holotype tibia of *Diamantinasaurus matildae* [37].

9.3. Fibula

The left fibula is almost complete (figure 25*t*–*z* and table 9), but has suffered substantial mediolateral distortion, particularly at its proximal and distal ends. Consequently, it is not possible to assess the presence or absence of the sole fibular autapomorphy of *Diamantinasaurus matildae*: the ridges and grooves on the medial surface near the mid-length [37]. Nevertheless, the fibula of AODF 0906 is similar to that of the *Diamantinasaurus matildae* holotype specimen (figure 25*s* and AA). The proximal end is comma-shaped, with an anteromedially directed crest, as in many somphospondylans [5,101,113]. A medial triangular scar appears to be present at the proximal end. The fairly prominent lateral trochanter, which is manifested as two parallel ridges, as in most titanosauriforms [100,144], is situated one-third of the way down the fibula shaft. In lateral view, the shaft appears sigmoidal, but this is only because of the substantial mediolateral (and oblique) taphonomic distortion to which this element has been subjected: it was likely straight in life. The right fibula is less complete, but better

Table 9. Measurements of the left fibula of AODF 0906 *Diamantinasaurus matildae*. An asterisk (*) indicates a tentative measurement based on an incomplete or distorted specimen.

measurements (mm)	left fibula	right fibula
proximodistal length	865	704*
maximum proximal mediolateral breadth	79	
maximum proximal anteroposterior width	223	
mid-shaft mediolateral breadth	63	
mid-shaft anteroposterior width	134	121*
mid-shaft minimum circumference	357	
maximum distal mediolateral breadth	93*	98
maximum distal anteroposterior width	>200*	136

preserved, than the left fibula. The proximal end is missing, but the mid-shaft and distal end are well represented; the distal end is practically identical to the subtriangular distal end of the holotype fibula (AODF 0603) of *Diamantinasaurus matildae* [37].

9.4. Astragalus

A worn element might represent that the worn dorsal portion of the right astragalus is present. However, very little of its morphology can be appraised.

9.5. Metatarsals

AODF 0906 preserves all five right metatarsals (figures 26 and 27; table 10). Metatarsal III is the longest, followed by IV, II, V and I. The fact that metatarsal I is shorter proximodistally than metatarsal V (ratio 0.8) is a feature that AODF 0906 shares with most macronarians other than *Opisthocoelicaudia* and *Rapetosaurus* [101]. The substantial disparity between the proximodistal lengths of metatarsal III and I (III: I ratio = 1.76, although taphonomic distortion might have slightly exaggerated the length of metatarsal III) is a feature that AODF 0906 shares with most titanosauriforms, other than *Notocolossus* wherein this ratio is 1.2 [236,244]. Metatarsal III is longer proximodistally than metatarsal IV (III: IV ratio = 1.09), as in most titanosauriforms (e.g. *Opisthocoelicaudia, Rapetosaurus, Tastavinsaurus*) other than *Alamosaurus, Epachthosaurus* and *Notocolossus* [236,244]. Metatarsals I and II appear to lack dorsolateral rugosities near their distal ends, distinguishing AODF 0906 from some diplodocoids [106].

9.5.1. Metatarsal I

Right metatarsal I (figure 27a-d,t,u) is almost complete, missing only a small section of the ventrolateral corner of the proximal end. In proximal view (figure 27b), the metatarsal has a convex medial margin and a lateral margin that is flat in its dorsal half and concave in its ventral half. The dorsal junction between the medial and lateral margins is manifested as a point situated near the lateral margin. When complete, the incomplete ventral margin was presumably broadly convex. The proximal surface is convex. In dorsal view (figure 27t), the distal half of metatarsal I shows effectively no flaring: it shares this morphology with some early-branching somphospondylans (e.g. Tastavinsaurus) and titanosaurs (e.g. Epachthosaurus, Mendozasaurus, Opisthocoelicaudia), but differs from other early-branching somphospondylans (e.g. Gobititan, Ligabuesaurus) and titanosaurs (e.g. Alamosaurus, Muyelensaurus, Notocolossus, Rapetosaurus, Saltasaurus), wherein ventrolateral expansion of this element results in significant distal flaring [101,245]. The dorsal surface is concave proximodistally, convex mediolaterally, and inclined such that it faces somewhat dorsomedially. The junction between the dorsal and lateral surfaces is a reasonably pronounced ridge. The lateral surface is essentially flat, faces slightly ventrolaterally, and is flared proximally and tapered distally (figure 27c). The ventral surface is incomplete proximolaterally but was evidently concave centrally and convex along its proximomedial and distal margins (figure 27u). The medial margin merges smoothly with both the dorsal and ventral margins (figure 27a). Although it has a subtle proximodistally elongate ridge at its mid-height, it appears to lack the median tubercle on this surface that is present in several brachiosaurids [6,246]. The distal

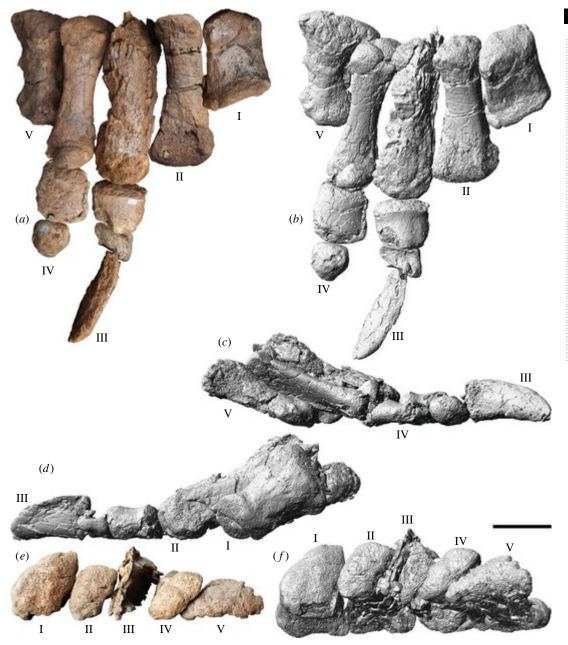


Figure 26. Diamantinasaurus matildae referred right pes (AODF 0906). (a-f) Articulated right pes in (a,b) dorsal, (c) lateral, (d) medial and (e,f) proximal views. (a) and (e) are photographs; (b-d) and (f) are three-dimensional models derived from surface scans. Scale = 100 mm.

end is 'D'-shaped, with the slightly concave ventral margin corresponding to the flat side of the 'D' and all other margins convex (figure 27d). The distal surface itself is mediolaterally concave and dorsoventrally convex either side of the midline, essentially forming two condyles for articulation with pedal phalanx I-1.

9.5.2. Metatarsal II

Right metatarsal II is practically complete (figure 27e–h,t,u), with only minor wear evident at the proximal and distal ends. The proximodistal length of metatarsal II is 30% that of the incomplete tibia. In proximal view (figure 27f), the metatarsal is broadly wedge-shaped. The dorsal margin is narrow and mediolaterally convex, and the medial margin is shallowly convex. The lateral margin is somewhat more complex, being divisible into a shallowly concave section that faces laterally (dorsal two-thirds) and a straight section that faces ventrolaterally (ventral one-third). In several other titanosaurs (e.g. Epachthosaurus, Mendozasaurus, Muyelensaurus), the proximal end of metatarsal II is also concave laterally; however, in most taxa it is straight [101]. The proximal surface is mostly concave, but hosts a subtle depression in its dorsal one-third.

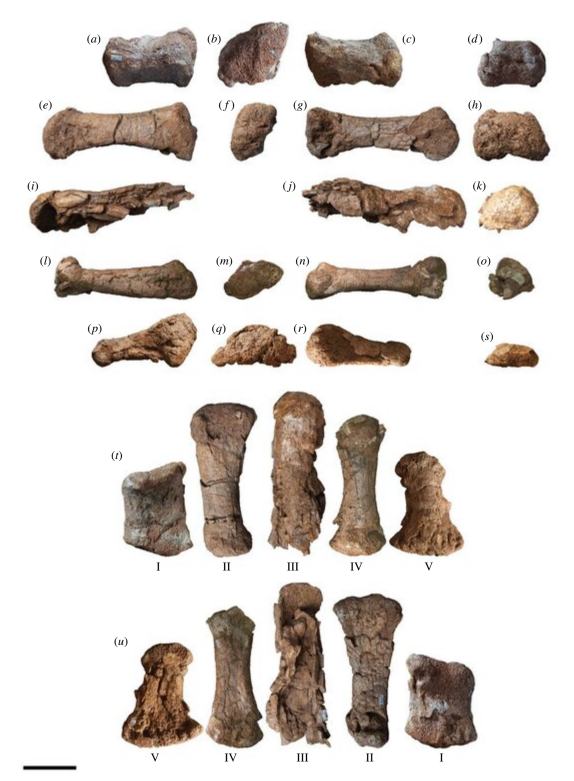


Figure 27. Diamantinasaurus matildae referred right metatarsals (AODF 0906). (a-d) Right metatarsal I in (a) medial, (b) proximal, (c) lateral and (d) distal views. (e-h) Right metatarsal II in (e) medial, (f) proximal, (g) lateral and (h) distal views. (i-k) Right metatarsal III in (i) medial, (j) lateral and (k) distal views. (l-o) Right metatarsal IV in (l) medial, (m) proximal, (n) lateral and (o) distal views. (p-s) Right metatarsal V in (p) medial, (q) proximal, (r) lateral and (s) distal views. (t-u) Right metatarsals I-V in (t) dorsal and (u) ventral (plantar) views. Scale bar = 100 mm.

In dorsal view, metatarsal II is flared at both the proximal and distal ends, with the flaring of the latter significantly more pronounced than the former. Although the distal one-quarter appears to have hosted a slight depression, this surface is otherwise proximodistally concave and almost entirely mediolaterally convex (figure 27t). The junction between the dorsal and lateral surfaces is manifested as a very subtle

Table 10. Measurements of the right metatarsals of AODF 0906 *Diamantinasaurus matildae*. An asterisk (*) indicates a tentative measurement based on an incomplete or distorted specimen.

	right meta	atarsals			
measurement (mm)	I I	II	III	IV	٧
proximodistal length	157	243	277*	255*	198
maximum proximal mediolateral width	85	82	_	98	135
maximum proximal dorsoventral height	109	91	_	73	65
mid-shaft mediolateral breadth	98	64	_	46	70*
maximum distal mediolateral width	110	110	96*	84	94
maximum distal dorsoventral height	74	77	65*	50*	40

ridge that is more pronounced distally. The lateral surface (figure 27g) is proximodistally concave, dorsoventrally convex, and twisted such that the proximal one-third faces slightly ventrolaterally to accommodate metatarsal III. Separated from the lateral surface by a subtle ridge, the ventral surface (figure 27u) varies in aspect along its length despite being broadly proximodistally concave: whereas the proximal half is mediolaterally convex and faces slightly laterally, the distal half faces entirely ventrally and is mediolaterally concave. The medial surface is essentially flat (figure 27e), and its junctions with both the dorsal and ventral surfaces are not marked by ridges despite being relatively abrupt. The ventral surface is convex (figure 27u). In distal view (figure 27h), the metatarsal is essentially trapezoidal, with the ventral margin mediolaterally wider than the dorsal one and the lateral and medial margins inclined inwards towards the dorsal margin. In AODF 0906, and in most somphospondylans, the distal articular surface does not extend on to the dorsal margin of the shaft; however, in some somphospondylans (e.g. Alamosaurus, Epachthosaurus, Euhelopus, Gobititan), the inverse is true [101,245]. The ventral margin of the distal end is concave, lacking the ventrolateral projection seen in the early-branching somphospondylans Dongbeititan, Gobititan and Tastavinsaurus, as well as the titanosaur Muyelensaurus [6].

9.5.3. Metatarsal III

Metatarsal III (figure 27i–k,t,u) is incomplete proximally and ventrally and has suffered substantial taphonomic distortion (dorsoventral compression). Consequently, its preserved proximodistal length might be a slight exaggeration. Nevertheless, it is probable that metatarsal III was at least 25% (and likely more than 30%) the length of the tibia. Few titanosaurs can be appraised for this feature, but those that can (e.g. *Epachthosaurus*, *Opisthocoelicaudia*) also have third metatarsals that are elongate relative to tibia length; among early-deriving somphospondylans, only *Tastavinsaurus* shares this feature [101,220]. Despite the distortion to which it has been subjected, it appears that the metatarsal III of AODF 0906 was less robust than that of *Savannasaurus elliottorum* [38,40].

9.5.4. Metatarsal IV

Right metatarsal IV is practically complete (figure 27*I*–*o,t,u*), but the distal end is fractured and slightly distorted. The proximal end is sub-rhomboidal (figure 27*m*), with essentially straight dorsomedial and dorsal margins that meet at a point, and a broadly convex ventrolateral margin which is kinked slightly dorsomedially at two-thirds its overall length. The proximal surface is almost entirely convex, with the exception of a shallow depression extending from the dorsomedial junction to the 'kink' on the ventrolateral margin; it is not clear if this corresponds to the embayment present in metatarsal IV in some titanosauriforms [5]. In dorsal view (figure 27*t*), both the proximal and distal ends are flared relative to the shaft, with the former more so than the latter as preserved. Both the dorsal and medial surfaces of the shaft are smoothly and shallowly convex mediolaterally along their respective lengths, with their junction only perceptible because it is slightly more strongly convex than either surface. Thus, the medial surface lacks the proximal concavity to accommodate metatarsal IV seen in some somphospondylans, including the early-branching forms *Chubutisaurus*, *Ligabuesaurus* and *Tastavinsaurus*, and the titanosaurs *Alamosaurus* and *Notocolossus* [5,174,243–245]. In dorsal view, both the medial and lateral margins are proximodistally concave, with the medial margin more strongly so than the lateral one. Along its length, the ventrolateral surface is shallowly concave, although this appears to have been exaggerated by the same distortion that

has affected the distal end. The junction between the dorsal and ventrolateral surfaces is more acute than that between the ventrolateral and medial surfaces, although the latter again seems to have been distorted. Despite the fact that the dorsal three-quarters of the distal end have been shifted dorsally relative to the ventral quarter, it appears to have been reniform *in vivo* (figure 270). The distal articular surface of metatarsal IV is approximately perpendicular to the shaft, not medially bevelled as in brachiosaurids [5].

9.5.5. Metatarsal V

Right metatarsal V is virtually complete (figure 27p-u), but has suffered slight dorsoventral deformation. In proximal view (figure 27q), the proximal end is triangular, with a very shallowly concave ventral margin and somewhat straighter dorsolateral and dorsomedial margins. It is also dorsally domed and dorsoventrally expanded relative to the shaft, contrasting with Tastavinsaurus, Muyelensaurus, and some saltasaurids, wherein it is unexpanded [38]. The proximal surface is convex in all directions. The mediolateral width of the proximal end of metatarsal V is slightly more than twice that of the distal end-similar to the condition seen among somphospondylans, including the early-branching form Tastavinsaurus and the titanosaurs Epachthosaurus and Alamosaurus, but not as pronounced as in Rapetosaurus where the mediolateral width of the proximal end is more than thrice that of the distal end [101]. In dorsal view (figure 27t), the proximal margin is shallowly and evenly mediolaterally convex, the medial and lateral margins are proximodistally concave, and the distal margin is broadly rounded. The dorsal surface of metatarsal V is mediolaterally convex along its length. Although the steepness of the convexity diminishes distally, the fact that it has this shape means that metatarsal V lacks true lateral or medial surfaces—each merges smoothly with the dorsal margin. However, the margin between the dorsal and ventral surfaces appears to have been manifested as a ridge on the lateral and medial sides. The ventral surface (figure 27u) is essentially flat, although the proximal and distal ends show some sign of wear. There is no evidence of a median tubercle or ridge on the ventral surface, distinguishing AODF 0906 from Epachthosaurus, Mendozasaurus, and saltasaurines [38]. The distal end is 'D'-shaped in distal view, with the flat side of the 'D' being the ventral margin. The distal surface is convex.

9.6. Pedal phalanges

Five pedal phalanges (including one ungual) are preserved in AODF 0906 (figure 28; table 11). All appear to be from the right foot, and they are interpreted as III-1, III-2, III-3, IV-1 and IV-2 (figure 26a). If these interpretations are correct, then digit III had three phalanges, as in brachiosaurids and *Epachthosaurus*, but unlike the early-branching somphospondylan *Gobititan*, the lognkosaurian titanosaurs *Mendozasaurus* and *Notocolossus*, and the saltasaurid titanosaur *Opisthocoelicaudia*, wherein only two phalanges are present in pedal digit III [244,247,248]. In AODF 0906, digit IV had two phalanges, as in neosauropods generally [8,100,106]. Given that all somphospondylans for which pedes are known have two phalanges (one non-ungual and one ungual) on pedal digits I and II, and none on V [2,244,249], we interpret the pedal phalangeal formula for *Diamantinasaurus* as being ?2-?2-3-2-?0.

9.6.1. Pedal phalanx III-1

Right pedal phalanx I-1 (figure 28*a*–*e*) is almost complete with the exception of a small proximomedial portion and a larger distolateral one. In proximal view (figure 28*d*), the phalanx is ovate, with the maximum dorsoventral height situated slightly lateral to the midline. The ventral margin of the proximal end is almost flat, whereas the dorsal margin is broadly convex and the lateral and medial margins more acutely so. In dorsal view (figure 28*a*), the phalanx is slightly flared proximally and would presumably have been similarly flared distally when complete. The dorsal surface is proximodistally concave and mediolaterally convex, merges smoothly with the more strongly dorsoventrally convex medial and lateral margins, and preserves a subtle bulge near the medial margin at the mid-length. The ventral surface (figure 28*c*) is shallowly concave, with the most pronounced but narrowest concavity at the distal end, where two subtle condyles are present. In distal view (figure 28*e*), the phalanx would presumably have been broadly 'D'-shaped when complete (albeit with the ventral margin shallowly concave rather than straight). In lateral (figure 28*b*) and medial views, the distal surface is inclined somewhat proximoventrally–distodorsally. It is divided into subtle lateral and medial condyles, which are divided by a shallow groove that is more pronounced ventrally than dorsally.

9.6.2. Pedal phalanx III-2

The pedal phalanx interpreted here as III-2 (figure 28*f*–*k*) is incompletely preserved but would have been relatively small, even when complete. In proximal view (figure 28*j*), the phalanx is D-shaped. The dorsal

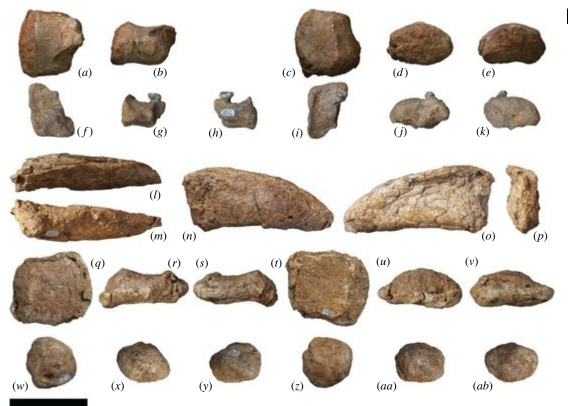


Figure 28. Diamantinasaurus matildae referred (AODF 0906) right pedal phalanges. (a-e) Right pedal phalanx III-1 in (a) dorsal, (b) lateral, (c) ventral, (d) proximal and (e) distal views. (f-k) Right pedal phalanx III-2 in (f) dorsal, (g) lateral, (h) medial, (i) ventral, (g) proximal and (k) distal views. (l-p) Right pedal ungual phalanx III-3 in (l) dorsal, (m) ventral, (n) lateral, (n) medial and (n) proximal views. (q-v) Right pedal phalanx IV-1 in (q) dorsal, (n) lateral, (n) medial, (n) unitary in (n) lateral, (n) medial views. (n) Right pedal phalanx IV-2 in (n) dorsal, (n) medial, (n) medial, (n) ventral, (n) medial, (n) medial,

Table 11. Measurements of the right pedal phalanges of AODF 0906 *Diamantinasaurus matildae*. An asterisk (*) indicates a tentative measurement based on an incomplete or distorted specimen.

	right peda	l phalanges			
measurement (mm)	III-1	III-2	III-3	IV-1	IV-2
proximodistal length	85	46	162	106	60
maximum proximal mediolateral width	93	65	25	84	60
maximum proximal dorsoventral height	57	41	69	43	46
mid-shaft mediolateral breadth	85	63		85	59
maximum distal mediolateral width	81	57		87	60
maximum distal dorsoventral height	54	31		37	48

surface (figure 28f) is proximodistally concave and mediolaterally convex, with rather acute proximal and distal flanges (visible in lateral (figure 28g) and medial (figure 28h) views); however, these might be a result of taphonomic distortion, given that this element was found in direct contact with another. The ventral surface is worn and incomplete (figure 28h), and the incomplete distal end is D-shaped (figure 28k).

9.6.3. Pedal ungual phalanx III-3

Right pedal ungual III-3 is complete but has suffered relatively minor mediolateral crushing (figure 28*l*–*p*). Relative to its dorsoventral height, it is quite elongate proximodistally, similar to the pedal unguals of some titanosaurs, including *Epachthosaurus*, *Bonatitan*, MUCPv-1533 (La Invernada titanosaur) and

Neuquensaurus [155,162,247,250], but unlike the relatively proximodistally shorter and dorsoventrally taller unguals of others, including Bonitasaura, Lognkosauria, Malawisaurus, Mnyamawamtuka, Opisthocoelicaudia and Rapetosaurus [115,225,236,244,251–254]. The ungual in AODF 0906 lacks the strong curvature seen in the pedal unguals of the brachiosaurid Cedarosaurus [255] and the non-titanosaurian somphospondylan Ligabuesaurus [174]. In proximal view (figure 28p), the ungual is significantly taller dorsoventrally than it is wide mediolaterally, with the lateral margin somewhat concave and the medial one essentially straight. The proximal articular surface is bevelled proximolaterally relative to the long axis of the ungual. In lateral (figure 28n) and medial views (figure 28o), the ungual shows a gradual but distinct distal taper, the proximal margin is straight, the dorsal margin is convex, and the ventral surface is concave. The dorsal surface (figure 281) is manifested as a sharp ridge, although this appears to have been exaggerated somewhat by post mortem deformation. The lateral surface (figure 28n) is dorsoventrally convex but undulates proximodistally, such that the mid-section is convex and the proximal and distal ends show some degree of concavity. The medial surface (figure 280) is dorsoventrally and proximodistally convex, although the dorsal two-thirds have suffered deformation such that they are now partly concave. The medial surface contributes more to the ventral surface than the lateral one, with the junction between the two manifested as a ridge that more or less follows the lateral margin. Neither the medial nor lateral surface has a vascular groove, differentiating it from several titanosauriform taxa, including Paludititan nalatzensis [256]. The lack of vascular grooves also differentiates this ungual from the manual unguals of the Diamantinasaurus matildae holotype specimen (AODF 0603) and juvenile referred specimen (AODF 0663) [9,37,44]. The ventral margin appears to show a small tuberosity near the distal end, a feature that characterizes most somphospondylans, including the non-titanosaurians Euhelopus, Tangvayosaurus, Tastavinsaurus and Gobititan, and the titanosaurs Malawisaurus, Epachthosaurus, Mendozasaurus and Muyelensaurus [38,101,220].

9.6.4. Pedal phalanx IV-1

Pedal phalanx IV-1 is complete but has suffered from slight dorsoventral compression (figure 28q-v); nevertheless, it was probably somewhat dorsoventrally compressed in life, as is the case in pedal phalanges IV-1 of *Bonitasaura* [257] and *Mendozasaurus* [236]. In proximal view (figure 28u), the phalanx is D-shaped with a flat ventral surface. The dorsal surface (figure 28q) is mediolaterally convex and very shallowly proximodistally concave. The lateral and medial margins are shallowly concave proximodistally in dorsal view, and the former has a subtle node at the mid-length. The distal end is convex in dorsal view. The medial (figure 28s) and lateral (figure 28r) surfaces are acutely dorsoventrally convex. The ventral surface is concave (figure 28t). The distal end is slightly crushed, but the distal surface would have been convex. In distal view (figure 28v), the phalanx is D-shaped but flattened, with the apex slightly lateral to the midline.

9.6.5. Pedal phalanx IV-2

Pedal phalanx IV-2 is rounded and relatively featureless (figure 28w–ab). In proximal view (figure 28aa), the phalanx is shaped like a tall 'D', with the flat margin being the ventral one. The proximal surface is proximodorsally–distoventrally inclined, as can be seen in lateral view (figure 28x). Whereas the lateral and medial margins are straight in dorsal view, the distal margin is convex. The dorsal surface is proximodistally concave and medialaterally convex. Both the lateral (figure 28x) and medial (figure 28y) surfaces are smoothly proximodistally concave, but the lateral is dorsoventrally convex and the medial is dorsoventrally concave (such that it looks almost 'kinked'). The ventral surface (figure 28z) is convex. The distal surface is rounded (figure 28ab).

10. Phylogenetic results

Analysis under equal weights results in 300 960 most parsimonious trees (MPTs) of length 2679 steps. The topology (figure 29) is essentially unchanged from that presented in Poropat *et al.* [41]. AODF 0906 is recovered as a member of Diamantinasauria, forming a polytomy with the *Diamantinasaurus matildae* holotype (AODF 0603), AODF 0836, *Savannasaurus* and *Sarmientosaurus*. Diamantinasauria has a Bremer support of 3 and is recovered as a non-lithostrotian titanosaur clade. It would take trees three steps longer to place Diamantinasauria outside of Titanosauria.

Analysis using extended implied weights produces 588 MPTs of length 139.9 steps. As with the equal weights analysis, the same five operational taxonomic units (OTUs) form a diamantinasaurian polytomy.



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Figure 29. Equal weights phylogenetic analysis. Agreement subtree showing the relationships within Titanosauriformes based on the results of the equal weights phylogenetic analysis. Brachiosauridae has been collapsed into a single lineage. The names of Australian taxa/specimens are in boldface.

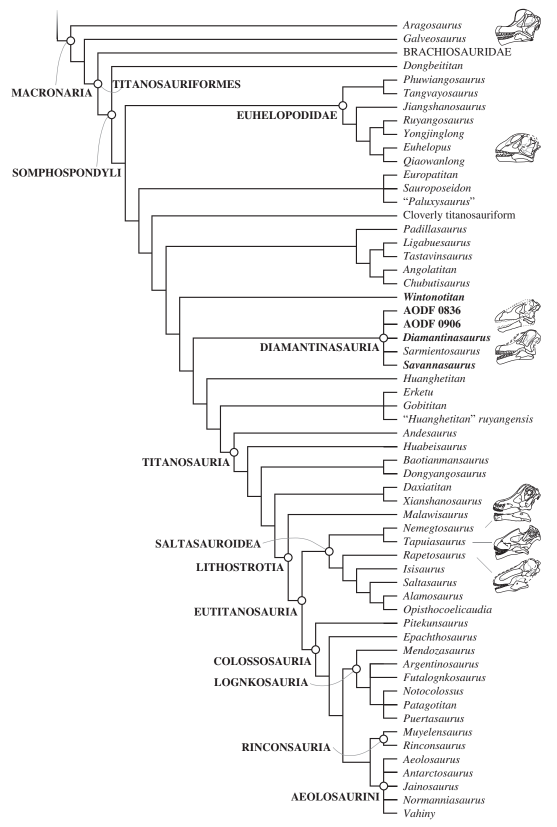


Figure 30. Extended implied weights phylogenetic analysis. Agreement subtree showing the relationships within Titanosauriformes based on the results of the equal weights phylogenetic analysis. Brachiosauridae has been collapsed into a single lineage. The names of Australian taxa/specimens are in boldface.

The overall topology (figure 30) is little changed from the equivalent analysis in Poropat *et al.* [41]; however, Diamantinasauria is recovered outside of Titanosauria, with a small number of East Asian taxa placed as closer relatives to the titanosaurian radiation (figure 30).

Analyses were also run with AODF 0603, AODF 0836 and AODF 0906 combined as a single OTU. The topology of the trees was identical, but Diamantinasauria remained a polytomy.

11. Discussion

11.1. The phylogenetic composition and position of Diamantinasauria

As noted by Carballido *et al.* [1], the taxa comprising the clade Diamantinasauria have remained relatively consistent since the initial publication of *Savannasaurus elliottorum* and the referred specimen of *Diamantinasaurus matildae* (AODF 0836) by Poropat *et al.* [38]. *Sarmientosaurus musacchioi* was first resolved within Diamantinasauria by Poropat *et al.* [41], and this result is replicated herein—with additional support from features of the AODF 0906 skull, notably the presence of a posterior tongue-like process on the quadratojugal, which was identified as an autapomorphy of *Sarmientosaurus* by Martínez *et al.* [23], but can now be regarded as a diamantinasaurian synapomorphy. The only other taxa to have been resolved within Diamantinasauria are *Wintonotitan wattsi* and *Australotitan cooperensis*, by Hocknull *et al.* [34], and both of these taxa derive from the same stratigraphic unit as *Diamantinasaurus* and *Savannasaurus* (i.e. the Winton Formation in the Eromanga Basin). However, we note that *Wintonotitan* was only resolved within Diamantinasauria by Hocknull *et al.* [34] when they ran modified analyses in which large numbers of taxa (non-macronarian taxa or post-Cenomanian taxa) or characters (all cranial and axial characters) were excluded. As such, we contend that there is currently limited support for the inclusion of *Wintonotitan* within Diamantinasauria.

Whereas the content of Diamantinasauria has remained fairly consistent, the phylogenetic position of the clade within Somphospondyli has not [1]. Prior to the description of Savannasaurus elliottorum [38], Diamantinasaurus was generally regarded as a lithostrotian titanosaur, specifically either as a member of Opisthocoelicaudiinae within Saltasauridae [9,37], the sister taxon to Saltasauridae [9,21], or the sister taxon to Tapuiasaurus [37]. However, in other analyses it was resolved within Titanosauria but just outside Lithostrotia, or within Somphospondyli but outside Titanosauria [101]. The inclusion of Savannasaurus elliottorum and an additional Diamantinasaurus specimen (AODF 0836) within an updated and expanded phylogenetic analysis based on that of Mannion et al. [101] saw the two Diamantinasaurus specimens and Savannasaurus consistently resolved as a clade, which was situated near the base of Titanosauria in equal weights analyses, but within Lithostrotia (and close to Saltasauridae) in implied weights analyses [38). Subsequent analyses based on this dataset have generally returned the same result [12,90,236,258–260], with a few notable exceptions [6,90,249,261], as discussed by Poropat et al. [40]. An early-branching position for Diamantinasauria within Titanosauria has received additional support in several recent studies [34,40,41,43]. Nevertheless, it is plausible that the presence of several seemingly plesiomorphic titanosaurian characters [40,41] simply indicates that Diamantinasauria occupies a position outside Titanosauria, rather than an early-branching position within, as recovered in our analyses applying extended implied weighting. This interpretation is supported by at least one newly identified character state that would be plesiomorphic within Titanosauria or a slightly more inclusive clade—the presence of five, rather than at least six, sacral vertebrae in Diamantinasaurus matildae. We discuss the possible implications of this feature, as well as aspects pertaining to the pes and skull, below.

11.2. Implications of the presence of five sacral vertebrae in Diamantinasauria

Currently, Diamantinasaurus is the only diamantinasaurian for which we can evaluate the number of sacral vertebrae, with three specimens consistently demonstrating that the sacrum of this genus was composed of only five vertebrae. As noted above, no other titanosaur is characterized by fewer than six sacral vertebrae. By itself, this might merely indicate that five sacral vertebrae is the plesiomorphic condition in titanosaurs, given that we do not know the sacral counts in other early diverging titanosaurs, including Andesaurus [262], Choconsaurus [159] and Mnyamawamtuka [254]. However, as noted above, several somphospondylans that almost certainly lie outside of Titanosauria are characterized by six-vertebrae sacra. Of those somphospondylans that are known to have sacra composed of five vertebrae, most of these taxa (i.e. Sauroposeidon, Sibirotitan and Tastavinsaurus) are typically recovered as relatively early diverging members of the clade (e.g. [219]; this study), which could be interpreted as retention of the plesiomorphic eusauropod condition prior to the acquisition of a sixth sacral vertebra at a more nested node within Somphospondyli. Nevertheless, Huanghetitan liujiaxiaensis, which is also characterized by a five-vertebrae sacrum [218], is consistently recovered as one of the closest relatives of Titanosauria in our analyses. Furthermore, more detailed evaluation of somphospondylan interrelationships (figures 29 and 30) indicates that some lineages characterized by six-vertebrae sacra (namely taxa placed within Euhelopodidae) are

typically recovered as more distantly related to Titanosauria than some taxa with five-vertebrae sacra (e.g. *Sauroposeidon*). This suggests that there have either been a number of independent acquisitions of a six-vertebrae sacrum among Somphospondyli and/or that there have been numerous reversals to a five-vertebrae sacrum. The fact that six-vertebrae sacra have also been described from non-titanosauriform sauropods—notably, a specimen of the early-branching macronarian *Camarasaurus* [263], the enigmatic neosauropod *'Apatosaurus' minimus* [264–266], and the non-neosauropodan eusauropod *Klamelisaurus gobiensis* [267]—further highlights the likely high levels of plasticity of this characteristic. As such, we suggest that the number of sacral vertebrae is of relatively limited importance in determining whether Diamantinasauria lies within or outside of Titanosauria.

11.3. Implications of the diamantinasaurian pes for titanosaurian pedal evolution

Several previous studies have discussed the evolution of the titanosauriform pes [2,244,247,249, 251,253,268,269]. These authors have noted that there is a trend towards pedal phalangeal reduction in the clade, particularly within Titanosauria (table 12 in appendix). All titanosaurs have only two phalanges (including an ungual) on pedal digits I and II. Somphospondylans that unequivocally lie outside of Titanosauria, and that preserve near-complete pedes, are rare, but Tangvayosaurus and Tastavinsaurus have both been reconstructed with three phalanges on digit II [239,296]. By contrast, the non-titanosaurian somphospondylan Gobititan has only two phalanges on this digit [290]. Most titanosaurs have two phalanges on pedal digit III, including Mendozasaurus, Notocolossus, Opisthocoelicaudia, UNCUYO-LD 313 (Agua del Padrillo titanosaur) and MUCPv-1533 (La Invernada titanosaur). However, some early deriving forms, such as Epachthosaurus and (probably) Diamantinasaurus, have three phalanges on this digit. Three or more phalanges also characterize digit III of Tangvayosaurus and Tastavinsaurus, whereas Gobititan only has two phalanges on this digit. Nearly all somphospondylans have only two phalanges on digit IV; the only known possible exception is Opisthocoelicaudia, which only preserves one phalanx on this digit [225], although it has been inferred to have possessed a second based on the morphology of the distal end of that phalanx [249]. All of the few titanosaurs that preserve near-complete pedes lack phalanges on pedal digit V. Gobititan, Tangvayosaurus and Tastavinsaurus all have one or two phalanges on pedal digit V [239,290,296].

If the above interpretation of the pedal phalanges of AODF 0906 is correct, and irrespective of whether Diamantinasauria occupies an early-branching position within Titanosauria, or lies just outside of Titanosauria, then *Diamantinasaurus* seemingly fits neatly into the pattern of phalangeal reduction seen within Somphospondyli. Nevertheless, more specimens of taxa just outside of Titanosauria, as well as early members of this clade, are required to test this, including whether phalangeal number might be relatively plastic.

11.4. Implications of the diamantinasaurian skull for titanosaurian cranial evolution

The evolution of titanosaurian sauropod skull disparity is poorly understood. Few taxa are known from partial or complete skulls, and those that are date either to the late Early Cretaceous (Aptian–Albian: *Tapuiasaurus* [21,22] and *Malawisaurus* [115,120,302]), the early Late Cretaceous (Cenomanian: *Sarmientosaurus* [23]), or the latest Cretaceous (Maastrichtian: *Nemegtosaurus* [18,20] and *Rapetosaurus* [16,17]). The absence of titanosaur skull material prior to the Aptian and relative scarcity from the Turonian–Campanian interval—the incomplete crania of *Antarctosaurus* [145,162], *Bonitasaura* [141] and *Quaesitosaurus* [19], and the embryonic skulls from Auca Mahuevo [129,130] notwithstanding—remains the greatest impediment to understanding titanosaurian skull evolution.

The skulls of early-branching titanosaurs, typified by *Diamantinasaurus* and *Sarmientosaurus* [23], are quite different from those of phylogenetically nested titanosaurs [17,20–22]. The skulls of diamantinasaurians have a more undulatory lateral profile (with the back of the skull taller and somewhat raised relative to the snout), more robust dentigerous elements, and compressed-cone—chisel-like teeth. In all of these ways, the skulls of these early-branching titanosaurs are similar to those of brachiosaurids [24–26,133]. Current evidence implies that diamantinasaurians, as well as other early-branching titanosaurs, such as *Choconsaurus* [159], had rounded jaws, similar to those of *Camarasaurus* [118] and brachiosaurids [24–26].

More phylogenetically nested titanosaurs have elongate, low skulls with pencil- or chisel-like teeth confined to the anterior part of the snout [17,20–22]. In all of these ways, their skulls are reminiscent of those of diplodocoids [161,303], so much so that the skulls of the titanosaurs *Antarctosaurus*, *Nemegtosaurus* and *Quaesitosaurus* were once thought to be from diplodocoids [18,19,100,106,162,304]. However, the titanosaurian affinities of these skulls were demonstrated beyond reasonable doubt through a detailed reappraisal of the specimens [20,305], and confirmed through the subsequent

discovery of titanosaurian postcrania at the *Nemegtosaurus* type site [306]. In the interim, the discovery of near-complete skulls associated with undisputed titanosaurian postcranial remains (all attributed to *Rapetosaurus*) settled the issue [3,17].

Among 'derived' titanosaurs, there is evidence for two main jaw shapes: jaws with rounded ends, like those of *Ampelosaurus, Karongasaurus, Malawisaurus, Mansourasaurus, Nemegtosaurus, Quaesitosaurus, Rapetosaurus* and *Tapuiasaurus* [17–22,115,160,168]; and squared-off jaws, like those of *Antarctosaurus, Baalsaurus, Bonitasaura* and *Brasilotitan* [141,145,162–165]. This disparity parallels that seen in diplodocids [105]: some have rounded jaws, such as the dicraeosaurids *Suuwassea emilieae* and *Dicraeosaurus hansemanni* [25,166]; some have squared-off jaws, such as the diplodocids *Apatosaurus louisae, Diplodocus longus* and *Tornieria africana*, and the rebbachisaurid *Lavocatisaurus agrioensis* [25,213,303,307]; and others have squared-off and transversely expanded jaws, namely the rebbachisaurid *Nigersaurus* [110,308,309].

Whereas several brachiosaurids are known from skulls of varying levels of completeness (e.g. 'Brachiosaurus' [24], Giraffatitan [25], Abydosaurus [26]), few non-titanosaurian somphospondylan taxa are represented by any cranial remains at all. Several non-titanosaurian somphospondylans preserve only teeth, including Astrophocaudia slaughteri [293], Borealosaurus wimani [310], Europatitan eastwoodi [311], Huabeisaurus allocotus [223,312], Sibirotitan astrosacralis [219] and Yongjinglong datangi [173]. Ligabuesaurus leanzai preserves teeth and a fragmentary maxilla [174,291,313], whereas teeth and a braincase are preserved in both Mongolosaurus haplodon [152,314] and Tambatitanis amicitiae [151,315]. Teeth are preserved in several specimens of Phuwiangosaurus sirindhornae, with one specimen also preserving a partial skull [82,149,215,316–319]. Euhelopus zdanskyi preserves a nearly complete skull, along with teeth, but lacks the braincase [111–114,320]; furthermore, it is possible that it lies outside Neosauropoda [267]. By far the most complete nontitanosaurian somphospondylan skull is the holotype specimen of Liaoningotitan sinensis, although this has only been briefly described [321]. This skull has suffered substantial taphonomic distortion, but the dentaries are effectively mirror images of each other, implying that they probably reflect their original morphology; if so, then Liaoningotitan had jaws with somewhat squared-off ends, paralleling some diplodocids, rebbachisaurids, and some titanosaurs, and implying even greater plasticity in sauropod jaw morphology.

The similarity between the skulls of the diamantinasaurians *Diamantinasaurus* and *Sarmientosaurus* implies morphological conservatism in this clade of early-branching titanosaurs (figure 31), starkly contrasting with the disparity seen in the snout shapes of more phylogenetically nested titanosaurs. The similarities between the skulls of *Diamantinasaurus* and *Sarmientosaurus* and those of brachiosaurids are quite striking, as are the differences they present when compared with the skulls of the non-titanosaurian somphospondylan *Liaoningotitan*, and those of other titanosaurs, including the stratigraphically older *Tapuiasaurus* and *Malawisaurus*. Although the disparity in maxillary and dentary alveolus count (nine maxilla, 11 dentary in *Diamantinasaurus*; 11 or 12 maxilla, 13 dentary in *Sarmientosaurus*), and the differing number of ossified exits on each side for cranial nerves V and XII (two for CN V, one for CN XII in *Diamantinasaurus*; three for CN V, two for CN XII in *Sarmientosaurus*) indicate some disparity between the two taxa, a similar level of disparity has been identified within some sauropod genera. *Camanasaurus*—one of the few sauropod genera known from multiple skulls—shows variability in all but one of these characters: maxillary alveolus count ranges from 8 to 10 [118], dentary alveolus count ranges from 12 to 13 [118], and either one or two exits for CN XII can be present on each side [118,138,153,154].

The relative conservatism seen in the skulls of diamantinasaurian titanosaurs might be a reflection of the restricted spatio-temporal range of the clade (as presently understood). Both *Sarmientosaurus* and *Diamantinasaurus* lived at approximately the same palaeolatitude (approx. 50° S for *Sarmientosaurus*, approx. 48° S for *Diamantinasaurus* [322]) during the Cenomanian–Turonian, which is regarded as the warmest interval the Earth has endured in the past 150 Myr [323]. By contrast, *Tapuiasaurus* and *Malawisaurus*, which are geologically older (Aptian), but more phylogenetically nested than Diamantinasauria, were discovered at sites that during the Aptian would have been at approximately 15° S and approximately 30° S, respectively [322].

Recent work on Cenomanian–Turonian floras in South America and Zealandia [324] has strongly supported the hypothesis that several, largely latitudinally controlled floral provinces existed in Gondwana during the early Late Cretaceous [325]. The latitudinal band at 50° S was dominated by gymnosperms in both South America and Australia; however, whereas cheirolepidacean gymnosperms were dominant in the former region (constituting more than 80% of the samples studied), gymnosperms other than cheirolepidaceans were most abundant in northern Australia (although cheirolepidaceans still comprised approx. 25% of the flora), and angiosperms had become a notable component of the flora as well [324]. The plant macrofossil record of the Winton Formation indicates that gymnosperms and angiosperms were co-dominant [326]; the findings of Santamarina *et al.* [324] corroborate this. Given that diamantinasaurians have been interpreted as mid-level feeders (*sensu* Whitlock [105]) based on tooth



Figure 31. Life restoration of the head and neck of *Diamantinasaurus matildae*, based on AODF 0906 (referred skull) and AODF 0836 (referred partial skull and anterior cervical vertebrae). Artwork © Elena Marian.

wear analyses [43,327], and given the variety of plants available within the 1–10 m height bracket to diamantinasaurians in northeast Australia (summarized in [43]), it is tempting to attribute at least some of the subtle differences between the skulls of *Sarmientosaurus* and *Diamantinasaurus*—namely those related to alveolus count and tooth morphology—to subtle differences in their diets. The marked difference between the skulls of diamantinasaurians and more derived titanosaurs presumably reflects a greater degree of dietary divergence, but this requires further testing.

12. Conclusion

The new specimen of the sauropod dinosaur *Diamantinasaurus matildae* described herein preserves numerous skeletal elements not previously known for this taxon, primarily a partial skull and an incomplete pes. When this new specimen is considered in conjunction with the holotype specimen (presacral vertebrae, sacrum, pectoral girdle, forelimbs, pelvic girdle, right hind limb), and the two previously referred specimens AODF 0836 (partial skull, presacral vertebrae, appendicular elements) and AODF 663 (presacral vertebrae, appendicular elements), the osteology of *Diamantinasaurus* is now among the most completely known of any somphospondylan.

The skull of *Diamantinasaurus matildae* is herein shown to be strikingly similar to that of the approximately coeval *Sarmientosaurus musacchioi* from Argentina. This supports our previous proposal [41] of a close evolutionary relationship between these taxa (forming the clade Diamantinasauria), and is borne out with renewed support in the revised phylogenetic analyses presented herein. Although the position of Diamantinasauria within Somphospondyli remains uncertain, our analyses imply that it lies near the base of Titanosauria: probably just within (meaning that diamantinasaurians were early-branching titanosaurs), or possibly just outside (meaning they were non-titanosaurian somphospondylans), that clade. Future discoveries in the Winton Formation will hopefully fill the remaining gaps in our knowledge of the skeleton of *Diamantinasaurus matildae*, thereby enhancing its importance for future research into the early evolution of Titanosauria.

Data accessibility. The electronic supplementary material, comprising three-dimensional models (derived from surface scans) of the AODF 0906 skull, hindquarters and right pes, three-dimensional models (derived from surface scans) of the AODF 0836 skull, and the CT and Synchrotron data of the AODF 0906 skull elements, is available from MorphoSource: https://www.morphosource.org/projects/000486235.

The data are provided in electronic supplementary material [328].

Authors' contributions. S.F.P.: conceptualization, data curation, formal analysis, investigation, methodology, project administration, visualization, writing—original draft, writing—review and editing; P.D.M.: formal analysis, methodology, validation, writing—original draft, writing—review and editing; S.L.R.: data curation, investigation, software, writing—review and editing; R.J.D.: data curation, investigation, software, visualization, writing—review and editing; J.J.B.: data curation, formal analysis, investigation, resources, software, writing—review and editing; T.S.: data curation, resources, writing—review and editing; D.A.E.: data curation, funding acquisition, resources, software, writing—review and editing.

All authors gave final approval for publication and agreed to be held accountable for the work performed therein. **Conflict of interest declaration.** We declare we have no competing interests.

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Appendix

Characters 1–548 (and their scorings) follow those of Mannion *et al.* [90], and those for C549–552 follow those of Poropat *et al.* [41], except where stated below. However, C550, as proposed by Poropat *et al.* [41], was incorrectly worded. This character statement is modified as follows (note that the scores are unaffected):

C550. Cranial nerve V, number of external openings: two or more on each side (0); one on each side (1) ([23,41]; modified here).

We also add the following new characters (C### denotes the character number):

Table 12. Cretaceous titanosauriform taxa for which pedal elements are known. Reports only known to the authors from published abstracts have been omitted. L, left; I, lower; m, middle; mt, metatarsal; pp, pedal phalanx; pu, pedal ungual; R, right; u, upper.

							lengun	
taxon	specimen	locality	formation	age range	pedal elements present	phalangeal formula	formula	reference(s)
Lithostrotia indet.	TMM 45890-2	Texas, USA	Black Peaks	u. Maastrichtian	R pu	<i>د</i> .	د .	Fronimos & Lehman [270]
Rapetosaurus krausei	FMNH PR 2209	Mahajanga, Madagascar	Маеvarano	Maastrichtian	L mt I—IV, R mt V, four pp (including two pu)	2-1-1-1-1	<i>i-;-;-</i> ;-1	Curry Rogers [251]
Rapetosaurus krausei	UA 9998	Mahajanga, Madagascar	Maevarano	Maastrichtian	R mt l	į	į	Curry Rogers [251]
Titanosauria indet. (formerly ? Antarctosaurus sp.)	GSI K27/509	Maharashtra, India	Lameta	Maastrichtian	L mt l	į	,	Huene & Matley [271]
Alamosaurus sanjuanensis	TMM 43621 1	Texas, USA	Black Peaks	Maastrichtian	Mt	į	į	Lehman & Coulson [272]
Alamosaurus sanjuanensis	TMM 42495	Texas, USA	Javelina / Black Peaks	Maastrichtian	Mt	į	į	Lehman & Coulson [272]
Alamosaurus sanjuanensis	TMM 43598	Texas, USA	Javelina / Black Peaks	Maastrichtian	Mt II	į	į	Woodward & Lehman [273]
?Alamosaurus sanjuanensis	NMMNH P-49967	New Mexico, USA	Ojo Alamo	Maastrichtian	R mt I–V, five R pp	į	;	D'Emic <i>et al.</i> [245]
Paludititan nalatzensis UBB NVM1	UBB NVM1	Haţeg Basin, Romania	Sânpetru	Maastrichtian	two pu	į	į	Csiki <i>et al.</i> [256]

(Continued.)

Table 12. (Continued.)

taxon	specimen	locality	formation	age range	pedal elements present	phalangeal formula	ungual formula	reference(s)
Opisthocoelicaudia skarzynskii	Z. PAL MgD-1/48	Ömnögovi, Mongolia	Nemegt	Maastrichtian	R mt I–V, L mt I–II and IV, seven R pp (including three pu)	2-2-2-1-0 [225] 2-2-2-2-0 (inferred by González Riga <i>et al.,</i> [249] on the basis of the morphology of pp IV-1)	1-1-1-0 0	Borsuk-Białynicka [225]
Saltasaurus loricatus	PVL 4017 various	Salta, Argentina	Lecho	?Campanian— Maastrichtian	seven mt (from more than one individual)	?	į	Powell [144,145]; S.F.P. and P.D.M. <i>pers. obs.</i> 2013
Aeolosaurus sp.	MPCA-Pv 27178	Río Negro, Argentina	Allen	Campanian— Maastrichtian	mt l	į	į	García & Salgado [274]
Bonatitan reigi	'Individual A' (MACN PV RN 821, 1061 and unnumbered)	Rio Negro, Argentina	Allen	Campanian– Maastrichtian	L mt I, R mt III, mt V	<u>.</u>	į	Martinelli & Forasiepi [146]; Salgado et al. [155]; S.F.P. and P.D.M. pers. obs. 2013
Bonatitan reigi	Individual D' (MACN PV RN 821, 1061 and unnumbered)	Rio Negro, Argentina	Allen	Campanian— Maastrichtian	mt I, III (×2), V (×2), pu	}	į	Martinelli & Forasiepi [146]; Salgado et al. [155]; S.F.P. and P.D.M. pers. obs. 2013
Titanosauria indet.	MUCPv-1533 ('La Invernada titanosaur')	Neuquén, Argentina	Allen	Campanian— Maastrichtian	L mt I—V, eight pp (induding three pu)	2-2-2-0	1-1-1-0-	González Riga et al. [247,253]

(Continued.)

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Table 12. (Continued.)

MDE G various Agentina four Administration (possibly from more inclindual) ? ? ratiosaurus MDE G various Santa Cruz, Argentina Cerro Fortaleza (Ampanian - Infriedual) R mt -H.I. R pu Ir 2 ? ? ? có MDE G various Languedro- Portaleza (Ampanian - Infriedual) Manter Rouges Campanian - Infriedual R mt -H.I. R pu Ir 2 ? ? ? có MDE G various Languedro- Pour Rouges Campanian - Infriedual mt, pp ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? <t< th=""><th>taxon</th><th>specimen</th><th>locality</th><th>formation</th><th>age range</th><th>pedal elements present</th><th>phalangeal formula</th><th>ungual formula</th><th>reference(s)</th></t<>	taxon	specimen	locality	formation	age range	pedal elements present	phalangeal formula	ungual formula	reference(s)
roam MPM PV 1156 Santa Cruz, Argentina Cerro Fortaleza (Ampanian – Masastrichtan Manes Rouges (Ampanian – mr, pp R mt I–II, R pu I–2 ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ?	ostrotia indet. formerly <i>Laplatasaurus</i> rraukanicus)	MLP CS 1203, 1217, 2002	Río Negro, Argentina	Allen	Campanian— Maastrichtian	R mt II and ?V, L pu (possibly from more than one individual)	¿	<i>`</i>	Huene [162]; Gallina & Otero [275]
MOE G various Languedoc- Mannes Rouges Campanian- mt, pp ? ? ? ? ? ? % % % %	dnoughtus schrani	MPM PV 1156	Santa Cruz, Argentina	Cerro Fortaleza	Campanian— Maastrichtian	R mt I–II, R pu I-2	į	į	Lacovara <i>et al.</i> [252]; Ullmann & Lacovara [276]
ace MCNA 14474 Alava, Spain Vitoria u. Campanian—I. L mt I—IV ? ? MPCA-Pv 27100 Río Negro, Los Alamitos u. Campanian—I. L mt I—IV ? ? MPCA-Pv 27100 Río Negro, Los Alamitos u. Campanian—I. L mt I—IV ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? </td <td>Ampelosaurus atadis</td> <td>MDE G various</td> <td>Languedoc- Roussillon, France</td> <td>Marnes Rouges Inférieures</td> <td>Campanian— Maastrichtian</td> <td>mt, pp</td> <td>į</td> <td>į</td> <td>Le Loeuff [168,277]</td>	Ampelosaurus atadis	MDE G various	Languedoc- Roussillon, France	Marnes Rouges Inférieures	Campanian— Maastrichtian	mt, pp	į	į	Le Loeuff [168,277]
MPCA-Pv 27100 Rio Negro, Argentina Los Alamitos u. Campanian—I. Lmt I—IV ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? ? </td <td>nosaurus astibiae</td> <td>MCNA 14474</td> <td>Alava, Spain</td> <td>Vitoria</td> <td>u. Campanian—I. Maastrichtian</td> <td>L mt III</td> <td>į</td> <td>į</td> <td>Díez Díaz et al. [278]</td>	nosaurus astibiae	MCNA 14474	Alava, Spain	Vitoria	u. Campanian—I. Maastrichtian	L mt III	į	į	Díez Díaz et al. [278]
UNPSJB-PV 1051 Chubut, Argentina Lago Colhué Gampanian—? two mt and one pp ? ? ? Huapi Maastrichtian Phagus MPM 125R São Paulo, Brazil Adamantina u. Campanian pu ? ? ? Rhône, France sandstones MMS/VBN.93.10 Bouches du Begudian' u. Campanian L mt l Rhône, France sandstones Andron MUVP 2000 Dakhla Oasis, Quseir Campanian R mt I, II or IV, and III ? ? Egypt Argentina Argentina	osaurus sp.	MPCA-Pv 27100	Río Negro, Argentina	Los Alamitos	u. Campanian—I. Maastrichtian	L mt I–IV	į	į	Salgado <i>et al.</i> [97]
ophagus MPM 125R São Paulo, Brazil Adamantina u. Campanian L mt l ? ? MMS/VBN.93.10 Bouches du Begudian' u. Campanian L mt l ? ? Shahinae MUVP 2000 Dakhla Oasis, Quseir Campanian R mt l, II or IV, and III ? ? Egypt Egypt Anacleto I. Campanian L mt I–IV (and V?) ? ?	ostrotia indet.	UNPSJB-PV 1051	Chubut, Argentina	Lago Colhué Huapi	Campanian—? Maastrichtian	two mt and one pp	į	į	lbiricu <i>et al.</i> [279]
MMS/VBN.93.10 Bouches du Begudian' u. Campanian L mt l Rhône, France sandstones shahinae MUVP 2000 Dakhla Oasis, Quseir Campanian R mt I, II or IV, and III ? ? ? Egypt MACN 6904 Río Negro, Anacleto I. Campanian L mt I–IV (and V?) ? ?	lotitan nemophagus	MPM 125R	São Paulo, Brazil	Adamantina	u. Campanian	nd	į	į	Machado <i>et al.</i> [164]
s <i>shahinae</i> MUVP 2000 Dakhla Oasis, Quseir Campanian R mt I, II or IV, and III ? ? ? Egypt MACN 6904 Río Negro, Anacleto I. Campanian L mt I–IV (and V?) ? ?	ganosaurus elauciensis	MMS/VBN.93.10	Bouches du Rhône, France	`Begudian' sandstones	u. Campanian	L mt I	,	į	Garcia <i>et al.</i> [280]; Díez Díaz <i>et al.</i> [13]
MACN 6904 Río Negro, Anacleto I. Campanian L mt I—IV (and V?) ? ? ? ?	sourasaurus shahinae	MUVP 2000	Dakhla Oasis, Egypt	Quseir	Campanian	R mt I, II or IV, and III	į	į	Sallam <i>et al.</i> [160]
	uns	MACN 6904		Anacleto	l. Campanian	L mt I–IV (and V?)	į	;	Huene [162]; Powell [145]

taxon	specimen	locality	formation	age range	pedal elements present	phalangeal formula	ungual formula	reference(s)
cf. Bonitasaura sp. (formerly Laplatasaurus araukanicus)	MLP Av. 2102, 2111, 2113	Neuquén, Argentina	Anadeto	l. Campanian	L mt I, R mt III—IV (possibly from more than one individual)	<i>`</i>	۲.	Huene [162]; Gallina & Otero [275]
Neuquensaurus australis (including N. robustus)	MLP CS various	Río Negro, Argentina	Anacleto	I. Campanian	mt I (x3), II (x4), III (x3) IV (x4) and V (x4), seven pp (including two pu) (from more than one individual)	į	į.	Huene [162]; Powell [145]; Otero [240]
Neuquensaurus australis	MCS 10	Río Negro, Argentina	Anacleto	l. Campanian	one mt	į	į	Salgado <i>et al.</i> [226]; Otero [240]
Lithostrotia indet.	MACN	Río Negro, Argentina	Anacleto	l. Campanian	two mt	į	į	Huene [162]; Gallina & Otero [275]
Bonitasaura salgadoi	MPCA-Pv 460	Rio Negro, Argentina	Bajo de la Cama	Santonian	L mt I–III and V, R mt I–II and IV–V, three L pp (induding one pu)	'	į	Apesteguía [163]; Gallina and Apesteguía [257]
Notocolossus gonzalezparejasi	UNCUYO-LD 302	Mendoza, Argentina	Plottier	u. Coniacian—I. Santonian	R mt I–V, eight pp (including three pu)	2-2-2-0	1-1-1-0- 0	González Riga <i>et al.</i> [244,253]
Titanosauria indet.	UNCUYO-LD 313 (Agua del Padrillo titanosaur)	Mendoza, Argentina	Plottier	u. Coniacian–I. Santonian	ier u. Coniacian—I. L mt I—V, eight pp 2-2-2-0 1-1-1-0- González Riga <i>et al.</i> Santonian (including three pu) 0 [244,253]	2-2-2-0	1-1-1-0- 0	González Riga <i>et al.</i> [244,253]

 Table 12. (Continued.)

(Continued.)

Table 12. (Continued.)

taxon	specimen	locality	formation	age range	pedal elements present	phalangeal formula	ungual formula	reference(s)
Kaijutitan maui	MAU-Pv-CM-522	Neuquén, Argentina	Sierra Barossa	u. Coniacian	R mt II	į	;	Filippi <i>et al.</i> [281]
Mendozasaurus neguyelap	IANIGLA-PV <i>77–79,</i> 100, 153	Mendoza, Argentina	Sierra Barossa	m.—u. Coniacian	mt I (×3), II, III (×2), IV (×2) and V (×4), 10 pp (including three pu) (from more than one individual)	į	į	González Riga [236,282]
Saltasauridae indet.	FC-DPV 1900	Paysandú, Uruguay	Guichón	Turonian— Santonian	several mt and pp	į	;	Soto <i>et al.</i> [283]
Muyelensaurus pecheni	MRS Pv various	Neuquén, Argentina	Portezuelo	m.—u. Turonian	15 mt and 11 pp (from more than one individual)	į	į.	Calvo <i>et al.</i> [116]
Titanosauria indet.	IZANUZ 27, 248 and 636; ZIN PH 672/16	Navoi, Uzbekistan	Bissekty	Turonian	$mt \mid (x,2) \text{ and } \ (x,2)$ (probably from more than one individual)	į	į	Sues <i>et al.</i> [284]
Epachthosaurus sciuttoi	UNPSJB-PV 920	Chubut, Argentina	Lower Bajo Barreal	u. Cenomanian—I. Turonian	L and R mt I—V, 18 pp (including six pu)	2-2-3-2-0	1-1-1-0- 0	Martinez <i>et al.</i> [250]; S.F.P. and P.D.M. <i>pers.</i> obs. 2013
Savannasaurus elliottorum	AODF 0660	Queensland, Australia	Winton	Cenomanian	L mt III	į	į	Poropat <i>et al.</i> [38,40]
Diamantinasaurus matildae	AODF 0906	Queensland, Australia	Winton	Cenomanian	R mt I—V, five R pp (including one pu)	?-?-3-2-0	?-?-1-0- 0	This paper

(Continued.)

 Table 12. (Continued.)

taxon	specimen	locality	formation	age range	pedal elements present	phalangeal formula	ungual formula	reference(s)
Titanosauriformes indet. (='Acanthopholis' platypus)	CAMSM B55449— 55461	Cambridgeshire, United Kingdom	Upper Greensand	l. Cenomanian	L mt I–V and one pp	į	¿	Seeley [285–287]); Le Loeuff [288]
Sonorasaurus thompsoni	ASDM 500	Arizona, USA	Turney Ranch	u. Albian—I. Cenomanian	R mt I–II and IV–V, six pp (including two pu)	į	į	Ratkevich [289]; D'Emic <i>et al.</i> [246]
Gobititan shenzhouensis	IVPP 12579	Gansu, China	Xinminbao Group	I.—m. Albian	L mt I–V, 10 pp (including three pu)	2-2-2-2	1-1-1-0- 0	You <i>et al.</i> [290]
Ligabuesaurus leanzai	MCF-PVPH-233	Neuquén, Argentina	Lohan Cura	I. Albian	R mt I–V, R pp I-1–2 and II-1 (including one pu)	24:1-1:1	1-?-? ?-?-?	Bonaparte <i>et al.</i> [291]; Bellardini <i>et al.</i> [174]
Cedarosaurus weiskopfae (=Pleurocoelus sp.)	FMNH PR <i>977</i>	Texas, USA	Paluxy	I. Albian	L mt I–V, 11 L pp (including four pu)	2-3-4-2-0 [292] 2-3-3- 3-1 [293]	1-1-1-1 0	Langston (294]; Gallup [292]; D'Emic [293]
Мпуатаматtика тоуоwаткіа	RRBP 05834	Rukwa Rift Basin, Tanzania	Galula	Aptian— Cenomanian	L mt I–II and IV–V, R mt III, three pp (including two pu)	į	į.	Gorscak & O'Connor [254]
Sauroposeidon proteles (= Paluxysaurus jonesí)	FWMSH 938-10-16	Texas, USA	Twin Mountains	Aptian—Albian	R mt I and II	į	;	Rose [175]; D'Emic [293]
Sauroposeidon proteles (= Paluxysaurus jonesí)	FWMSH 93B-10-26	Texas, USA	Twin Mountains	Aptian–Albian	?L mt IV	į	<i>:</i>	Rose [175]; D'Emic [293]
Chubutisaurus insignis MPEF-PV 1129 Chubut, Argenti	MPEF-PV 1129	Chubut, Argentina	Cerro Barcino	u. Aptian—I. Albian	L mt IV	į	į	Carballido <i>et al.</i> [243]

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taxon	specimen	locality	formation	age range	pedal elements present	phalangeal formula	formula	reference(s)
Tapuiasaurus macedoi	MZSP-PV 807	Minas Gerais, Brazil	Quiricó	Aptian	L mt l–V, several pp	2-2-2-1-1	1-1-1-0-	Zaher <i>et al.</i> [21]; Navarro [295]
Malawisaurus dixeyi	MAL 145	Dinosaur Beds	Karonga, Malawi	Aptian	?mt III	į	į	Gomani [115]
Malawisaurus dixeyi	MAL 210–213	Dinosaur Beds	Karonga, Malawi	Aptian	R mt V, R pp l-1, L pu l-2 and ll-2	į	į	Gomani [115]
Tangvayosaurus hoffeti	TV2-1-40	Savannakhet, Laos	Grès supérieurs	Aptian	L mt I–III, nine pp (including three pu)	2-3-3-2-1	1-1-1-0-	Allain <i>et al.</i> [296]
Titanosauriformes indet. (=Astrodon johnstoni,	USNM various	Maryland, USA	Arundel	l. Aptian	six mt and 13 pp (only one positively	į	į	Marsh [297]; Lull [298]; Carpenter
=Preurocoerus artus, = P. nanus)					identined as pedal)			& HOWEII [123]
Tastavinsaurus sanzi	9/66 ZAW	Teruel, Spain	Хегт	l. Aptian	R mt I–V, L mt IV, seven pp (induding four pu)	2-3-3?-2?-1?	1-1-1-0- 0	Canudo <i>et al.</i> [220]; Royo-Torres [221]
Tastavinsaurus sanzi	CPT- various	Teruel, Spain	Forcall	l. Aptian	L mt I–V, 12 L pp (including three pu)	2-3-4-2-1	1-1-1-0- 0	Royo-Torres <i>et al.</i> [239]
Venenosaurus dicrocei	DMNH 40932	Utah, USA	Cedar Mountain (Poison Strip Member)	Barremian—I. Aptian	R mt I–II and IV	į	'	Tidwell <i>et al.</i> [299]; D'Emic <i>et al.</i> [246]
Phuwiangosaurus sirindhornae	SM K11-0001– 0168	Khon Kaen, Thailand	Sao Khua	Barremian	three ?mt, two phalanges (might be manual)	į	į	Suteethorn <i>et al.</i> Suteethorn <i>et al.</i> [149]
								(Continued.)

 Table 12. (Continued.)

taxon	specimen	locality	formation	age range	pedal elements present	phalangeal formula	ungual formula	reference(s)
Dongbeititan dongi	DNHM D2867	Liaoning, China	Yixian (Jianshangou Member)	Barremian	L mt I–V, pu I and II	¿	~.	Wang <i>et al.</i> [300]
Fukuititan nipponensis	FPDM-V8468	Fukui, Japan	Kitadani	Barremian	pp ?	į	ć	Azuma & Shibata [301]
Cedarosaurus weiskopfae	DMNH 39045	Utah, USA	Cedar Mountain (Yellow Cat Member)	Valanginian	right mt I–II and V and four R pp (including three pu)	. 1-1-1- 1-1-1-	1-1-1- ?-?	Tidwell <i>et al.</i> [255]; D'Emic <i>et al.</i> [246]
Euhelopus zdanskyi PMU 24706 Shandong, China	PMU 24706	Shandong, China	Mengyin	Berriasian	R mt I—IV, five R pp (including one definite and one possible pu)	? Wiman [111]; Wilson & Updurch [113]	į	Wiman [111]; Wilson & Upchurch [113]

- C553: postorbital, lateral surface hosts nutrient foramen near the junction of the squamosal and jugal processes: absent (0); present (1) (new character).
- C554: squamosal, quadrate facet is: undivided (0); divided such that it is bilobate or trilobate (1) (new character).
- C555: quadratojugal, posterior tongue-like process: absent or weakly developed (0); present (1).
- C556: quadratojugal and quadrate, horizontal ridge present across both elements anterior to their articulation point (lateral surface of quadrate, medial surface of quadratojugal: absent (0); present (1) (new character).

Several character scores were changed in existing OTUs. Below, C### denotes the character number, the first value in the parentheses (before the arrow) indicates the original score, and the second value (after the arrow) in the parentheses denotes the new score; sources external to this paper are cited where necessary.

Diamantinasaurus matildae AODF 0603: C24 (? \rightarrow 0); C108 (? \rightarrow 1; [43]); C109 (? \rightarrow 0&1; [43]); C110 (? \rightarrow 0; [43]); C111 (? \rightarrow 1; [43]); C113 (? \rightarrow 1; [43]); C173 (0 \rightarrow 1); C221 (? \rightarrow 0; [41]); C222 (? \rightarrow 0; [41]); C260 (0 \rightarrow 1); C321 (? \rightarrow 1; [43]); C347 (? \rightarrow 0); C402 (? \rightarrow 0; [43]); C408 (? \rightarrow 1); C454 (? \rightarrow 0; [43]); C513 (? \rightarrow 1; [44]); C546 (? \rightarrow 0; [43]).

Diamantinasaurus matildae AODF 0836 (including AODF 2298 [tooth]): C108 (? → 1; [43]); C173 (1 → ?); C321 (? → 1; [43]); C402 (? → 0; [43]); C454 (? → 0; [43]); C546 (? → 0; [43]).

Sarmientosaurus musacchioi: C76 (0 \rightarrow ?); C435 (1 \rightarrow 0).

Phuwiangosaurus sirindhornae: C315 (? \rightarrow 1; [149]); C316 (0 \rightarrow 1; [82]); C317 (1 \rightarrow 0; [82]); C550 (1 \rightarrow 0; [82]); C551 (? \rightarrow 1; [82]).

References

- Carballido JL, Otero A, Mannion PD, Salgado L, Pérez Moreno A. 2022 Titanosauria: a critical reappraisal of its systematics and the relevance of the South American record. In South American sauropodomorph dinosaurs: record, diversity and evolution (eds A Otero, JL Carballido, D Pol), pp. 269–298. Cham, Switzerland: Springer.
- Mannion PD, Carballido JL, Curry Rogers K, Díez Díaz V, Poropat SF. In press. Macronaria. In The Dinosauria (third edition) (eds PJ Makovicky, PM Barrett, MT Carrano, DB Weishampel).
 Cambridge, UK: Cambridge University Press.
- Curry Rogers KA. 2005 Titanosauria: a phylogenetic overview. In *The sauropods:* evolution and paleobiology (eds KA Curry Rogers, JA Wilson), pp. 50–103. Berkeley, CA: University of California Press.
- Wilson JA. 2006 An overview of titanosaur evolution and phylogeny. In Actas de las III jornadas internacionales sobre paleontología de dinosaurios y su entorno (ed. Colectivo Arqueológico-Paleontológico Salense), pp. 169—190. Salas de los Infantes, Burqos, España.
 - 5. D'Emic MD. 2012 Early evolution of titanosauriform sauropod dinosaurs. *Zool. J. Linnean Soc.* **166**, 624–671. (doi:10.
 1111/j.1096-3642.2012.00853.x)
- Mannion PD, Upchurch P, Schwarz D, Wings O. 2019 Taxonomic affinities of the putative titanosaurs from the Late Jurassic Tendaguru Formation of Tanzania: phylogenetic and biogeographic implications for eusauropod dinosaur evolution. *Zool. J. Linnean Soc.* 185, 784—909. (doi:10.1093/zoolinnean/zly068)
- Gallina PA, Canale JI, Carballido JL. 2021 The earliest known titanosaur sauropod dinosaur.

- *Ameghiniana* **58**, 35–51. (doi:10.5710/AMGH. 20.08.2020.3376)
- Upchurch P, Barrett PM, Dodson P. 2004
 Sauropoda. In *The Dinosauria (second edition)* (eds DB Weishampel, P Dodson, H Osmólska), pp. 259–322. Berkeley: University of California Press.
- Hocknull SA, White MA, Tischler TR, Cook AG, Calleja ND, Sloan T, Elliott DA. 2009 New mid-Cretaceous (latest Albian) dinosaurs from Winton, Queensland, Australia. PLoS ONE 4, e6190. (doi:10.1371/journal.pone.0006190)
- Cerda IA, Paulina Carabajal A, Salgado L, Coria RA, Reguero MA, Tambussi CP, Moly JJ. 2012 The first record of a sauropod dinosaur from Antarctica. Naturwissenschaften 99, 83–87. (doi:10.1007/s00114-011-0869-x)
- Gorscak E, O'Connor PM, Roberts EM, Stevens NJ. 2017 The second titanosaurian (Dinosauria: Sauropoda) from the middle Cretaceous Galula Formation, southwestern Tanzania, with remarks on African titanosaurian diversity. J. Vertebr. Paleontol. 37, e1343250. (doi:10. 1080/02724634.2017.1343250)
- Averianov A, Efimov V. 2018 The oldest titanosaurian sauropod of the Northern Hemisphere. *Biol. Commun.* 63, 145–162. (doi:10.21638/spbu03.2018.301)
- Díez Díaz V, Garcia G, Pereda Suberbiola X, Jentgen-Ceschino B, Stein K, Godefroit P, Valentin X. 2018 The titanosaurian dinosaur Atsinganosaurus velauciensis (Sauropoda) from the Upper Cretaceous of southern France: new material, phylogenetic affinities, and palaeobiogeographical implications. Cretaceous Res. 91, 429–456. (doi:10. 1016/j.cretres.2018.06.01)
- Gallina PA, González Riga BJ, Ortíz David LD.
 2022 Time for giants: titanosaurs from the

- Berriasian—Santonian age. In South American sauropodomorph dinosaurs: record, diversity and evolution (eds A Otero, JL Carballido, D Pol), pp. 299—340. Cham, Switzerland: Springer.
- Santucci RM, Filippi LS. 2022 Last titans: titanosaurs from the Campanian—Maastrichtian age. In South American sauropodomorph dinosaurs: record, diversity and evolution (eds A Otero, JL Carballido, D Pol), pp. 341–391. Cham, Switzerland: Springer.
- Curry Rogers K, Forster CA. 2001 The last of the dinosaur titans: a new sauropod from Madagascar. *Nature* 412, 530–534. (doi:10. 1038/35087566)
- Curry Rogers K, Forster CA. 2004 The skull of Rapetosaurus krausei (Sauropoda: Titanosauria) from the Late Cretaceous of Madagascar. J. Vertebr. Paleontol. 24, 121–144. (doi:10. 1671/A1109-10)
- Nowiński A. 1971 Nemegtosaurus mongoliensis

 n. gen., n. sp., (Sauropoda) from the uppermost

 Cretaceous of Mongolia. Palaeontol. Pol. 25, 57–81.
- Kurzanov SM, Bannikov AF. 1983 A new sauropod from the Upper Cretaceous of Mongolia. *Paleontol. J.* 1983, 90–96.
- Wilson JA. 2005 Redescription of the Mongolian sauropod Nemegtosaurus mongoliensis Nowinski (Dinosauria: Saurischia) and comments on Late Cretaceous sauropod diversity. J. Syst. Paleontol. 3, 283–318. (doi:10.1017/51477201905001628)
- Zaher H et al. 2011 A complete skull of an Early Cretaceous sauropod and the evolution of advanced titanosaurians. PLoS ONE 6, e16663. (doi:10.1371/journal.pone.0016663)
- Wilson JA, Pol D, Carvalho AB, Zaher H. 2016
 The skull of the titanosaur *Tapuiasaurus macedoi* (Dinosauria: Sauropoda), a basal

- titanosaur from the Lower Cretaceous of Brazil. Zool. J. Linnean Soc. **178**, 611–662. (doi:10. 1111/zoj.12420)
- Martínez RDF, Lamanna MC, Novas FE, Ridgely RC, Casal GA, Martínez JE, Vita JR, Witmer LM.
 2016 A basal lithostrotian titanosaur (Dinosauria: Sauropoda) with a complete skull: implications for the evolution and paleobiology of Titanosauria. PLoS ONE 11, e0151661. (doi:10.1371/journal.pone.0166272)
- D'Emic MD, Carrano MT. 2020 Redescription of brachiosaurid sauropod dinosaur material from the Upper Jurassic Morrison Formation, Colorado, USA. Anatom. Rec. 303, 732–758. (doi:10.1002/ ar.24198)
- Janensch W. 1935–1936 Die Schädel der Sauropoden Brachiosaurus, Barosaurus und Dicraeosaurus aus den Tendaguru-Schichten Deutsch-Ostafrikas. Palaeontographica, Supplement 38. VII 2, 145–298.
- Chure D, Britt BB, Whitlock JA, Wilson JA. 2010
 First complete sauropod dinosaur skull from the
 Cretaceous of the Americas and
 the evolution of sauropod dentition.
 Naturwissenschaften 97, 379–391. (doi:10.1007/s00114-010-0650-6)
- Longman HA. 1933 A new dinosaur from the Queensland Cretaceous. Memoirs Queensland Mus. 10. 131–144.
- Coombs Jr WP, Molnar RE. 1981 Sauropoda (Reptilia, Saurischia) from the Cretaceous of Queensland. Memoirs Queensland Mus. 20, 351– 373
- Molnar RE. 2001 A reassessment of the phylogenetic position of Cretaceous sauropod dinosaurs from Queensland, Australia. In VII linternational Symposium on Mesozoic Terrestrial Ecosystems: Asociacion Paleontológica Argentina publicación especial no. 7 (eds HA Leanza), pp. 139–144. Asociacion Paleontologica Argentina, Buenos Aires
- Molnar RE. 2010 Taphonomic observations on eastern Australian Cretaceous sauropods. Alcheringa 34, 421–429. (doi:10.1080/ 03115518.2010.497258)
- Molnar RE. 2011 New morphological information about Cretaceous sauropod dinosaurs from the Eromanga Basin, Queensland, Australia. Alcheringa 35, 329–339. (doi:10.1080/03115518.2011.533978)
- Molnar RE. 2011 Sauropod (Saurischia: Dinosauria) material from the Early Cretaceous Griman Creek Formation of the Surat Basin, Queensland. Australia. Alcheringa 35, 303–307. (doi:10.1080/03115518.2010.533975)
- Molnar RE, Salisbury SW. 2005 Observations on Cretaceous sauropods from Australia. In Thunderlizards: the sauropodomorph dinosaurs (eds V Tidwell, K Carpenter), pp. 454–465. Bloomington & Indianapolis, IA: Indiana University Press.
- Hocknull SA, Wilkinson M, Lawrence RA, Konstantinov V, Mackenzie S, Mackenzie R. 2021 A new giant sauropod, *Australotitan cooperensis* gen. et sp. nov., from the mid-Cretaceous of Australia. *PeerJ* 9, e11317. (doi:10.7717/ peeri.11317)
- Frauenfelder TG, Campione NE, Smith ET, Bell PR.
 2021 Diversity and palaeoecology of Australia's southern-most sauropods, Griman Creek Formation

- (Cenomanian), New South Wales, Australia. *Lethaia* **54**, 354–367. (doi:10.1111/let.1240**)**
- Poropat SF, Mannion PD, Upchurch P, Hocknull SA, Kear BP, Elliott DA. 2015 Reassessment of the non-titanosaurian somphospondylan Wintonotitan wattsi (Dinosauria: Sauropoda: Titanosauriformes) from the mid-Cretaceous Winton Formation, Queensland, Australia. Papers Palaeontol. 1, 59–106. (doi:10.1002/spp2.1004)
- Poropat SF, Upchurch P, Mannion PD, Hocknull SA, Kear BP, Sloan T, Sinapius GHK, Elliott DA. 2015 Revision of the sauropod dinosaur Diamantinasaurus matildae Hocknull et al. 2009 from the middle Cretaceous of Australia: implications for Gondwanan titanosauriform dispersal. Gondwana Res. 27, 995—1033. (doi:10.1016/j.gr.2014.03.014)
- Poropat SF et al. 2016 New Australian sauropods shed light on Cretaceous dinosaur palaeobiogeography. Sci. Rep. 6, 34467. (doi:10. 1038/srep34467)
- Poropat SF, Nair JP, Syme CE, Mannion PD, Upchurch P, Hocknull SA, Cook AG, Tischler TR, Holland T. 2017 Reappraisal of Austrosaurus mckillopi Longman, 1933 from the Allaru Mudstone of Queensland, Australia's first named Cretaceous sauropod dinosaur. Alcheringa 41, 543–580. (doi:10.1080/03115518.2017.1334826)
- Poropat SF, Mannion PD, Upchurch P, Tischler TR, Sloan T, Sinapius GHK, Elliott JA, Elliott DA. 2020 Osteology of the wide-hipped titanosaurian sauropod dinosaur Savannasaurus elliottorum from the Upper Cretaceous Winton Formation of Queensland, Australia. J. Vertebr. Paleontol. 40, e1786836. (doi:10.1080/ 02724634.2020.1786836)
- Poropat SF, Kundrát M, Mannion PD, Upchurch P, Tischler TR, Elliott DA. 2021 Second specimen of the Late Cretaceous sauropod dinosaur Diamantinasaurus matildae provides new anatomical information on skull and neck evolution in early titanosaurs and the biogeographic origins of Australian dinosaur faunas. Zool. J. Linnean Soc. 192, 610–674. (doi:10.1093/zoolinnean/zlaa173)
- Poropat SF, White MA, Ziegler T, Pentland AH, Rigby SL, Duncan RJ, Sloan T, Elliott DA. 2021 A diverse Late Cretaceous vertebrate tracksite from the Winton Formation of Queensland, Australia. PeerJ 9, e11544. (doi:10.7717/ peerj.11544)
- Poropat SF, Frauenfelder TG, Mannion PD, Rigby SL, Pentland AH, Sloan T, Elliott DA. 2022 Sauropod dinosaur teeth from the lower Upper Cretaceous Winton Formation of Queensland, Australia, and the global record of early titanosauriforms. R. Soc. Open Sci. 9, 220381. (doi:10.1098/rsos.220381)
- Rigby SL, Poropat SF, Mannion PD, Pentland AH, Sloan T, Rumbold SJ, Webster CB, Elliott DA. 2022 A juvenile *Diamantinasaurus matildae* (Dinosauria: Titanosauria) from the Upper Cretaceous Winton Formation of Queensland, Australia, with implications for sauropod ontogenetic growth. *J. Vertebr. Paleontol.* 41, e2047991. (doi:10.1080/ 02724634.2021.2047991)
- 45. Navarro BA *et al.* 2022 A new nanoid titanosaur (Dinosauria: Sauropoda) from the Upper

- Cretaceous of Brazil. *Ameghiniana* **59**, 317–354. (doi:10.5710/AMGH.25.08.2022.3477)
- Vine RR. 1964 Mackunda, Queensland. 1:250
 000 geological series sheet SF54-11. Canberra,
 Australia: Bureau of Mineral Resources, Geology and Geophysics.
- Vine RR, Casey DJ. 1967 Winton, Queensland. 1:250 000 geological series sheet SF54-12.
 Canberra, Australia: Bureau of Mineral Resources, Geology and Geophysics.
- Senior BR, Mond A, Harrison PL. 1978
 Geology of the Eromanga Basin. Bureau of Mineral Resources. Geol. Geophys. Bull. 167, 1–102
- Fletcher TL, Moss PT, Salisbury SW. 2018 The palaeoenvironment of the Upper Cretaceous (Cenomanian—Turonian) portion of the Winton Formation, Queensland, Australia. *PeerJ* 6, e5513. (doi:10.7717/peeri.5513)
- Cook AG. 2012 Cretaceous faunas and events, northern Eromanga Basin, Queensland. *Episodes* 35, 153–159. (doi:10.18814/epiiugs/2012/ v35i1/014)
- Cook AG, Bryan SE, Draper JJ. 2013
 Postorogenic Mesozoic basins and magmatism.
 In Geology of Queensland (ed. PA Jell), pp.
 515–575. Brisbane: Geological Survey of Queensland.
- Berrell RW, Alvarado-Ortega J, Yabumoto Y, Salisbury SW. 2014 The first record of the ichthyodectiform fish Cladocyclus from eastern Gondwana: a new species from the Lower Cretaceous of Queensland, Australia. Acta Palaeontol. Pol. 59, 903–920. (doi:10.4202/app. 2012.0019)
- Kemp A. 1991 Australian Mesozoic and Cainozoic lungfish. In Vertebrate palaeontology of Australasia (eds P Vickers-Rich, JM Monaghan, RF Baird, TH Rich), pp. 465–496. Melbourne: Pioneer Design Studio.
- Kemp A. 1997 Four species of Metaceratodus (Osteichthyes: Dipnoi, Family Ceratodontidae) from Australian Mesozoic and Cainozoic deposits. J. Vertebr. Paleontol. 17, 26–33. (doi:10.1080/02724634.1997.10010949)
- Kemp A, Berrell RW. 2013 Lungfish as environmental indicators. In Mesozoic fishes 5 – global diversity and evolution (eds G Arratia, H-P Schultze, MVH Wilson), pp. 499–508. Munich, Germany: Verlag Dr. Friedrich Pfeil.
- Scanlon JD, Hocknull SA. 2008 A dolichosaurid lizard from the latest Albian (mid-Cretaceous) Winton Formation, Queensland, Australia. In Proceedings of the Second Mosasaur Meeting. Fort Hays Studies, special issue 3 (ed. MJ Everhart), pp. 131–136. Hays, Kansas: Fort Hays State University.
- Kear BP. 2016 Cretaceous marine amniotes of Australia: perspectives on a decade of new research. Memoirs Mus. Victoria 74, 17–28. (doi:10.24199/j.mmv.2016.74.03)
- Salisbury S, Molnar RE, Frey E, Willis PMA. 2006
 The origin of modern crocodyliforms: new evidence from the Cretaceous of Australia.
 Proc. R. Soc. B 273, 2439–2448. (doi:10.1098/rspb.2006.3613)
- Syme CE, Salisbury SW. 2018 Taphonomy of *Isisfordia duncani* specimens from the Lower Cretaceous (upper Albian) portion of the Winton

- Formation, Isisford, central-west Queensland. R. Soc. Open Sci. 5, 171651. (doi:10.1098/rsos.17165)
- White MA et al. 2022 Abdominal contents reveal Cretaceous crocodyliforms ate dinosaurs. Gondwana Res. 106, 281–302. (doi:10.1016/j. gr 2022 01 016)
- Pentland AH, Poropat SF, Tischler TR, Sloan T, Elliott RA, Elliott HA, Elliott JA, Elliott DA. 2019 Ferrodraco lentoni gen. et sp. nov., a new ornithocheirid pterosaur from the Winton Formation (Cenomanian–lower Turonian) of Queensland, Australia. Sci. Rep. 9, 13454. (doi:10.1038/s41598-019-49789-4)
- Pentland AH et al. 2022 The osteology of Ferrodraco lentoni, an anhanguerid pterosaur from the mid-Cretaceous of Australia. J. Vertebr. Paleontol. 41, e2038182. (doi:10.1080/ 02724634.2021.2038182)
- Pentland AH, Poropat SF, White MA, Rigby SL, Vickers-Rich P, Rich TH, Elliott DA. 2022 New anhanguerian (Pterosauria: Pterodactyloidea) remains from the Early Cretaceous of Queensland, Australia. Alcheringa 46, 188–197. (doi:10.1080/03115518.2022.2065028)
- Leahey LG, Salisbury SW. 2013 First evidence of ankylosaurian dinosaurs (Ornithischia: Thyreophora) from the mid-Cretaceous (late Albian–Cenomanian) Winton Formation of Queensland, Australia. Alcheringa 37, 249–257. (doi:10.1080/03115518.2013.743703)
- Hocknull SA, Cook AG. 2008 Hypsilophodontid (Dinosauria, Ornithischia) from latest Albian, Winton Formation, central Queensland. Mem. Queensland Mus. 52, 212.
- Herne MC, Nair JP, Evans AR, Tait AM. 2019 New small-bodied omithopods (Dinosauria, Neomithischia) from the Early Cretaceous Wonthaggi Formation (Strzelecki Group) of the Australian-Antarctic rift system, with revision of *Qantassaurus* intrepidus Rich and Vickers-Rich, 1999. J. Paleontol. 93, 543–584. (doi:10.1017/jpa.2018.95)
- Salisbury SW, Herne MC, Lamanna MC, Nair JP, Syme C, Witmer LM. 2019 An exceptionally preserved small-bodied ornithopod dinosaur from the Lower Cretaceous (upper Albian) Winton Formation of Isisford, Central-Western Queensland, Australia, and the diversification of Gondwanan ornithopods. J. Vertebr. Paleontol. Program Abstr. 2019, 185.
- Thulborn RA, Wade M. 1979 Dinosaur stampede in the Cretaceous of Queensland. *Lethaia* 12, 275–279. (doi:10.1111/j.1502-3931.1979. th01008 x)
- Thulborn RA, Wade M. 1984 Dinosaur trackways in the Winton Formation (mid-Cretaceous) of Queensland. Memoirs Queensland Mus. 21, 413–517.
- Thulborn RA, Wade M. 1989 A footprint as a history of movement. In *Dinosaur tracks* and traces (eds DD Gillette, MG Lockley), pp. 51–56. Cambridge, UK: Cambridge University Press.
- Romilio A, Salisbury SW. 2011 A reassessment of large theropod dinosaur tracks from the mid-Cretaceous (late Albian–Cenomanian) Winton Formation of Lark Quarry, central-western Queensland, Australia: a case for mistaken identity. Cretaceous Res. 32, 135–142. (doi:10. 1016/j.cretres.2010.11.003)

- Romilio A, Salisbury SW. 2014 Large dinosaurian tracks from the Upper Cretaceous (Cenomanian— Turonian) portion of the Winton Formation, Lark Quarry, central-western Queensland, Australia: 3D photogrammetric analysis renders the 'stampede trigger' scenario unlikely. Cretaceous Res. 51, 186–207. (doi:10.1016/j.cretres.2014.06.003)
- Romilio A, Tucker RT, Salisbury SW. 2013
 Reevaluation of the Lark Quarry dinosaur
 Tracksite (late Albian—Cenomanian Winton
 Formation, central-western Queensland,
 Australia): no longer a stampede? J. Vertebr.
 Paleontol. 33, 102—120. (doi:10.1080/02724634.
 2012.694591)
- Thulborn RA. 2013 Lark Quarry revisited: a critique of methods used to identify a large dinosaurian track-maker in the Winton Formation (Albian–Cenomanian), western Queensland, Australia. Alcheringa 37, 312–330. (doi:10.1080/03115518.2013.748482)
- Thulborn T. 2017 Behaviour of dinosaurian track-makers in the Winton Formation (Cretaceous, Albian-Cenomanian) at Lark Quarry, western Queensland, Australia: running or swimming? *Ichnos* 24, 1–18. (doi:10.1080/ 10420940.2015.1129326)
- White MA, Bell PR, Cook AG, Poropat SF, Elliott DA. 2015 The dentary of Australovenator wintonensis (Theropoda, Megaraptoridae); implications for megaraptorid dentition. PeerJ 3, e1512. (doi:10.7717/peerj.1512)
- White MA, Cook AG, Hocknull SA, Sloan T, Sinapius GHK, Elliott DA. 2012 New forearm elements discovered of holotype specimen Australovenator wintonensis from Winton, Queensland, Australia. PLoS ONE 7, e39364. (doi:10.1371/journal.pone.0039364)
- White MA et al. 2013 New Australovenator hind limb elements pertaining to the holotype reveal the most complete neovenatorid leg. PLoS ONE 8, e68649. (doi:10.1371/journal.pone.0068649)
- White MA, Bell PR, Poropat SF, Pentland AH, Rigby SL, Cook AG, Sloan T, Elliott DA. 2020 New theropod remains and implications for megaraptorid diversity in the Winton Formation (lower Upper Cretaceous), Queensland, Australia. R. Soc. Open Sci. 7, 191462. (doi:10.1098/rsos. 191462)
- Lallensack JN, Romilio A, Falkingham PL. 2022 A machine learning approach for the discrimination of theropod and ornithischian dinosaur tracks. J. R. Soc. Interface 19, 20220588. (doi:10.1098/rsif.2022.0588)
- Gureyev TE, Nesterets Y, Ternovski D, Thompson D, Wilkins SW, Stevenson AW, Sakellariou A, Taylor JA. 2011 Toolbox for advanced X-ray image processing. In Proc. of SPIE Optical Engineering + Applications 8141:81410B.
 San Diego, CA, 21–24 August 2011 SPIE.
- Kaikaew S, Suteethorn V, Deesri U, Suteethorn S. 2023 The endocast of *Phuwiangosaurus* sirindhornae from the Lower Cretaceous of Thailand. Cretaceous Res. 144, 105434. (doi:10. 1016/j.cretres.2022.105434)
- Riggs ES. 1903 Brachiosaurus altithorax, the largest known dinosaur. Am. J. Sci. 15(Series 4), 299–306. (doi:10.2475/ajs.s4-15.88.299)
- Riggs ES. 1904 Structure and relationships of opisthocoelian dinosaurs. Part II, the

- Brachiosauridae. *Field Columbian Museum Geol. Ser.* **2**, 229–248.
- Taylor MP. 2009 A re-evaluation of *Brachiosaurus altithorax* Riggs 1903 (Dinosauria, Sauropoda) and its generic separation from *Giraffatitan brancai* (Janensch 1914). *J. Vertebr. Paleontol.* 29, 787–806. (doi:10.1671/039.029.0309)
- Maidment SCR, Muxworthy A. 2019 A chronostratigraphic framework for the Upper Jurassic Morrison Formation, western U.S.A. J. Sediment. Res. 89, 1017–1038. (doi:10.2110/ isr.2019.54)
- Goloboff PA, Farris JS, Nixon KC. 2008 TNT: a free program for phylogenetic analysis. *Cladistics* 24, 774–786. (doi:10.1111/j.1096-0031.2008. 00217 x)
- Goloboff PA, Catalano SA. 2016 TNT version 1.5, including a full implementation of phylogenetic morphometrics. *Cladistics* 32, 221–238. (doi:10. 1111/cla.12160)
- Goloboff P, Morales M. 2023 TNT version 1.6, with a graphical interface for MacOs and Linux, including new routines in parallel. *Cladistics* 39, 144–153. (doi:10.1111/cla.12524)
- Mannion PD, Upchurch P, Jin X, Zheng W. 2019
 New information on the Cretaceous sauropod dinosaurs of Zhejiang Province, China: impact on Laurasian titanosauriform phylogeny and biogeography. R. Soc. Open Sci. 6, 191057. (doi:10.1098/rsos.191057)
- Mannion PD, Tschopp E, Whitlock JA. 2021
 Anatomy and systematics of the diplodocoid *Amphicoelias altus* supports high sauropod dinosaur diversity in the Upper Jurassic Morrison Formation of the USA. *R. Soc. Open Sci.* 8, 210377. (doi:10.1098/rsos.210377)
- 92. Goloboff PA. 2014 Extended implied weighting. *Cladistics* **30**, 260–272. (doi:10.1111/cla.12047)
- Goloboff PA, Torres A, Arias JS. 2018 Weighted parsimony outperforms other methods of phylogenetic inference under models appropriate for morphology. Cladistics 34, 407–437. (doi:10.1111/cla.12205)
- Marsh OC. 1878 Principal characters of American Jurassic dinosaurs: Part I. Am. J. Sci. 16(series 3), 411–416. (doi:10.2475/ajs.s3-16.95.411)
- Wilson JA, Sereno PC. 1998 Early evolution and higher-level phylogeny of sauropod dinosaurs.
 Society of Vertebrate Paleontology Memoir 5.
 J. Vertebr. Paleontol. 18, 1–68. (doi:10.1080/ 02724634.1998.10011115)
- Salgado L, Coria RA, Calvo JO. 1997 Evolution of titanosaurid sauropods. I: phylogenetic analysis based on the postcranial evidence. *Ameghiniana* 34, 3–32.
- Salgado L, Coria RA, Calvo JO. 1997 Presencia del genero Aeolosaurus (Sauropoda, Titanosauridae) en la Formacion Los Alamitos, Cretácico Superior de la Provincia de Rio Negro, Argentina. Geociências 2, 44–49.
- Bryan SE, Cook AG, Allen CM, Siegel C, Purdy DJ, Greentree JS, Uysal IT. 2012 Early—mid Cretaceous tectonic evolution of eastern Gondwana: from silicic LIP magmatism to continental rupture. *Episodes* 35, 142–152. (doi:10.18814/epiiugs/2012/v35i1/013)
- Tucker RT, Roberts EM, Hu Y, Kemp AIS, Salisbury SW. 2013 Detrital zircon age constraints for the Winton Formation,

- Queensland: contextualizing Australia's Late Cretaceous dinosaur faunas. *Gondwana Res.* **24**, 767–779. (doi:10.1016/j.gr.2012.12.009)
- Upchurch P. 1998 The phylogenetic relationships of sauropod dinosaurs. *Zool. J. Linnean Soc.* 124, 43–103. (doi:10.1111/j.1096-3642.1998. tb00569.x)
- Mannion PD, Upchurch P, Barnes RN, Mateus O. 2013 Osteology of the Late Jurassic Portuguese sauropod dinosaur *Lusotitan atalaiensis* (Macronaria) and the evolutionary history of basal titanosauriforms. *Zool. J. Linnean Soc.* 168, 98–206. (doi:10.1111/zoj.12029)
- Juárez Valieri R, Ríos Díaz SD. 2014
 Reinterpretation of the rostral skull shape of Malawisaurus dixeyi (Haughton) (Sauropoda, Titanosauria). Ameghiniana 51, 15.
- 103. Gorscak, E. 2016. Descriptive and comparative morphology of African titanosaurian sauropods: new information on the evolution of Cretaceous African continental faunas: In College of Arts and Sciences, Vol. Ph.D., pp. 470. Ohio University.
- Wilson JA. 2002 Sauropod dinosaur phylogeny: critique and cladistic analysis. *Zool. J. Linnean Soc.* 136, 217–276. (doi:10.1046/j.1096-3642. 2002.00029.x)
- WhitlockJA.2011 Inferences of diplodocoid (Sauropoda: Dinosauria) feeding behavior from snout shape and microwear analyses. *PLoS ONE* 6, e18304. (doi:10.1371/journal.pone.0018304)
- Upchurch P. 1995 The evolutionary history of sauropod dinosaurs. *Phil. Trans. Biol. Sci.* 349, 365–390. (doi:10.1098/rstb.1995.0125)
- Ksepka DT, Norell MA. 2010 The illusory evidence for Asian Brachiosauridae: new material of *Erketu ellisoni* and a phylogenetic reappraisal of basal Titanosauriformes. *Am. Mus. Novitates* 3700, 1–27. (doi:10.1206/3700.2)
- Mannion PD, Upchurch P, Mateus O, Barnes RN, Jones MEH. 2012 New information on the anatomy and systematic position of *Dinheirosaurus lourinhanensis* (Sauropoda: Diplodocoidea) from the Late Jurassic of Portugal, with a review of European diplodocoids. *J. Syst. Paleontol.* 10, 521–551. (doi:10.1080/14772019.2011.595432)
- McIntosh JS. 1990 Sauropoda. In *The Dinosauria* (eds DB Weishampel, P Dodson, H Osmólska), pp. 345–401. Berkeley: University of California Press.
- Sereno PC, Wilson JA, Witmer LM, Whitlock JA, Maga A, Ide 0, Rowe TA. 2007 Structural extremes in a Cretaceous dinosaur. PLoS ONE 2, e1230. (doi:10.1371/journal.pone.0001230)
- 111. Wiman C. 1929 Die Kreide-Dinosaurier aus Shantung. *Palaeontol. Sin. (Series C)* **6**, 1–67.
- Mateer NJ, McIntosh JS. 1985 A new reconstruction of the skull of Euhelopus zdanskyi (Saurischia: Sauropoda). Bull. Geol. Inst. Upsala (new series) 11, 124–132.
- WilsonJA, Upchurch P. 2009 Redescription and reassessment of Euhelopus zdanskyi (Dinosauria: Sauropoda) from the Early Cretaceous of China. J. Syst. Paleontol. 7, 199–239. (doi:10.1017/ S1477201908002691)
- 114. Poropat SF, Kear BP. 2013 Photographic atlas and three-dimensional reconstruction of the holotype skull of Euhelopus zdanskyi with description of additional cranial elements. PLoS

- ONE **8**, e79932. (doi:10.1371/journal.pone. 0079932)
- Gomani EM. 2005 Sauropod dinosaurs from the Early Cretaceous of Malawi, Africa. *Palaeontol. Electron.* 8, 27A.
- Calvo JO, González Riga BJ, Porfiri JD. 2007 A new titanosaur sauropod from the Late Cretaceous of Neuquén, Patagonia, Argentina. Arquivos do Museu Nacional, Rio de Janeiro 65, 485–504.
- Barrett PM, Upchurch P. 1994 Feeding mechanisms of *Diplodocus*. In *Aspects of* sauropod paleobiology. Gaia, vol. 10 (eds MG Lockley, VF Santos, CA Meyer, AP Hunt), pp. 195–203. Lisbon, Portugal: Museu Nacional de História Natural.
- Madsen Jr JH, McIntosh JS, Berman DS. 1995
 Skull and atlas-axis complex of the Upper Jurassic sauropod Camarasaurus Cope (Reptilia: Saurischia). Bull. Carnegie Mus. Nat. Hist. 31, 1–115. (doi:10.5962/p.240778)
- Marpmann JS, Carballido JL, Sander PM, Knötschke N. 2015 Cranial anatomy of the Late Jurassic dwarf sauropod *Europasaurus holgeri* (Dinosauria, Camarasauromorpha): ontogenetic changes and size dimorphism. *J. Syst. Paleontol.* 13, 221–263. (doi:10.1080/14772019.2013.87507#
- Jacobs LL, Winkler DA, Downs WR, Gomani EM.
 1993 New material of an Early Cretaceous titanosaurid sauropod dinosaur from Malawi.
 Palaeontology 36, 523-534.
- Kundrát M, Coria RA, Manning TW, Snitting D, Chiappe LM, Nudds J, Ahlberg PE. 2020 Specialized craniofacial anatomy of a titanosaurian embryo from Argentina. *Curr. Biol.* 30, 1–7. (doi:10.1016/j.cub.2020.07.091)
- Lull RS. 1911 The Reptilia of the Arundel Formation. In Lower Cretaceous (ed. Maryland Geological Survey), pp. 173–178. Baltimore, MD: Maryland Geological Survey, Johns Hopkins Press
- 123. Carpenter K, Tidwell V. 2005 Reassessment of the Early Cretaceous sauropod Astrodon johnsoni Leidy 1865 (Titanosauriformes). In Thunderlizards: the sauropodomorph dinosaurs (eds V Tidwell, K Carpenter), pp. 78–114. Bloomington & Indianapolis, IA: Indiana University Press.
- 124. Sciutto JC, Martínez RD. 1994 Un nuevo yacimiento fosilifero de la Formacion Bajo Barreal (Cretacico Tardio) y su fauna de sauropodos. Naturalia Patagónica, Ciencias de la Tierra 2. 77–47
- Carballido JL, Salgado L, Pol D, Canudo JI, Garrido A. 2012 A new basal rebbachisaurid (Sauropoda, Diplodocoidea) from the Early Cretaceous of the Neuquén Basin; evolution and biogeography of the group. *Hist. Biol.* 24, 631– 654. (doi:10.1080/08912963.2012.672416)
- Edmund AG. 1957 On the special foramina in the jaws of many ornithischian dinosaurs.
 Paleontol. Contrib. R. Ontario Mus. 48, 1–14.
- 127. Jin L, Chen J, Zan S, Butler RJ, Godefroit P. 2010 Cranial anatomy of the small ornithischian dinosaur *Changchunsaurus parvus* from the Quantou Formation (Cretaceous: Aptian— Cenomanian) of Jilin Province, northeastern China. J. Vertebr. Paleontol. 30, 196–214. (doi:10.1080/02724630903412372)
 - 128. Chiappe LM, Salgado L, Coria RA. 2001 Embryonic skulls of titanosaur sauropod

- dinosaurs. *Science* **293**, 2444–2446. (doi:10. 1126/science.1063723)
- Salgado L, Coria RA, Chiappe LM. 2005
 Osteology of the sauropod embryos from the Upper Cretaceous of Patagonia. Acta Palaeontol. Pol. 50, 79–92.
- García RA, Salgado L, Coria RA, Chiappe LM.
 2010 Osteología embrionaria de saurópodos titanosaurios de Neuquén (Argentina): aspectos ontogenéticos y evolutivos. *Ameghiniana* 47, 409–430. (doi:10.5710/AMGH.v47i4.4)
 - Whitlock JA. 2011 A phylogenetic analysis of Diplodocoidea (Saurischia: Sauropoda).
 Zool. J. Linnean Soc. 161, 872–915. (doi:10. 1111/j.1096-3642.2010.00665.x)
- Gauthier J. 1986 Saurischian monophyly and the origin of birds. In *The origin of birds and the* evolution of flight. Memoirs of the California Academy of Sciences, vol. 8 (ed. K Padian), pp. 1–55. San Francisco, CA: California Academy of Sciences.
- Carpenter K, Tidwell V. 1998 Preliminary description of a *Brachiosaurus* skull from Felch Quarry 1, Garden Park, Colorado. *Modern Geol.* 23. 69–84.
- Filippi LS, García RA, Garrido AC. 2011 A new titanosaur sauropod dinosaur from the Upper Cretaceous of North Patagonia, Argentina. Acta Palaeontol. Pol. 56, 505–520. (doi:10.4202/app. 2010.0019)
- Gilmore CW. 1925 A nearly complete articulated skeleton of *Camarasaurus*, a saurischian dinosaur from Dinosaur National Monument. *Memoirs Carnegie Mus.* 10, 347–384. (doi:10. 5962/ p.217807)
- 136. White TE. 1958 The braincase of *Camarasaurus lentus* (Marsh). *J. Paleontol.* **32**, 477–494.
- McIntoshJS, Miles CA, Cloward KC, Parker JR. 1996
 A new nearly complete skeleton of Camarasaurus.
 Bull. Gunma Mus. Nat. Hist. 1,1–87.
- Chatterjee S, Zheng Z. 2005 Neuroanatomy and dentition of Camarasaurus lentus. In Thunderlizards: the sauropodomorph dinosaurs (eds V Tidwell, K Carpenter), pp. 199–211. Bloomington & Indianapolis, IA: Indiana University Press.
- Wiersma K, Sander PM. 2017 The dentition of a well-preserved specimen of *Camarasaurus* sp.: implications for function, tooth replacement, soft part reconstruction, and food intake. *PalZ* 91, 145–161. (doi:10.1007/s12542-016-0332-6)
- WoodruffDC, Foster JR. 2017 The first specimen of *Camarasaurus* (Dinosauria: Sauropoda) from Montana: the northernmost occurrence of the genus. *PLoS ONE* 12, e0177423. (doi:10.1371/ journal.pone.0177423)
- GallinaPA, Apesteguía S. 2011 Cranial anatomy and phylogenetic position of the titanosaurian sauropod Bonitasaura salgadoi. Acta Palaeontol. Pol. 56, 45— 60. (doi:10.4202/app.2010.0011)
- 142. Carballido JL, Pol D, Parra Ruge ML, Padilla Bernal S, Páramo-Fonseca ME, Etayo-Sema F. 2015 A new Early Cretaceous brachiosaurid (Dinosauria, Neosauropoda) from northwestem Gondwana (Villa de Leiva, Colombia). J. Vertebr. Paleontol. 35, e980505. (doi:10.1080/02724634.2015.98050)
- 143. Tschopp E, Mateus O. 2013 The skull and neck of a new flagellicaudatan sauropod from the Morrison Formation and its implication for the evolution and ontogeny of diplodocid dinosaurs.

- *J. Syst. Paleontol.* **11**, 853–888. (doi:10.1080/14772019.2012.746589)
- 144. Powell JE. 1992 Osteologia de Saltasaurus loricatus (Sauropoda—Titanosauridae) del Cretácico Superior del noroeste Argentino. In Los dinosaurios y su entorno biótico: Actas del Segundo Curso de Paleontologia in Cuenca (eds JL Sanz, AD Buscalioni), pp. 165–230. Cuenca, Spain: Instituto 'Juan de Valdés'.
- Powell JE. 2003 Revision of South American titanosaurid dinosaurs: palaeobiological, palaeobiogeographical and phylogenetic aspects. Rec. Queen Vic. Mus. 111, 1–173.
- 146. Martinelli AG, Forasiepi AM. 2004 Late Cretaceous vertebrates from Bajo de Santa Rosa (Allen Formation), Río Negro province, Argentina, with the description of a new sauropod dinosaur (Titanosauridae). Revista del Museo Argentino de Ciencias Naturales 6, 257–305. (doi:10.22179/REVMACN.6.88)
- 147. Suteethorn S, Le Loeuff J, Buffetaut E, Suteethorn V, Talumbook C, Chonglakmani C. 2009 A new skeleton of *Phuwiangosaurus* sirindhornae (Dinosauria, Sauropoda) from NE Thailand. In *Late Palaeozoic and Mesozoic* ecosystems in SE Asia. Geological Society Special Publication, vol. 315 (eds E Buffetaut, G Cuny, J Le Loeuff, V Suteethorn), pp. 189–215. London, UK: Geological Society.
- 148. Salgado L, Calvo JO. 1992 Cranial osteology of Amargasaurus cazaui Salgado & Bonaparte (Sauropoda, Dicraeosauridae) from the Neocomian of Patagonia. Ameghiniana 29, 337–346.
- 149. Amiot R, Buffetaut E, Lécuyer C, Fernandez V, Fourel F, Martineau F, Suteethorn V. 2009 Oxygen isotope composition of continental vertebrate apatites from Mesozoic formations of Thailand; environmental and ecological significance. In Late Palaeozoic and Mesozoic ecosystems in SE Asia. Geological Society Special Publications, vol. 315 (eds E Buffetaut, G Cuny, J Le Loeuff, V Suteethorn), pp. 271–283. London, UK: Geological Society.
- Curry Rogers K, Wilson JA. 2014 Vahiny depereti, gen. et sp. nov., a new titanosaur (Dinosauria, Sauropoda) from the Upper Cretaceous Maevarano Formation, Madagascar. J. Vertebr. Paleontol. 34, 606–617. (doi:10.1080/02724634. 2013.822874)
- Saegusa H, Ikeda T. 2014 A new titanosauriform sauropod (Dinosauria: Saurischia) from the Lower Cretaceous of Hyogo, Japan. *Zootaxa* 3848, 1–66. (doi:10.11646/zootaxa.3848.1.1)
- 152. Mannion PD. 2011 A reassessment of Mongolosaurus haplodon Gilmore, 1933, a titanosaurian sauropod dinosaur from the Early Cretaceous of Inner Mongolia, People's Republic of China. J. Syst. Paleontol. 9, 355–378. (doi:10. 1080/14772019.2010.527379)
- Paulina Carabajal A. 2012 Neuroanatomy of titanosaurid dinosaurs from the Upper Cretaceous of Patagonia, with comments on endocranial variability within Sauropoda. *Anat. Rec.* 295, 2141–2156. (doi:10.1002/ ar.22572)
- 154. Witmer LM, Ridgely RC, Dufeau DL, Semones MC. 2008 Using CT to peer into the past: 3D visualization of the brain and ear regions of birds, crocodiles, and nonavian dinosaurs. In Anatomical

- imaging: towards a new morphology (eds H Endo, R Frey), pp. 67–87. Tokyo, Japan: Springer.
- Salgado L, Gallina PA, Paulina Carabajal A. 2015
 Redescription of *Bonatitan reigi* (Sauropoda: Titanosauria), from the Campanian— Maastrichtian of the Río Negro Province (Argentina). *Hist. Biol.* 27, 525—548. (doi:10. 1080/08912963.2014.894038)
- Sander PM, Mateus O, Laven T, Knötschke N. 2006
 Bone histology indicates insular dwarfism in a new Late Jurassic sauropod dinosaur. *Nature* 441, 739–741. (doi:10.1038/nature04633)
- Mannion PD, Allain R, Moine O. 2017 The earliest known titanosauriform sauropod dinosaur and the evolution of Brachiosauridae. PeerJ 5, e3217. (doi:10.7717/peerj.3217)
- Remes K. 2009 Taxonomy of Late Jurassic diplodocid sauropods from Tendaguru (Tanzania). Fossil Rec. 12, 23–46. (doi:10.1002/ mmnq.200800008)
- Simón E, Salgado L, Calvo JO. 2018 A new titanosaur sauropod from the Upper Cretaceous of Patagonia, Neuquén Province, Argentina. Ameghiniana 55, 1–29. (doi:10.5710/AMGH.01. 08.2017.3051)
- Sallam HM et al. 2018 New Egyptian sauropod reveals Late Cretaceous dinosaur dispersal between Europe and Africa. Nat. Ecol. Evol. 2, 445–451. (doi:10.1038/s41559-017-0455-5)
- Tschopp E, Mateus O, Benson RBJ. 2015 A specimen-level phylogenetic analysis and taxonomic revision of Diplodocidae (Dinosauria, Sauropoda). PeerJ 3, e857. (doi:10.7717/peerj.857)
- Huene FV. 1929 Los Saurisquios y Ornitisquios del Cretáceo Argentina. Anales Museo de La Plata 3, 1–196.
- Apesteguía S. 2004 Bonitasaura salgadoi gen. et sp. nov.: a beaked sauropod from the Late Cretaceous of Patagonia. Naturwissenschaften 91, 493–497. (doi:10.1007/s00114-004-0560-6)
- 164. Machado EB, d L, Avilla S, Nava WR, d D, Campos A, Kellner AWA. 2013 A new titanosaur sauropod from the Late Cretaceous of Brazil. Zootaxa 3701, 301–321. (doi:10.11646/zootaxa. 3701.3.1)
- 165. Calvo JO, González Riga B. 2019 Baalsaurus mansillai gen. et sp. nov. a new titanosaurian sauropod (Late Cretaceous) from Neuquén, Patagonia, Argentina. Anais da Academia Brasileira de Ciências 91, e20180661. (doi:10. 1590/0001-3765201820180661)
- Whitlock JA, Harris JD. 2010 The dentary of Suuwassea emilieae (Sauropoda: Diplodocoidea). J. Vertebr. Paleontol. 30, 1637–1641. (doi:10. 1080/02724634.2010.501452)
- Osborn HF, Mook CC. 1921 Camarasaurus, Amphicoelias, and other sauropods of Cope. Memoirs Am. Mus. Nat. Hist. (New Series) 3, 247–387.
- 168. Le Loeuff J. 2005 Osteology of Ampelosaurus atacis (Titanosauria) from southern France. In Thunder-lizards: the sauropodomorph dinosaurs (eds V Tidwell, K Carpenter), pp. 115–137. Bloomington & Indianapolis, IA: Indiana University Press.
- Oelrich TM. 1956 The anatomy of the head of Ctenosaura pectinata (Iguanidae). Museum of Zoology, University of Michigan, Miscellaneous Publications 94, 1–122.

- Bell CJ, Mead JI, Swift SL. 2009 Cranial osteology of *Moloch horridus* (Reptilia: Squamata: Agamidae). *Rec. Western Aust. Mus.* 25, 201–237. (doi:10.18195/issn.0312-3162. 25(2).2009.201-237)
- 171. Calvo JO. 1994 Jaw mechanics in sauropod dinosaurs. In Aspects of sauropod paleobiology. Gaia, vol. 10 (eds MG Lockley, VF Santos, CA Meyer, AP Hunt), pp. 183–193. Lisbon, Portugal: Museu Nacional de História Natural.
- Barrett PM, Hasegawa Y, Manabe M, Isaji S, Matsuoka H. 2002 Sauropod dinosaurs from the Lower Cretaceous of eastern Asia: taxonomic and biogeographical implications. *Palaeontology* 45, 1197–1217. (doi:10.1111/1475-4983.00282)
- Li L-G, Li D-Q, You H-L, Dodson P. 2014 A new titanosaurian sauropod from the Hekou Group (Lower Cretaceous) of the Lanzhou-Minhe Basin, Gansu Province, China. PLoS ONE 9, e85979. (doi:10.1371/journal.pone.0085979)
- 174. Bellardini F, Coria RA, Pino DA, Windholz GJ, Baiano MA, Martinelli AG. 2022 Osteology and phylogenetic relationships of *Ligabuesaurus leanzai* (Dinosauria: Sauropoda) from the Early Cretaceous of the Neuquén Basin, Patagonia, Argentina. *Zool. J. Linnean Soc.* 196, 1333. (doi:10.1093/zoolinnean/zlac003)
- Rose PJ. 2007 A new titanosauriform sauropod (Dinosauria: Saurischia) from the Early Cretaceous of central Texas and its phylogenetic relationships. *Palaeontol. Electron.* 10, 8A.
- Díez Díaz V, Tortosa T, Le Loeuff J. 2013
 Sauropod diversity in the Late Cretaceous of southwestern Europe: the lessons of odontology. Annales de Paléontologie 99, 119–129. (doi:10.1016/j.annpal.2012.12.002)
- Hill RV, D'Emic MD, Bever GS, Norell MA. 2015 A complex hyobranchial apparatus in a Cretaceous dinosaur and the antiquity of avian paraglossalia. *Zool. J. Linnean Soc.* 175, 892– 909. (doi:10.1111/zoj.12293)
- Yoshida J, Kobayashi Y, Norell MA. 2023
 An ankylosaur larynx provides insights for birdlike vocalization in non-avian dinosaurs. Commun. Biol. 6, 152. (doi:10.1038/s42003-023-04513-x)
- Cabreira SF et al. 2016 A unique Late Triassic dinosauromorph assemblage reveals dinosaur ancestral anatomy and diet. Curr. Biol. 26, 3090–3095. (doi:10.1016/j.cub.2016.09.040)
- 180. Müller RT, Langer MC, Bronzati M, Pacheco CP, Cabreira SF, Dias-Da-Silva S. 2018 Early evolution of sauropodomorphs: anatomy and phylogenetic relationships of a remarkably wellpreserved dinosaur from the Upper Triassic of southern Brazil. *Zool. J. Linnean Soc.* 184, 1187–1248. (doi:10.1093/zoolinnean/zly009)
- Kermack D. 1984 New prosauropod material from South Wales. Zool. J. Linnean Soc. 82, 101–117. (doi:10.1111/j.1096-3642.1984. tb00538.x)
- 182. Galton PM, Kermack D. 2010 The anatomy of Pantydraco caducus, a very basal sauropodomorph dinosaur from the Rhaetian (Upper Triassic) of South Wales, UK. Revue de Paléobiologie 29, 341–404.
- Müller RT. 2020 Craniomandibular osteology of *Macrocollum itaquii* (Dinosauria: Sauropodomorpha) from the Late Triassic of

- southern Brazil. *J. Syst. Paleontol.* **18**, 805–841. (doi:10.1080/14772019.2019.1683902)
- 184. McPhee BW, Bittencourt JS, Langer MC, Apaldetti C, Da Rosa ÁAS. 2020 Reassessment of Unaysaurus tolentinoi (Dinosauria: Sauropodomorpha) from the Late Triassic (early Norian) of Brazil, with a consideration of the evidence for monophyly within non-sauropodan sauropodomorphs. J. Syst. Paleontol. 18, 259— 293. (doi:10.1080/14772019.2019. 1602856)
- 185. Beccari V, Mateus O, Wings O, Milàn J, Clemmensen LB. 2021 Issi saaneq gen. et sp. nov.—a new sauropodomorph dinosaur from the Late Triassic (Norian) of Jameson Land, Central East Greenland. Diversity 13, 561. (doi:10.3390/d13110561)
- Jaekel O. 1914 Über die Wirbeltierfunde in der oberen Trias von Halberstadt. Paläontologische Zeitschrift 1, 155–215. (doi:10.1007/ BF03160336)
- 187. Fürbringer M. 1922 Das Zungenbein der Wirbeltiere insbesondere der Reptilien ung Vögel. Nachgelassene Untersuchungen über systematische Phylogenie mi besonderer Berücksichtigung der Wurzel der Säugetiere. Abhandlungen der Heidelberger Akademie der Wissenschaften 11. 1–164
- Huene FV. 1928 Lebensbild des Saurischier-Vorkommens im obersten Keuper von Trossingen in Württemberg. *Palaeobiologica* 1, 103–116.
- Huene FV. 1932 Die fossile Reptil-Ordnung Saurischia, ihre Entwicklung und Geschichte. Monographien zur Geologie und Paläontologie (Series 1) 4, 1–361.
- Janensch W. 1932 Das Zungenbein der Dinosaurier. Sitzungsbericht der Gesellschaft Naturforschender Freunde zu Berlin 1932, 229–234.
- 191. Galton PM. 1985 Cranial anatomy of the prosauropod dinosaur Plateosaurus from the Knollenmergel (Middle Keuper, Upper Triassic) of Germany. II. All the cranial material and details of soft-part anatomy. Geologica et Palaeontol. 19, 119–159.
- Lallensack JN, Teschner EM, Pabst B, Sander PM.
 New skulls of the basal sauropodomorph Plateosaurus trossingensis from Frick,
 Switzerland: is there more than one species?
 Acta Palaeontol. Pol. 66, 1–28. (doi:10.4202/app.00804.2020)
- Carroll RL, Galton PM. 1977 'Modern' lizard from the Upper Triassic of China. Nature 266, 252–255. (doi:10.1038/266252a0)
- Evans SE, Milner AR. 1989 Fulengia, a supposed early lizard reinterpreted as a prosauropod dinosaur. Palaeontology 32, 223–230.
- Sekiya T. 2010 A new prosauropod dinosaur from Lower Jurassic in Lufeng of Yunnan. Global Geol. 29, 6–15.
- Galton PM. 1976 Prosauropod dinosaurs (Reptilia: Saurischia) of North America. *Postilla* 169, 1–98.
- Fabbri M, Navalón G, Mongiardino Koch N, Hanson M, Petermann H, Bhullar B-A. 2021 A shift in ontogenetic timing produced the unique sauropod skull. Evolution 75, 819–831. (doi:10. 1111/evo.14190)

- Martínez RN. 2009 Adeopapposaurus mognai, gen. et sp. nov. (Dinosauria: Sauropodomorpha), with comments on adaptations of basal Sauropodomorpha. J. Vertebr. Paleontol. 29, 142–164. (doi:10. 1671/039.029.0102)
- Apaldetti C, Martínez RN, Alcober OA, Pol D. 2011 A new basal sauropodomorph (Dinosauria: Saurischia) from Quebrada del Barro Formation (Marayes-El Carrizal Basin), northwestern Argentina. PLoS ONE 6, e26964. (doi:10.1371/journal.pone.0026964)
- Gow CE, Kitching JW, Raath MA. 1990 Skulls of the prosauropod dinosaur Massospondylus carinatus Owen in the collections of the Bernard Price Institute for Palaeontological Research. Palaeontol. Africana 27, 45–58.
- Sues H-D, Reisz RR, Hinic S, Raath MA. 2004
 On the skull of Massospondylus carinatus
 Owen 1854 (Dinosauria: Sauropodomorpha) from the Elliot and Clarens formations
 (Lower Jurassic) of South Africa. Annals Carnegie Mus. 73, 239–257. (doi:10.5962/p.316084)
- Zhang Y, Yang Z. 1995 A new complete osteology of Prosauropoda in Lufeng Basin, Yunnan, China, 100 pp. Kunming, China: Yunnan Publishing House of Science and Technology.
- Yates AM. 2007 The first complete skull of Melanorosaurus Haughton (Sauropodomorpha: Anchisauria). In Evolution and palaeobiology of early sauropodomorph dinosaurs. Special papers in palaeontology, vol. 77 (eds PM Barrett, DJ Batten), pp. 9–55.
- WangY-M, You H-L, Wang T. 2017 A new basal sauropodiform dinosaur from the Lower Jurassic of Yunnan Province, China. Sci. Rep. 7, 41881. (doi:10.1038/srep41881)
- Wang Y-M, Wang T, Yang Z-W, You H-L. 2020 Cranium and vertebral column of *Xingxiulong chengi* (Dinosauria: Sauropodomorpha) from the Early Jurassic of China. *Anat. Rec.* 303, 772–789. (doi:10.1002/ar.24305)
- Barrett PM, Upchurch P, Zhou X-D, Wang X-L.
 2007 The skull of *Yunnanosaurus huangi* Young,
 1942 (Dinosauria: Prosauropoda) from the
 Lower Lufeng Formation (Lower Jurassic) of
 Yunnan, China. *Zool. J. Linnean Soc.* 150,
 319–341. (doi:10.1111/j.1096-3642.2007.00290.
 x)
- Zhang Y. 1988 The Middle Jurassic dinosaurian fauna from Dashanpu, Zigong, Sichuan. Vol. III. Sauropod dinosaurs (1) Shunosaurus. 89 pp. Chengdu, Sichuan: Sichuan Publishing House of Science and Technology.
- Dong Z, Zhou S, Zhang Z. 1983 Dinosaurs from the Jurassic of Sichuan. *Palaeontol. Sin. (Series* () 162. 1–145.
- Zhang Y, Li K, Zeng Q. 1998 A new species of sauropod dinosaur from the Upper Jurassic of Sichuan Basin, China. J. Chengdu Univers. Technol. 25, 61–68.
- Woodruff DC, Carr TD, Storrs GW, Waskow K, Scannella JB, Nordén KK, Wilson JP. 2018 The smallest diplodocid skull reveals cranial ontogeny and growth-related dietary changes in the largest dinosaurs. Sci. Rep. 8, 14341. (doi:10.1038/s41598-018-32620-x)

- Tschopp E, Mateus O. 2017 Osteology of Galeamopus pabsti sp. nov. (Sauropoda: Diplodocidae), with implications for neurocentral closure timing, and the cervicodorsal transition in diplodocids. PeerJ 5, e3179. (doi:10.7717/peerj.3179)
- Marsh OC. 1883 Principal characters of American Jurassic dinosaurs: Part VI: restoration of Brontosaurus. Am. J. Sci. 26(series 3), 81–85. (doi:10.2475/ajs.s3-26.152.81)
- Canudo JI, Carballido JL, Garrido A, Salgado L.
 2018 A new rebbachisaurid sauropod from the Aptian—Albian, Lower Cretaceous Rayoso Formation, Neuquén, Argentina. Acta Palaeontol. Pol. 63, 679—691. (doi:10.4202/app. 00524.2018)
- 214. Laven TA. 2001 Kraniale osteologie eines sauropoden (reptilia, saurischia) aus dem oberjura norddeutschlands und dessen phylogenetische stellung: In institut für geowissenschaften, Vol. Diplomarbeit, pp. 171. Mainz, Germany: Johannes-Gutenberg-Universität Mainz.
- 215. Martin V, Suteethorn V, Buffetaut E. 1999 Description of the type and referred material of Phuwiangosaurus sirindhornae Martin, Buffetaut and Suteethorn, 1994, a sauropod from the Lower Cretaceous of Thailand. Oryctos 2, 39–91.
- 216. Romer AS. 1956 *Osteology of the reptiles*, 772 pp. Chicago, IL: University of Chicago Press.
- Wilson JA. 2011 Anatomical terminology for the sacrum of sauropod dinosaurs. *Contrib. Mus. Paleontol. Univ. Michigan* 32, 59–69.
- You H, Li D, Zhou L, Ji Q. 2006 Huanghetitan liujiaxiaensis, a new sauropod dinosaur from the Lower Cretaceous Hekou Group of Lanzhou Basin, Gansu Province. China. Geol. Rev. 52, 668–674.
- Averianov A, Ivantsov S, Skutschas P, Faingertz A, Leshchinskiy S. 2018 A new sauropod dinosaur from the Lower Cretaceous Ilek Formation, Western Siberia, Russia. Géobios 51, 1–14. (doi:10.1016/j.geobios.2017.12.004)
- Canudo JI, Royo-Torres R, Cuenca-Bescós G.
 2008 A new sauropod: *Tastavinsaurus sanzi* gen. et sp. nov. from the Early Cretaceous (Aptian) of Spain. *J. Vertebr. Paleontol.* 28, 712–731. (doi:10.1671/0272-4634(2008)28[712:ANSTSG]2. 0.CO:2)
- 221. Royo-Torres R. 2009 El saurópodo de Peñarroya de Tastavins. *Monografías Turolenses* **6**, 1–548.
- 222. Lü J, Azuma Y, Chen R, Zheng W, Jin X. 2008 A new titanosauriform sauropod from the early Late Cretaceous of Dongyang, Zhejiang Province. Acta Geol. Sin. 82, 225–235.
- 223. D'Emic MD, Mannion PD, Upchurch P, Benson RBJ, Pang Q, Cheng Z. 2013 Osteology of Huabeisaurus allocotus (Sauropoda: Titanosauriformes) from the Upper Cretaceous of China. PLoS ONE 8, e69375. (doi:10.1371/ journal.pone.0069375)
- 224. Lü J, Pu H, Xu L, Jia S, Zhang J, Shen C. 2014 Osteology of the giant sauropod dinosaur Ruyangosaurus giganteus Lü et al., 2009, 224 pp. Beijing, China: Geological Publishing House.
- Borsuk-Białynicka M. 1977 A new camarasaurid sauropod *Opisthocoelicaudia skarzynskii* gen. n., sp. n. from the Upper Cretaceous of Mongolia. *Palaeontol. Pol.* 37, 5–64.

- Salgado L, Apesteguía S, Heredia SE. 2005 A new specimen of *Neuquensaurus australis*, a Late Cretaceous saltasaurine titanosaur from north Patagonia. *J. Vertebr. Paleontol.* 25, 623–634. (doi:10.1671/0272-4634(2005)025[0623:ANSONA]2.0.CO;2)
- 227. D'Emic MD, Wilson JA. 2011 New remains attributable to the holotype of the sauropod dinosaur Neuquensaurus australis, with implications for saltasaurine systematics. Acta Palaeontol. Pol. 56, 61–73. (doi:10.4202/app. 2009.0149)
- Coria RA, Filippi LS, Chiappe LM, García R, Arcucci AB. 2013 Overosaurus paradasorum gen. et sp. nov., a new sauropod dinosaur (Titanosauria: Lithostrotia) from the Late Cretaceous of Neuquén, Patagonia, Argentina. Zootaxa 3683, 357–376. (doi:10.11646/zootaxa. 3683.4.2)
- Wedel MJ. 2003 The evolution of vertebral pneumaticity in sauropod dinosaurs. *J. Vertebr. Paleontol.* 23, 344–357. (doi:10.1671/0272-4634(2003)023[0344:TEOVPl]2.0.CO;2)
- Wedel MJ, Taylor MP. 2013 Caudal pneumaticity and pneumatic hiatuses in the sauropod dinosaurs Giraffatitan and Apatosaurus. PLoS ONE 8, e78213. (doi:10.1371/journal.pone. 0078213)
- Díez Díaz V, Demuth OE, Schwarz D, Mallison H.
 2020 The tail of the Late Jurassic sauropod Giraffatitan brancai: digital reconstruction of its epaxial and hypaxial musculature, and implications for tail biomechanics. Front. Earth Sci. 8, 160. (doi:10.3389/feart.2020.00160)
- Calvo JO, Salgado L. 1995 Rebbachisaurus tessonei sp. nov. a new Sauropoda from the Albian— Cenomanian of Argentina: new evidence on the origin of the Diplodocidae. Gaia 11, 13–33.
- 233. Powell JE. 1987 The Late Cretaceous fauna of Los Alamitos, Patagonia, Argentina. Part VI–The titanosaurids. Revista del Museo Argentino de Ciencias Naturales «Bernardino Rivadavia» e Instituto Nacional de Investigacion de las Ciencias Naturales 3, 147–153.
- Santucci RM, Arruda-Campos AC. 2011 A new sauropod (Macronaria, Titanosauria) from the Adamantina Formation, Bauru Group, Upper Cretaceous of Brazil and the phylogenetic relationships of Aeolosaurini. *Zootaxa* 3085, 1–33. (doi:10.11646/zootaxa.3085.1.1)
- Díez Díaz V, Mocho P, Páramo A, Escaso F, Marcos-Fernández F, Sanz JL, Ortega F. 2016 A new titanosaur (Dinosauria, Sauropoda) from the Upper Cretaceous of Lo Hueco (Cuenca, Spain). Cretaceous Res. 68, 49–60. (doi:10. 1016/ i.cretres.2016.08.001)
- González Riga BJ, Mannion PD, Poropat SF, Ortíz David LD, Coria JP. 2018 Osteology of the Late Cretaceous Argentinean sauropod dinosaur Mendozasaurus neguyelap: implications for basal titanosaur relationships. Zool. J. Linnean Soc. 184, 136–181. (doi:10.1093/zoolinnean/zlx103)
- Klinkhamer AJ, Mallison H, Poropat SF, Sinapius GHK, Wroe S. 2018 Three-dimensional musculoskeletal modelling of the sauropodomorph hind limb: the effect of postural change on muscle leverage. *Anatom. Rec.* 301, 2145–2163. (doi:10.1002/ar.23950)

- Carballido JL, Pol D, Otero A, Cerda IA, Salgado L, Garrido AC, Ramezzani J, Cúneo NR, Krause MJ. 2017 A new giant titanosaur sheds light on body mass evolution among sauropod dinosaurs. Proc. R. Soc. B 284, 20171219. (doi:10.1098/rspb.2017.1219)
- 239. Royo-Torres R, Alcalá L, Cobos A. 2012 A new specimen of the Cretaceous sauropod *Tastavinsaurus sanzi* from El Castellar (Teruel, Spain), and a phylogenetic analysis of the Laurasiformes. *Cretaceous Res.* 34, 61–83. (doi:10.1016/j.cretres.2011.10.005)
- Otero A. 2010 The appendicular skeleton of Neuquensaurus, a Late Cretaceous saltasaurine sauropod from Patagonia, Argentina. Acta Palaeontol. Pol. 55, 399–426. (doi:10.4202/app. 2009.0099)
- Cerda IA, Zurriaguz VL, Carballido JL, Gonzalez R, Salgado L. 2021 Osteology, paleohistology and phylogenetic relationships of Pellegrinisaurus powelli (Dinosauria: Sauropoda) from the Upper Cretaceous of Argentinean Patagonia. Cretaceous Res. 128, 104957. (doi:10. 1016/j.cretres.2021.104957)
- Wilson JA, Carrano MT. 1999 Titanosaurs and the origin of 'wide-gauge' trackways: a biomechanical and systematic perspective on sauropod locomotion. *Paleobiology* 25, 252– 267. (doi:10.1017/S0094837300026543)
- Carballido JL, Pol D, Cerda I, Salgado L. 2011
 The osteology of *Chubutisaurus insignis* Del
 Corro, 1975 (Dinosauria: Neosauropoda) from
 the 'Middle' Cretaceous of Central Patagonia,
 Argentina. *J. Vertebr. Paleontol.* 31, 93–110.
 (doi:10.1080/02724634.2011.539651)
- González Riga BJ, Lamanna MC, Ortiz David LD, Calvo JO, Coria JP. 2016 A gigantic new dinosaur from Argentina and the evolution of the sauropod hind foot. Sci. Rep. 6, 19165. (doi:10. 1038/srep19165)
- D'Emic MD, Wilson JA, Williamson TE. 2011 A sauropod dinosaur pes from the latest Cretaceous of North America and the validity of Alamosaurus sanjuanensis (Sauropoda, Titanosauria). J. Vertebr. Paleontol. 31, 1072–1079. (doi:10.1080/02724634.2011. 595856)
- 246. D'Emic MD, Foreman BZ, Jud NA. 2016 Anatomy, systematics, paleoenvironment, growth, and age of the sauropod dinosaur Sonorasaurus thompsoni from the Cretaceous of Arizona, USA. J. Paleontol. 90, 102–132. (doi:10.1017/jpa.2015.67)
- González Riga BJ, Calvo JO, Porfiri JD. 2008 An articulated titanosaur from Patagonia (Argentina): new evidence of neosauropod pedal evolution. *Palaeoworld* 17, 33–40. (doi:10.1016/j.palwor.2007.08.003)
- Nair JP, Salisbury SW. 2012 New anatomical information on *Rhoetosaurus brownei* Longman, 1926, a gravisaurian sauropodomorph dinosaur from the Middle Jurassic of Queensland, Australia. *J. Vertebr. Paleontol.* 32, 369–394. (doi:10.1080/02724634. 2012.622324)
- 249. González Riga BJ, Lamanna MC, Otero A, Ortíz David LD, Kellner AWA, Ibiricu LM. 2019 An overview of the appendicular skeletal anatomy of South American titanosaurian sauropods,

- with definition of a newly recognized clade. Anais da Academia Brasileira de Ciências **91**, e20180374.
- (doi:10.1590/0001-3765201920180374)
- Martínez R, Giménez O, Rodríguez J, Luna M, Lamanna MC. 2004 An articulated specimen of the basal titanosaurian (Dinosauria: Sauropoda) Epachthosaurus sciuttoi from the early Late Cretaceous Bajo Barreal Formation of Chubut Province, Argentina. J. Vertebr. Paleontol. 24, 107–120. (doi:10.1671/9.1)
- Curry Rogers K. 2009 The postcranial osteology of *Rapetosaurus krausei* (Sauropoda: Titanosauria) from the Late Cretaceous of Madagascar. *J. Vertebr. Paleontol.* 29, 1046– 1086. (doi:10.1671/039.029.0432)
- Lacovara KJ et al. 2014 A gigantic, exceptionally complete titanosaurian sauropod dinosaur from southern Patagonia, Argentina. Sci. Rep. 4, 6196. (doi:10.1038/srep06196)
- 253. González Riga BJ, Ortíz David LD, Tomaselli MB, Candeiro CRdA, Coria JP, Prámparo M. 2015 Sauropod and theropod dinosaur tracks from the Upper Cretaceous of Mendoza (Argentina): trackmakers and anatomical evidences. *J. South Am. Earth Sci.* 61, 134–141. (doi:10.1016/j. jsames.2014.11.006)
- 254. Gorscak E, O'Connor PM. 2019 A new African titanosaurian sauropod dinosaur from the middle Cretaceous Galula Formation (Mtuka Member), Rukwa Rift Basin, southwestern Tanzania. PLoS ONE 14, e0211412. (doi:10. 1371/journal.pone.0211412)
- Tidwell V, Carpenter K, Brooks W. 1999 New sauropod from the Lower Cretaceous of Utah, USA. Oryctos 2, 21–37.
- Csiki Z, Codrea V, Jipa-Murzea C, Godefroit P.
 2010 A partial titanosaur (Sauropoda, Dinosauria) skeleton from the Maastrichtian of Nălaţ-Vad, Haţeg Basin, Romania. Neues Jahrbuch für Geologie und Paläontologie, Abhandlungen 258, 297—324. (doi:10.1127/ 0077-7749/2010/0098)
- Gallina PA, Apesteguía S. 2015 Postcranial anatomy of *Bonitasaura salgadoi* (Sauropoda, Titanosauria) from the Late Cretaceous of Patagonia. *J. Vertebr. Paleontol.* 35, e924957. (doi:10.1080/02724634.2014.924957)
- 258. Royo-Torres R, Fuentes C, Meijide M, Meijide-Fuentes F, Meijide-Fuentes M. 2017 A new Brachiosauridae sauropod dinosaur from the lower Cretaceous of Europe (Soria Province, Spain). Cretaceous Res. 80, 38–55. (doi:10. 1016/j.cretres.2017.08.012)
- Mocho P, Pérez-García A, Martin Jiménez M, Ortega F. 2019 New remains from the Spanish Cenomanian shade light on the Gondwanan origin of European Early Cretaceous titanosaurs. Cretaceous Res. 95, 164—190. (doi:10.1016/j. cretres.2018.09.016)
- Mocho P, Royo-Torres R, Ortega F. 2019
 A new macronarian sauropod from the Upper Jurassic of Portugal. J. Vertebr. Paleontol. 39, e1578782. (doi:10.1080/02724634.2019. 1578782)
- Junior JC, Marinho TS, Martinelli AG, Langer MC.
 2019 Osteology and systematics of *Uberabatitan ribeiroi* (Dinosauria; Sauropoda): a Late Cretaceous titanosaur from Minas Gerais, Brazil.

- Zootaxa **4577**, 401–438. (doi:10.11646/zootaxa. 4577 3 1)
- 262. Mannion PD, Calvo JO. 2011 Anatomy of the basal titanosaur (Dinosauria, Sauropoda) Andesaurus delgadoi from the mid-Cretaceous (Albian-early Cenomanian) Río Limay Formation, Neuquén Province, Argentina: implications for titanosaur systematics. Zool. J. Linnean Soc. 163, 155–181.
- 263. Tidwell V, Stadtman K, Shaw A. 2005 Agerelated characteristics found in a partial pelvis of Camarasaurus. In Thunder-lizards: the sauropodomorph dinosaurs (eds V Tidwell, K Carpenter), pp. 180–186. Bloomington & Indianapolis, IA: Indiana University Press.
- Osborn HF. 1904 Manus, sacrum, and caudals of Sauropoda. Bull. Am. Mus. Nat. Hist. 20, 181– 190
- MookCC.1917 Criteria for the determination of species in the Sauropoda, with description of a new species of Apatosaurus. Bull. Am. Mus. Nat. Hist. 38. 355–360.
- 266. Taylor MP, Wedel MJ. 2012 Re-evaluating 'Apatosaurus' minimus, a bizarre Morrison Formation sauropod with diplodocoid and macronarian features: In 60th Annual Symposium of Vertebrate Palaeontology and Comparative Anatomy, 10th–15th September 2012 (eds M Friedman, G Lloyd), pp. 23, Oxford, UK: University of Oxford.
- Moore AJ, Upchurch P, Barrett PM, Clark JM, Xu X. 2020 Osteology of Klamelisaurus gobiensis (Dinosauria, Eusauropoda) and the evolutionary history of Middle-Late Jurassic Chinese sauropods. J. Syst. Paleontol. 20, 1299–1393. (doi:10.1080/14772019.2020.1759706)
- 268. Bonnan MF. 2005 Pes anatomy in sauropod dinosaurs: implications for functional morphology, evolution, and phylogeny. In *Thunder-lizards: the sauropodomorph dinosaurs* (eds V Tidwell, K Carpenter), pp. 346–380. Bloomington & Indianapolis, IA: Indiana University Press.
- Fowler DW, Hall LE. 2011 Scratch-digging sauropods, revisited. *Hist. Biol.* 23, 27–40. (doi:10.1080/08912963.2010.504852)
- Fronimos JA, Lehman TM. 2014 New specimens of a titanosaur sauropod from the Maastrichtian of Big Bend National Park, Texas. J. Vertebr. Paleontol. 34, 883–899. (doi:10.1080/02724634. 2014.840308)
- Huene FV, Matley CA. 1933 The Cretaceous Saurischia and Ornithischia of the central provinces of India. *Palaeontol. Indica* 21, 1–74.
- 272. Lehman TM, Coulson AB. 2002 A juvenile specimen of the sauropod dinosaur Alamosaurus sanjuanensis from the Upper Cretaceous of Big Bend National Park, Texas. J. Paleontol. 76, 156–172. (doi:10.1666/0022-3360(2002)076
 0156:AJSOTS>2.0.CO;2)
- Woodward HN, Lehman TM. 2009 Bone histology and microanatomy of Alamosaurus sanjuanensis (Sauropoda: Titanosauria) from the Maastrichtian of Big Bend National Park, Texas. J. Vertebr. Paleontol. 29, 807–821. (doi:10. 1671/039.029.0310)
- 274. García RA, Salgado L. 2013 The titanosaur sauropods from the late Campanian-early Maastrichtian Allen Formation of Salitral

- Moreno, Río Negro, Argentina. *Acta Palaeontol. Pol.* **58**, 269–284. (doi:10.4202/app.2011.0055)
- 275. Gallina PA, Otero A. 2015 Reassessment of Laplatasaurus araukanicus (Sauropoda: Titanosauria), from the Late Cretaceous of Patagonia, Argentina. Ameghiniana 52, 487— 501. (doi:10.5710/AMGH.08.06.2015.2911)
- 276. UllmannPV,Lacovara KJ. 2016 Appendicular osteology of *Dreadnoughtus schrani*, a giant titanosaurian (Sauropoda, Titanosauria) from the Upper Cretaceous of Patagonia, Argentina. J. Vertebr. Paleontol. 36, e1225303. (doi:10. 1080/02724634.2016.1225303)
- 277. Le Loeuff J. 1995 Ampelosaurus atacis (nov. gen., nov. sp.), un nouveau Titanosauridae (Dinosauria, Sauropoda) du Crétacé supérieur de la Haute Vallée de l'Aude (France). Comptes Rendus de l'Académie des Sciences Series IIA -Earth Planet. Sci. 321, 693–699.
- 278. Díez Díaz V, Pereda Suberbiola X, Sanz JL. 2013 Appendicular skeleton and dermal armour of the Late Cretaceous titanosaur *Lirainosaurus* astibiae (Dinosauria: Sauropoda) from Spain. Palaeontol. Electron. 16.2.19A, 1–18.
- 279. Ibiricu LM, Casal GA, Martínez RD, Luna M, González Svoboda E, Cerda IA. 2017 New materials of lithostrotian titanosaurs (Dinosauria: Sauropoda) from the Upper Cretaceous of central Patagonia. Cretaceous Res. 73, 25—39. (doi:10.1016/j.cretres.2016.12.015)
- 280. Garcia G, Amico S, Fournier F, Thouand E, Valentin X. 2010 A new titanosaur genus (Dinosauria, Sauropoda) from the Late Cretaceous of southern France and its paleobiogeographic implications. Bulletin de la Société Géologique de France 181, 269–277. (doi:10.2113/qssqfbull.181.3.269)
- Filippi LS, Salgado L, Garrido AC. 2019 A new giant basal titanosaur sauropod in the Upper Cretaceous (Coniacian) of the Neuquén Basin, Argentina. Cretaceous Res. 100, 61–81. (doi:10. 1016/j.cretres.2019.03.008)
- González Riga BJ. 2003 A new titanosaur (Dinosauria, Sauropoda) from the Upper Cretaceous of Mendoza Province, Argentina. Ameghiniana 40, 155–172.
- Soto M, Perea D, Cambiaso A. 2012 First sauropod (Dinosauria: Saurischia) remains from the Guichón Formation, Late Cretaceous of Uruguay. J. South Am. Earth Sci. 33, 68–79. (doi:10.1016/j.jsames.2011.08.003)
- Sues H-D, Averianov A, Ridgely RC, Witmer LM.
 Titanosauria (Dinosauria, Sauropoda) from the Upper Cretaceous (Turonian) Bissekty Formation of Uzbekistan. J. Vertebr. Paleontol.
 689145. (doi:10.1080/02724634.2014.
- 285. Seeley HG. 1869 Index to the fossil remains of Aves, Ornithosauria and Reptilia, from the Secondary System of strata arranged in the Woodwardian Museum of the University of Cambridge, 143 pp. Cambridge, UK: Cambridge University Press.
- Seeley HG. 1871 On Acanthopholis platypus (Seeley), a pachypod from the Cambridge Greensand. Ann. Mag. Nat. Hist. (Series 4) 8, 305–318. (doi:10.1080/00222937108696494)
- 287. Seeley HG. 1876 On *Macrurosaurus semnus* (Seeley), a long tailed animal with procedous

- vertebræ from the Cambridge Upper Greensand, preserved in the Woodwardian Museum of the University of Cambridge. *Q. J. Geol. Soc. Lond.* **32**, 440–444. (doi:10.1144/GSLJGS.1876.032. 01-04.50)
- 288. Le Loeuff J. 1993 European titanosaurids. *Revue de Paléobiologie* **7**, 105–117.
- Ratkevich R. 1998 New Cretaceous brachiosaurid dinosaur, Sonorasaurus thompsoni gen. et sp. nov., from Arizona. J. Arizona-Nevada Acad. Sci. 31, 71–82.
- You H, Tang F, Luo Z. 2003 A new basal titanosaur (Dinosauria: Sauropoda) from the Early Cretaceous of China. Acta Geol. Sin. 77, 424–429.
- Bonaparte JF, González Riga BJ, Apesteguía S.
 2006 Ligabuesaurus leanzai gen. et sp. nov.
 (Dinosauria, Sauropoda), a new titanosaur from the Lohan Cura Formation (Aptian, Lower Cretaceous) of Neuquén, Patagonia, Argentina. Cretaceous Res. 27, 364–376. (doi:10.1016/j. cretres.2005.07.004)
- Gallup MR. 1989 Functional morphology of the hindfoot of the Texas sauropod *Pleurocoelus* sp. indet. In *Paleobiology of the dinosaurs* (ed. JO Farlow), pp. 71–74. Boulder, CO: Geological Society of America.
- D'Emic MD. 2013 Revision of the sauropod dinosaurs of the Lower Cretaceous Trinity Group, southern USA, with the description of a new genus. J. Syst. Paleontol. 11, 707–726. (doi:10. 1080/14772019.2012.667446)
- 294. Langston Jr W. 1974 Non-mammalian Comanchean tetrapods. *Geosci. Man* **3**, 77–102.
- 295. Navarro BA. 2019 Osteologia pós-craniana e relações filogenéticas do titanossauro do Cretáceo Inferior *Tapuiasaurus macedoi* Zaher et al. 2011: In Departamento de Zoologia do Instituto de Biociências, Vol. Master of Science, Universidade de São Paulo, São Paulo.
- 296. Allain R et al. 1999 Un nouveau genre de dinosaure sauropode de la formation des Grès supérieurs (Aptien-Albien) du Laos. Comptes Rendus de l'Académie des Sciences - Series IIA -Earth Planet. Sci. 329, 609–616.
- Marsh OC. 1896 The dinosaurs of North America.
 133–416 pp. Washington, DC: United States Geological Survey.
- 298. Lull RS. 1911 Vertebrata (in Systematic paleontology of the Lower Cretaceous deposits of Maryland; eds RS Lull, WB Clark, EW Berry). In Lower Cretaceous (ed. Maryland Geological Survey), pp. 183–211. Baltimore, MD: Maryland Geological Survey, Johns Hopkins Press.
- 299. Tidwell V, Carpenter K, Meyer S. 2001 New titanosauriform (Sauropoda) from the Poison Strip Member of the Cedar Mountain Formation (Lower Cretaceous), Utah. In Mesozoic vertebrate life: New research inspired by the paleontology of Philip J. Currie (eds DH Tanke, K Carpenter), pp. 139–165. Bloomington & Indianapolis, IA: Indiana University Press.
- 300. Wang X, You H, Meng Q, Gao C, Chang X, Liu J. 2007 Dongbeititan dongi, the first sauropod dinosaur from the Lower Cretaceous Jehol Group of western Liaoning Province, China. Acta Geol. Sin. 81, 911–916. (doi:10.1111/j.1755-6724. 2007 tho1013 x)

- Azuma Y, Shibata M. 2010 Fukuititan nipponensis, a new titanosauriform sauropod from the Early Cretaceous Tetori Group of Fukui Prefecture, Japan. Acta Geol. Sin. 84, 454–462. (doi:10.1111/j.1755-6724.2010.00268.x)
- 302. Andrzejewski KA, Polcyn MJ, Winkler DA, Gomani Chindebvu E, Jacobs LL. 2019 The braincase of *Malawisaurus dixeyi* (Sauropoda: Titanosauria): a 3D reconstruction of the brain endocast and inner ear. PLoS ONE 14, e0211423
- Berman DS, McIntosh JS. 1978 Skull and relationships of the Upper Jurassic sauropod Apatosaurus (Reptilia, Saurischia). Bull. Carnegie Mus. Nat. Hist. 8, 1–35. (doi:10.5962/p.228587)
- Upchurch P. 1999 The phylogenetic relationships of the Nemegtosauridae (Saurischia, Sauropoda). J. Vertebr. Paleontol. 19, 106–125. (doi:10.1080/02724634.1999.10011127)
- Salgado L, Calvo JO. 1997 Evolution of titanosaurid sauropods. II: the cranial evidence. Ameahiniana 34, 33–48.
- 306. Currie PJ, Wilson JA, Fanti F, Mainbayar B, Tsogtbaatar K. 2018 Rediscovery of the type localities of the Late Cretaceous Mongolian sauropods Nemegtosaurus mongoliensis and Opisthocoelicaudia skarzynskii: stratigraphic and taxonomic implications. Palaeogeogr. Palaeoclimatol. Palaeoecol. 494, 5–13. (doi:10. 1016/j.palaeo.2017.10.035)
- McIntosh JS, Berman DS. 1975 Description of the palate and lower jaw of *Diplodocus* (Reptilia: Saurischia) with remarks on the nature of the skull of *Apatosaurus*. *J. Paleontol.* 49, 187–199.
- Sereno PC et al. 1999 Cretaceous sauropods from the Sahara and the uneven rate of skeletal evolution among dinosaurs. Science 286, 1342–1347. (doi:10.1126/science.286. 5443.1342)
- 309. Sereno PC, Wilson JA. 2005 Structure and evolution of a sauropod tooth battery. In *The* sauropods: evolution and paleobiology (eds KA Curry Rogers, JA Wilson), pp. 157–177. Berkeley, CA: University of California Press.
- You H, Ji Q, Lamanna MC, Li J, Li Y. 2004 A titanosaurian sauropod dinosaur with opisthocoelous caudal vertebrae from the early

- Late Cretaceous of Liaoning Province, China. *Acta Geol. Sin.* **78**, 907–911.
- Torcida Fernández-Baldor F, Canudo JI, Huerta P, Moreno-Azanza M, Montero D. 2017 Europatitan eastwoodi, a new sauropod from the lower Cretaceous of Iberia in the initial radiation of somphospondylans in Laurasia. PeerJ 5, e3409. (doi:10.7717/peerj.3409)
- Pang Q, Cheng Z. 2000 A new family of sauropod dinosaur from the Upper Cretaceous of Tianzhen, Shanxi Province, China. Acta Geol. Sin. 74. 117–125.
- Martinelli AG, Garrido AC, Forasiepi AM, Paz ER, Gurovich Y. 2007 Notes on fossil remains from the Early Cretaceous Lohan Cura Formation, Neuquén Province, Argentina. Gondwana Res. 11, 537–552. (doi:10.1016/j.gr. 2006 07 007)
- GilmoreCW.1933 Two new dinosaurian reptiles from Mongolia with notes on some fragmentary specimens. Am. Mus. Novitates 679, 1–20.
- Saegusa H, Tomida Y. 2011 Titanosauriform teeth from the Cretaceous of Japan. Anais da Academia Brasileira de Ciências 83, 247–265.
- Martin V, Buffetaut E, Suteethorn V. 1994 A new genus of sauropod dinosaur from the Sao Khua Formation (Late Jurassic or Early Cretaceous) of northeastern Thailand. Comptes Rendus des Séances de l'Academie des Sci. 319, 1085—1092.
- Suteethorn V, Martin V, Buffetaut E, Triamwi Chanon S, Chaimanee Y. 1995 A new dinosaur locality in the Lower Cretaceous of northeastern Thailand. Comptes Rendus de l'Académie des Sci. 321, 1041–1047.
- 318. Buffetaut E, Suteethorn V, Le Loeuff J, Cuny G, Tong H, Khansubha S. 2002 A review of the sauropod dinosaurs of Thailand. In *Proceedings* of the Symposium on Geology of Thailand, Bangkok, 26–31 August 2002, (ed. N Mantajit), pp. 95–101 Bangkok, Thailand.
- Buffetaut E, Suteethorn V. 2004 Comparative odontology of sauropod dinosaurs from Thailand. Revue de Paléobiologie 9, 151–159.
- Poropat SF. 2013 Carl Wiman's sauropods: the Uppsala Museum of Evolution's collection. *GFF* 135, 104–119. (doi:10.1080/11035897.2012. 759268)

- Zhou C-F, Wu W-H, Sekiya T, Dong Z-M. 2018 A new titanosauriformes dinosaur from Jehol Biota of western Liaoning, China. Global Geol. 37, 327–333.
- Van Hinsbergen DJJ, de Groot LV, Van Schaik SJ, Spakman W, Bijl PK, Sluijs A, Langereis CG, Brinkhuis H. 2015 A paleolatitude calculator for paleoclimate studies. PLoS ONE 10, e0126946.
- O'Brien CL et al. 2017 Cretaceous sea-surface temperature evolution: constraints from TEX₈₆ and planktonic foraminiferal oxygen isotopes. Earth Sci. Rev. 172, 224–247. (doi:10.1016/j. earscirev.2017.07.012)
- 324. Santamarina PE, Barreda VD, Iglesias A, Varela AN, Mays C. 2022 Comparison of mid-Cretaceous (Cenomanian—Turonian) high-latitude palynofloras from Patagonia and New Zealand: richness, ecology, and provincialization. Palaeogeogr. Palaeoclimatol. Palaeoecol. 604, 111216. (doi:10.1016/j.palaeo.2022.111216)
- 325. Herngreen GFW, Kedves M, Rovnina LV, Smirnova SB. 1996 Cretaceous palynofloral provinces: a review. In *Palynology: principles and applications, Vol. 3* (eds J Jansonius, DC McGregor), pp. 1157–1188. Houston, TX: American Association of Stratigraphic Palynologists Foundation.
- McLoughlin S, Pott C, Elliott D. 2010 The Winton Formation flora (Albian–Cenomanian, Eromanga Basin): implications for vascular plant diversification and decline in the Australian Cretaceous. Alcheringa 34, 303–323. (doi:10. 1080/03115511003669944)
- Alvarez BN, Casal GA, Ibiricu LM, De Sosa Tomas A, Suarez CA. 2022 Insights into paleoecology of the Bajo Barreal Formation (Upper Cretaceous), Patagonia, Argentina. J. South Am. Earth Sci. 116, 103854. (doi:10.1016/j.jsames.2022. 103854)
- 328. Poropat SF, Mannion PD, Rigby SL, Duncan RJ, Pentland AH, Bevitt JJ, Sloan T, Elliott DA. 2023 A nearly complete skull of the sauropod dinosaur *Diamantinasaurus matildae* from the Upper Cretaceous Winton Formation of Australia and implications for the early evolution of titanosaurs. Figshare. (doi:10.6084/m9.figshare. c.6533948)