A new neornithischian dinosaur from the Upper Jurassic Tiaojishan Formation of northern China (#112698)

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A new neornithischian dinosaur from the Upper Jurassic Tiaojishan Formation of northern China

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The Middle and Late Jurassic Yanliao Biota is different from other contemporaneous fossil assemblages in that it lacks neornithischian dinosaurs. Here, we report a new, earlydiverging neornithischian, *Pulaosaurus ginglong* gen.et sp. nov. , from the Upper Jurassic Tiaojishan Formation of Qinglong, Hebei Province, of northern China. Diagnostic or noteworthy morphological characteristics of *P. ginglong* include: five premaxillary teeth; a small boss is present on the caudoventral corner of the dorsal ramus of the jugal; a nuchal crest is located along the parietal; the manus has five digits; a supra-acetabular crest is present on the ilium; the paired arytenoids are gracile and leaf-like in form; the obturator process along the ischium is located near the pubic peduncle; a notch-like shaped obturator opening is present within the pubis; a robust fibular condyle forms a dorsoventrally extending crest on the tibia; a subtriangular flange on the anterior surface of the astragalus extends dorsolaterally along three distal tarsals; three of the distal tarsals are unfused, including a small drop-shaped distal tarsal 3; distal tarsal 3 is pierced by a foramen. A phylogenetic analysis places *P. ginglong* as one of the earliest-diverging neornithischians yet described. Moreover, P. ginglong represents the second known dinosaur to preserve ossified laryngeal elements, thus suggesting that a bird-like vocalization evolved early in non-avian dinosaur evolution.

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1 A new neornithischian dinosaur from the Upper 2 Jurassic Tiaojishan Formation of northern China 3 4 5 Yunfeng Yang^{1,2}, James Logan King^{1,3,4}, Xing Xu^{1,5} 6 ¹ Institute of Vertebrate Paleontology and Paleoanthropology, Chinese Academy of Sciences, 7 8 Beijing, 100044, China 9 ² College of Earth and Planetary Sciences, University of Chinese Academy of Sciences, Beijing, 10 100049, China ³ Colorado Northwestern Community College, Craig, Colorado, 81625, United States 11 ⁴ School of Earth Sciences, University of Bristol, Bristol, BS8 1RJ, United Kingdom 12 ⁵ Centre for Vertebrate Evolutionary Biology, Yunnan University, Kunming, Yunnan, 650091, 13 14 China 15 16 Corresponding Author: 17 Xing Xu 18 No.142 Xizhimen outer Street, Beijing, 100044, China Email address: Xing Xu, xuxing@ivpp.ac.cn 19 20 **Abstract** 21 The Middle and Late Jurassic Yanliao Biota is different from other contemporaneous fossil 22 23 assemblages in that it lacks neornithischian dinosaurs. Here, we report a new, early-diverging neornithischian, *Pulaosaurus qinglong* gen.et sp. nov., from the Upper Jurassic Tiaojishan 24 25 Formation of Qinglong, Hebei Province, of northern China. Diagnostic or noteworthy 26 morphological characteristics of P. qinglong include: five premaxillary teeth; a small boss is 27 present on the caudoventral corner of the dorsal ramus of the jugal; a nuchal crest is located 28 along the parietal; the manus has five digits; a supra-acetabular crest is present on the ilium; the 29 paired arytenoids are gracile and leaf-like in form; the obturator process along the ischium is 30 located near the pubic peduncle; a notch-like shaped obturator opening is present within the 31 pubis; a robust fibular condyle forms a dorsoventrally extending crest on the tibia; a



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subtriangular flange on the anterior surface of the astragalus extends dorsolaterally along three distal tarsals; three of the distal tarsals are unfused, including a small drop-shaped distal tarsal 3; distal tarsal 3 is pierced by a foramen. A phylogenetic analysis places *P. qinglong* as one of the earliest-diverging neornithischians yet described. Moreover, *P. qinglong* represents the second known dinosaur to preserve ossified laryngeal elements, thus suggesting that a bird-like vocalization evolved early in non-avian dinosaur evolution.

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Introduction

The Middle-to-Late Jurassic-aged Yanliao Biota is one of the most significant Mesozoic, 40 41 terrestrial lagerstätte in China, with an age that ranges from 168 Ma to 157 Ma (Bai 2024; 42 Huang 2015; Liu 2022; Zhou & Wang 2017) and is comprised of fossil assemblages from the 43 Jiulongshan and the Tiaojishan Formations (Boyd 2015; Huang 2015; Zhou & Wang 2017). The 44 Daohugou Biota whose age was about 168-164 Ma represents the early stage of the Yanliao 45 Biota while the Linglongta Biota whose age was about 162-157 Ma represents its late stage 46 (Huang 2015; Zhou & Wang 2017). In total, there have been 54 genera and 58 species of 47 vertebrates reported from the Yanliao Biota, including 9 species of non-avian dinosaurs (Liu 48 2022; Zhou & Wang 2017). Because the Yanliao Biota preserves large amounts of vertebrate 49 material from many taxa, it offers insight to major palaeobiological events, such as the temporal 50 origin of birds and the early evolution of mammals (Liu 2022; Xu et al. 2022; Zhou & Wang 51 2017). However, non-avian dinosaurs found in the Yanliao Biota are all small-bodied theropods whereas Ornithischia is represented by only one species, which may possibly be from the Jehol 52 53 Biota (Zhou & Wang 2017). This is in stark contrast to other contemporaneous Chinese terrestrial faunas such as the Shishugou and Shaximiao faunas where body size and taxonomic 54 55 composition are more variable (Liu 2022; Xu et al. 2022).



56	Neornithischia is a significant group of dinosaurs whose earliest origin could be dated back to
57	the Middle Jurassic, represented by several early-diverging taxa including Sanxiasaurus,
58	Agilisaurus, Hexinlusaurus found in China (Fonseca et al. 2024; Li et al. 2019; Barrett and Han
59	et al. 2005). Besides China, there have been reports of neornithischian fossils from the Middle
60	Jurassic of Russia (Cincotta et al.), Scotland (Panciroli et al. 2025) and strata from other
61	geological times and countries. Neornithischia diverges rapidly into a number of taxa in
62	Cretaceous (Fonseca et al. 2024). The phylogeny of Neornithischia has been controversial in
63	recent years, especially for the early-diverging taxa referred as 'hypsilophodontids' (Boyd 2015;
64	Han et al. 2018; Dieudonné et al. 2020; Brown et al. 2022). There are two major hypotheses for
65	the phylogeny of Neornithischia: 1. most of the early-diverging neornithischians are grouped
66	within Cerapoda, as a paraphyletic group of early-diverging ornithopods (Brown et al. 2022); 2.
67	most of the early-diverging neornithischians are outside Cerapoda and within the clade
68	Thescelosauridae (Brown et al. 2022). Boyd (2015) and Han et al. (2018) agree on the former
69	hypothesis while Dieudonné et al. (2020) and Fonseca et al. (2024) agree on the latter
70	hypothesis; however, the controversy over the phylogeny of Neornithischia stems from the
71	differences in the morphological characters selected in the different phylogenetic analyses
72	(Brown et al. 2022), so the discovery of a new early-diverging neornithischian species helps to
73	clarify relationships among early-diverging neornithischian taxa.
74	Here, we describe a new specimen found in the Upper Jurassic Tiaojishan Formation, County
75	Qinglong, Province Hebei, China. This well-preserved specimen IVPP V30936 has both basal
76	and derived characteristics of Ornithischia and preserves cololites, hyoids, and an ossified
77	larynx. This new neornithischian specimen helps us understand the biodiversity of Yanliao Biota
78	and the phylogeny of early-diverging Neornithischia better.

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80	Institutional abbreviations
81	IVPP: Institute of Vertebrate Paleontology and Paleoanthropology, Beijing, China
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83 84	Materials & Methods The specimen V30936 was collected by Mr. Yong Wang and later transferred to the IVPP. It
85	is currently stored in IVPP. It was found in southern Shimen Gou, County Qinglong, Province
86	Hebei, People's Republic of China. The specimen is fully prepared and based on close
87	examinations and comparations of the osteology and matrix, the specimen was deemed authentic
88	The anatomical information of IVPP V30936 was acquired by personal observations. They
89	were supplemented by computed tomography (CT) scans of the pelvic girdle. The CT scans were
90	conducted by 160-Micro-CL(Computed Laminography)in IVPP. The CT data provides
91	insight into areas of the pelvic girdle obscured by the femora and the chevrons to gain more
92	complete anatomical data on IVPP 30936.
93	The electronic version of this article in Portable Document Format (PDF) will represent a
94	published work according to the International Commission on Zoological Nomenclature (ICZN),
95	and hence the new names contained in the electronic version are effectively published under that
96	Code from the electronic edition alone. This published work and the nomenclatural acts it
97	contains have been registered in ZooBank, the online registration system for the ICZN. The
98	ZooBank LSIDs (Life Science Identifiers) can be resolved and the associated information viewed
99	through any standard web browser by appending the LSID to the prefix http://zoobank.org/ . The
100	LSID for this publication is: urn:lsid:zoobank.org:pub:D3939AEC-9C5B-4397-9BA4-
101	47CB3F9DFEC8. The online version of this work is archived and available from the following

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103 104	Phylogenetic Analysis
105	To assess the phylogenetic position of IVPP V30936 within ornithischians, we ran the
106	phylogenetic analysis on the emended matrix dataset of Han et al. (2018), including IVPP
107	V30936 and Sanxiasaurus modaoxiensis, a recently reported early-diverging neornithischian
108	taxon. The coding of Sanxiasaurus was based on the description and images provided by Li et al.
109	(2019). For consistency, we followed the phylogenetic nomenclature of higher-level ornithischian
110	taxa as defined by Madzia et al. (2021).
111	The final dataset consists of 380 characters scored for 70 ingroup taxa and 4 outgroup taxa.
112	The analysis was conducted in TNT with each character equally weighted. The analysis is based
113	on a traditional search with 1000 random seeds and 1000 replications. The swapping algorithm is
114	tree bisection reconnection (TBR), with 100 trees to save per replication. The maximal memory
115	of trees was set to 100000 and zero-length branches collapsed.
116	Unstable taxa were identified by the command 'pruned trees' and include four taxa, such as
117	Pisanosaurus, Micropachycephalosaurus, Zephylosaurus, Yueosaurus. Having removed the
118	unstable taxa and repeating the steps described above, the reduced strict consensus tree was
119	gained and phylogenetic relationships of the remaining taxa were display.
120	Additionally, we carry out another analysis in TNT based on the emended dataset of Fonseca
121	et al. (2024) which consists of 943 characters and 173 taxa. Each character is equally weighted.
122	The analysis is based on a traditional search with 1000 random seeds and 1000 replications. The
123	swapping algorithm is tree bisection reconnection (TBR), with 100 trees to save per replication.
124	The maximal memory of trees was set to 100000 and zero-length branches collapsed.
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27	Results
28	Systematics Paleontology
29	Dinosauria Owen, 1842
30	Ornithischia Seeley, 1887
31	Neornithischia Cooper, 1985
32	Pulaosaurus qinglong gen.et sp. nov.
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34	Etymology
35	The generic name is derived from Chinese Pinyin for "Pulao", a mythical creature
36	resembling the Chinese dragon. According to Chinese legends, the "Pulao" engages in loud
37	shouting, thus referencing the possible bird-like vocalizations of this species. The specific name
38	is derived from the Chinese Pinyin for "Qinglong", which is the name of the county in Province
39	Hebei, China, where the specimen was found.
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41	Holotype
42	IVPP V30936 is comprised of a nearly complete skeleton prepared on a brownish-red
43	sandstone slab. The specimen preserves most of the skull and a complete postcranial skeleton.
44	Many skull elements are displaced and disarticulated due to mediolateral crushing. All cervical
45	vertebrae and dorsal vertebrae are compressed and obscured. The sacrum is overlapped by the
46	ilium. Only two proximal, five middle and 17 distal caudal vertebrae are well-preserved. Both
47	scapulae and the sternum are broken with only fragments being preserved. The left forelimb is
48	complete and articulated; however, the right humerus is not preserved and elements of the right
49	manus are disarticulated. Elements of the pelvic girdle are preserved although many
50	morphological characters are unknown due to overlapping and displacement of the available





elements. Most of the hindlimbs are preserved except for the right distal tarsals 2, 3, 4; left calcaneum; and phalange 1 of the left pes. Additionally, a pair of ceratobranchials and ossified arytenoids are also preserved ventrally adjacent to the mandible. Cololites (gut contents) are preserved in the abdominal cavity.

- Locality and Horizon
- 157 Southern Shimen Gou, County Qinglong, Province Hebei, People's Republic of China.
- 158 Tiaojishan Formation, Callovian-Oxfordian (Middle-Upper Jurassic)

160 Diagnosis

A small-bodied neornthischian dinosaur characterized by the combination of the following characteristics (autapomorphies preceded by an asterisk): five premaxillary teeth; a small boss is located on the caudoventral corner of the dorsal ramus of the jugal; a nuchal crest that is located on the parietal the mani have five digits; a supra-acetabular crest is located on the ilium; a pair of gracile, leaf-like arytenoids are present; the obturator process is near the pubic peduncle; the opening of the obturator on the pubis is notch-shaped; *a robust fibular condyle forms a dorsoventrally extending crest on the tibia; a subtriangular flange extends dorsolaterally on four distal tarsals; *three distal tarsals are unfused with distal tarsal 3 that is drop-shaped; distal tarsal 3 is pierced by a foramen.

Because our phylogenetic analysis recovered *Pulaosaurus qinglong* in a systematic position near that of the contemporaneous *Agilisaurus louderbacki* (Peng 1992) and *Hexinlusaurus multidens* (Barrett and Han et al. 2005), from Sichuan Province, southwestern China, we list additional taxonomical differences between *Pulaosaurus* and *Agilisaurus* and *Hexinlusaurus*.





Pulaosaurus qinglong differs from Agilisaurus louderbacki and Hexinlusaurus multidens in the
following characteristics: the relative length of the <i>Pulaosaurus</i> skull is more elongated while the
skull of Agilisaurus is foreshortened; all of the premaxillary teeth in Pulaosaurus are subequal to
each other in size while the middle premaxillary teeth are largest in Agilisaurus; a small boss is
present on the jugal of <i>Pulaosaurus</i> while there is no ornamentation on the jugal of <i>Agilisaurus</i>
or <i>Hexinlusaurus</i> ; the ventral edge of the jugal is straight in <i>Pulaosaurus</i> but caudoventrally
deflects in Agilisaurus and Hexinlusaurus; the orbital margin of the postorbital projects into the
orbit in Hexinlusaurus while it does not in Pulaosaurus and Agilisaurus; the frontals of
Pulaosaurus are shorter and more elongated than those of Hexinlusaurus and Agilisaurus; the
ventral process of the predentary is notably reduced in <i>Pulaosaurus</i> when compared to that of
Agilisaurus; the ossified tendons of Pulaosaurus are in basket-like arrangement of fusiform
tendons in caudal region while they are arranged longitudinally in Agilisaurus and
Hexinlusaurus; the ungual of manual digit I is sub-conical in Pulaosaurus while it is claw-like in
form in Agilisaurus and Hexinlusaurus a supra-acetabular crest is present on the ilium in
Pulaosaurus and Agilisaurus while it is absent in Hexinlusaurus; the obturator foramen in the
pubis is a notch-like shape in Pulaosaurus while it is a true foramen in Agilisaurus and
Hexinlusaurus; the ischial obturator process is near the pubic peduncle in Pulaosaurus while it is
located more distally from the pubic peduncle in Agilisaurus and Hexinlusaurus; the ratio
between the humerus and the femur in <i>Pulaosaurus</i> is shorter than that of <i>Hexinlusaurus and</i>
Agilisaurus; the astragalar ascending process is triangular in Pulaosaurus while it is finger-like
in Hexinlusaurus; there are three distal tarsals in Pulaosaurus while there are only two in
Agilisaurus and Hexinlusaurus.



Description and comparisons

The specimen is preserved on a brownish-red fine-grained sandstone slab (FIGURE 1) and is roughly mediolaterally compressed. The total length of this specimen is approximately 722mm (i.e. the length from the rostral end of the skull to the caudal end of the last preserved caudal vertebrate) while its skull length is about 8cm from the rostral end of the premaxilla to the caudal border of the parietal and the length of trunk (i.e. the length from the atlas to the caudal border of the pelvic girdle) is about 30cm. The neurocentral sutures of cervical and caudal vertebrae are not obliterated, which is a possible but not confirmed sign of an immature individual (Bertozzo et al. 2017). The ratio of the orbit diameter and the skull length is 37%, which is close to the ratio in immature specimens of *Jeholosaurus*, ranging from 40% to 50% in immature specimens and less than 33% in adult specimens (Barrett and Han, 2009). Because of this, we interpret IVPP V30936 to be an immature individual.

Skull

In lateral view, the skull shape (FIGURE 2, 3, Table 1) is elongate, low, and forms an obtuse triangle. Due to the displacement of the jugal, the exact position of postorbital bones and the shape of infratemporal fenestra are unknown. The supratemporal fenestra is sub-square in lateral view. The orbit is elliptical and is the largest cranial opening with a maximal diameter of which is about 3cm. Rostral to the orbit sits the subtriangular antorbital fossa observable in most early-diverging ornithischians including *Jeholosaurus*, *Lesothosaurus*, *Hexinlusaurus*, *Agilisaurus* (Norman et al. 2004; Barrett and Han 2009; Peng 1992; Sereno 1991; He et al. 1984), whose ventral margin is nearly at the same horizon of orbit ventral margin. All of the





mandibular elements are disarticulated from each other, so the overall shape of the mandible and the presence or shape of external mandibular fenestra is unknown.

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Premaxilla (FIGURE 2)—Only the left premaxilla is visible. In lateral view, the main body of the premaxilla is subrectangular and possesses two processes: a tapering and caudodorsaloriented maxillary process that forms the ventral border of the external nares and a reduced nasal process that forms the rostral border of the external nares. The ventral margin of the premaxilla is ventrally offset from the maxilla's ventral margin as seen in many ornithopods, which is considered a plesiomorphy in neornithischians (Norman et al. 2004). The dorsoventral distance between the premaxillary toothrow and the maxillary toothrow is about 2mm. In lateral view, the nasal process of the premaxilla is reduced and does not contact the nasals, possibly due to damage. The maxillary process (i.e., subnarial process) of the premaxilla is large and rostrodorsally contacts the nasal and posteriorly approaches the lacrimal. A premaxilla-lacrimal contact, which is observed in Changmiania (Yang et al. 2020) and Jeholosaurus (Barrett and Han 2009), may be absent or obscured in *Pulaosaurus*. The main body of the premaxilla is dorsoventrally concave with its caudal margin being dorsoventrally convex. The lateral surface of the premaxilla's main body is moderately concave and thickens along the oral margin. Rostral to the first premaxillary tooth, a short edentulous region develops anteriorly into a short beakshaped process, which is also present in *Jeholosaurus* (Barrett and Han 2009), *Changchunsaurus* (Jin et al. 2010), and *Hypsilophodon* (Norman et al. 2004). The rostral margin of the premaxillary main body is rugose, suggesting the presence of a rhamphotheca on the premaxilla. This character is considered a plesiomorphy among neornithischians and is observed in Lesothosaurus (Sereno 1991), Jeholosaurus (Barrett and Han 2009), Changchunsaurus (Jin et



242	al. 2010), and Hypsilophodon (Norman et al. 2004). The fossa-like depression along the
243	premaxilla-maxilla boundary is absent, whereas it is present in Haya (Barta and Norell 2021),
244	Orodromeus (Scheetz 1999), Jeholosaurus (Barrett and Han 2009) and Changchunsaurus (Jin et
245	al. 2010). There is no arched diastema between the premaxilla and the maxilla, which is a
246	synapomorphy within Heterodontosauridae (Norman et al. 2004). A premaxilla foramen,
247	observed in Haya (Berta and Norell 2021), Orodromeus (Scheetz 1999), Jeholosaurus (Barrett
248	and Han 2009), Changchunsaurus (Jin et al. 2010) and Agilisaurus (Peng 1992), is also absent.
249	Moreover, there is no prominent narial fossa and the ventral region of the premaxilla does not
250	flare laterally to form the floor of the narial fossa. These morphological characters suggest the
251	premaxilla of <i>Pulaosaurus</i> resembles that of early-diverging neornithischians (Norman et al.
252	2004).
253	Five bulbous and unserrated premaxillary teeth are present (FIGURE 2, 4), the same number as
254	Haya (Barta and Norell 2021), Orodromeus (Scheetz 1999), Changchunsaurus (Jin et al. 2010),
255	but different from the six found in Jeholosaurus (Barrett and Han 2009), Lesothosaurus (Sereno
256	1991) and Agilisaurus (Peng 1992); four in Convolosaurus marri (Andrzejewski et al. 2019).
257	The premaxillary tooth number of <i>Pulaosaurus</i> is rather primitive compared to later-diverging
258	than tooth counts of most early-diverging taxa within Neornithischia. The first premaxillary
259	tooth is close to the apex of the premaxilla with the second tooth arranged closely to the first.
260	The first premaxilla tooth is smaller than the following four teeth which are all subequal in size
261	to each other. In lateral view, the premaxillary tooth crown is recurved and spade-shaped while
262	the root is long and straight, with the crown moderately expanded mesiodistally and
263	labiolingually above the root. The labial surface of the tooth is smooth, and with no obvious
264	carinae visible. There are wear facets on the distal surfaces of the labial side of the 2nd, 3rd, and



5th premaxillary tooth. The premaxillary tooth row is slightly offset laterally from the maxillary
tooth row. No obvious diastema is present between the premaxillary tooth row and the maxillary
tooth row.
Maxilla—The left maxilla is preserved in lateral view (FIGURE 2, 3). It is an elongate, plate-
like, trapezoidal bone consisting of a straight, tooth-bearing ramus. The subtriangular ascending
process on the rostral end of the maxilla contacts the lacrimal dorsally. The premaxilla overlaps
the rostral region of the maxilla, so the ascending process on the rostral end of the maxilla that is
observed in many early-diverging neornithischians (Norman et al. 2004; Barrett and Han 2009;
Peng 1992; Sereno 1991; He et al. 1984) is not fully exposed. The maxilla forms the rostral and
ventral borders of the subtriangular antorbital fossa, the dorsal and caudal borders of which are
formed by the lacrimal. The maxilla comprises the entire ventral border of the antorbital fossa.
Due to taphonomic compression, it is unknown whether additional openings within the antorbital
fossa are present or not. It is also unknown whether the anterolateral boss that articulates with the
premaxilla is present or not because the rostral region of the maxilla is overlapped by the
premaxilla. The maxillary fenestra is absent. There is no slot in the maxilla for the lacrimal. The
tooth-bearing ramus mediolaterally widens and the maxillary tooth row is medially inset to form
a buccal emargination as seen in most ornithischians (Norman et al. 2004), which is a
plesiomorphy in neornithischians (Norman et al. 2004). A ridge extends along the dorsal border
of the caudal half of the buccal emargination, starting from the 7th maxillary tooth and extending
laterodorsally with its caudal end tapering and flaring laterally. Three or four nutrient foramina
lie on the maxillary lateral surface. It is uncertain how many nutrient foramina existed in life due
to poor preservation (FIGURE 2, 3).





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(FIGURE 2, 4). Except for the 8th replacement tooth, the apicobasal and mesiodistal lengths of the teeth increase from the mesially located teeth to the central tooth before decreasing distally. Each maxillary tooth is subtriangular with a distinct neck and a cingulum between the crown and the root, as is the case in many early-diverging ornithischians (Norman et al. 2004; Peng 1992; Sereno 1991; He et al. 1984). The labial surfaces of the maxillary teeth are relatively smooth with worn faces but no apicobasally extending ridges are present, contrary to Jeholosaurus (Barrett and Han 2009), Changchunsaurus (Jin et al. 2010), Lesothosaurus (Norman et al. 2004), and Tenontosaurus (Winkler & Murray 1997). The mesial and distal margins of the teeth are ornamented with simple denticles, which is the plesiomorphic condition that is observed in Jeholosaurus (Barrett and Han 2009), Lesothosaurus (Norman et al. 2004), and Agilisaurus (Peng 1992). The numbers of visible marginal denticles from the first to the tenth maxillary tooth are as follows: one denticle on the first tooth; five on the second; seven on the third and the fourth; nine on the fifth; six on the sixth; seven on the seventh and eighth; eight on the ninth and the tenth. The number of visible marginal denticles on the maxillary teeth of *Pulaosaurus* are apparently influenced by the growth stage of the tooth, the position of each tooth, and the erosion of tooth margin. Broadly, the morphology of the maxillary teeth of *Pulaosaurus* closely resembles those of early-diverging ornithischians. Nasal (FIGURE 2)—The left nasal is taphonomically compressed with its rostral end broken, so the internarial bar is not preserved. It is unknown whether both nasals were fused in life or not.

There are 10 maxillary teeth observed in this specimen, with all preserved in labial view

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process forming the dorsal border of the external nares. The main body of the nasal contacts the

In lateral view, the left nasal is hatchet-shaped with a tapering, caudodorsally extending caudal





310 maxillary process of the premaxilla ventrally while its caudal process contacts the lacrimal 311 ventrally, overlapping the prefrontal. The contact between the nasal and the frontal is not visible due to the displacement of the bones. In life, the nasals were excluded from the antorbital fossa. 312 No obvious narial fossa (e.g. Jeholosaurus [Barrett and Han 2009]) or nasal foramina is observed 313 314 in the specimen. 315 Lacrimal (FIGURE 2)— In lateral view, the lacrimal is an inverted L-shaped bone that is 316 comprised of a robust rostrodorsal process and the thinner caudoventral process. At the dorsal 317 318 margin of the lacrimal, the lacrimal contacts the nasal rostrodorsally and the prefrontal 319 caudaldorsally. The rostrodorsal process of the lacrimal contacts the maxilla to form the dorsal 320 border of the antorbital fossa and the caudoventral process of the lacrimal contacts the maxilla to 321 exclude the jugal from the antorbital fossa. The ventral border of the lacrimal overlaps the palatine. The tip of the lacrimal caudoventral process is situated rostral to the jugal and 322 caudodorsal to the maxilla. No lacrimal foramen is visible in lateral view. 323 324 325 Prefrontal (FIGURE 2)—In lateral view, the left prefrontal is a rod-like bone which rostrally 326 contacts the lacrimal and the nasal and overlaps the frontal caudally while forming the rostrodorsal margin of the orbit. Rostrally, the prefrontal has a descending ventral process that 327 328 rostrally articulates with the rod-like palpebral. 329 Palpebral (FIGURE 2)— Among ornithischians, the palpebral is a bone that emanates from the 330 rostrodorsal margin of the orbit and projects across the latter. This element is likely homologous 331 332 with the anterior supraorbital bone, which refers to one or multiple small osteological elements





that are incorporated into the dorsal rim of the orbit (Maidment and Porro 2010). However, the ornithischian palpebral is not homologous with similar elements reported from other reptilian groups (Nesbitt et al. 2013). The left palpebral is preserved in IVPP V30936. Similar to most ornithischians (Norman et al. 2004), the palpebral of *Pulaosaurus* is a rod-like bone with its rostral end articulating with the prefrontal. In IVPP V30936, the caudal end of the palpebral is overlapped by the displaced jugal. The length of the palpebral is more than 80% of the orbit's diameter and its caudal end projects into the orbit freely, which is a plesiomorphy of neornithischians (Han et al. 2018). The palpebral does not articulate with the postorbital as seen in *Agilisaurus* (Peng 1992; He et al. 1984). There is no indication of a postpalpebral observable in *Haya* (Barta and Norell 2021) or *Thescelosaurus* (Boyd 2014).

Jugal (FIGURE 2,3)— The left jugal is preserved, though it is taphonomically displaced. In lateral view, the jugal is a plate-like bone composed of three rami and it forms the caudoventral border of the orbit and the anteroventral border of the infratemporal fenestra. Due to its displacement, its points of contact with other bones remains uncertain. In contrast with *Yinlong* (Han et al. 2018), the jugal is excluded from the antorbital fossa as observed in *Jeholosaurus* (Barrett and Han 2009), *Haya* (Barta and Norell 2021), *Hexinlusaurus* (He et al. 1984), and *Agilisaurus* (Peng 1992). This condition is a plesiomorphy in neornithischians where the jugal is excluded from the antorbital fossa. The rostral process is in rostroventrally oriented and the dorsal process is in caudaldorsally oriented. Due to displacement, the jugal's articulating surface with the postorbital is exposed and the jugal is articulated with the postorbital with a 'finger-inrecess' joint. There is a small boss on the caudalventral corner of the jugal dorsal process. The boss is similar to, but much smaller than, the ones found in *Orodromeus* (Scheetz 1999). Similar





ornaments are also present in <i>Orodromeus</i> (Scheetz 1999), <i>Heterodontosaurus</i> (Jin et	al. 2010),
Zephyrosaurus (Jin et al. 2010), and marginocephalians (Jin et al. 2010) but absent in	
Ornithopoda (Jin et al. 2010). However, the boss on the jugal of <i>Pulaosaurus</i> is located	ed on the
dorsal process of the jugal while the jugal boss is located on the ventral margin of the	jugal in
Orodromeus (Scheetz 1999), Heterodontosaurus (Jin et al. 2010), and Zephyrosaurus	(Jin et al.
2010). The homology of these ornaments in different taxa is questionable and beyond	the scope
of this project. It is possible that the size of the jugal boss is ontogenetically variable	as is the
case in <i>Orodromeus</i> (Scheetz 1999). The dorsal branch of the caudal process is weak	ly expanded
caudodorsally while the lower, dorsoventrally wider branch expands caudoventrally.	A
bifurcated caudal process is a synapomorphy present in Lesothosaurus (Norman et al	2004),
Jeholosaurus (Barrett and Han 2009), Changchunsaurus (Jin et al. 2010), and psittace	osaurs (You
& Dodson 2004) but is absent in many early-diverging cerapodans including pachyce	ephalosaurs
(Jin et al. 2010) and Yinlong (Jin et al. 2010). There are no ornamentations on the late	eral side of
the jugal, which contrasts what is observed in <i>Jeholosaurus</i> (Barrett and Han 2009).	
Frontal (FIGURE 2)— In lateral view, the left frontal is an elongated, narrow, and for	rms an
arched shelf that comprises the rostrodorsal roof of the skull. This character is a synap	omorphy
in early-diverging neornithischians (Barrett and Han, 2009). The rostrodorsal margin	articulates
with the prefrontal, the rostroventral corner articulates with the palpebral, and it caud	ally
contacts the parietal. The contact between the frontal and the nasal is not visible due	to the
taphonomic displacement of the bones and the frontal contacts the postorbital caudov	entrally.
Although the exact ratio between frontal's length and width is unknown, the value is	estimated to
be about 2.5. The estimated length and width ratio is smaller than that of <i>Jeholosauru</i>	s (Barrett





and Han 2009), *Agilisaurus* (Peng 1992), *Hypsilophodon* (Barrett and Han 2009), and *Zephyrosaurus* (Jin et al. 2010) but greater than that of early-diverging ceratopsians (Barrett and Han 2009), *Hexinlusaurus* (Barrett and Han 2009), *Orodromeus* (Scheetz 1999) and *Thescelosaurus* (Barta and Norell 2021; Barrett and Han 2009). The lateral margin thins and flares laterally, thus forming the orbital margin. The orbital margin is not as rugose as other ornithischians. There exists a groove on the rostrodorsal margin of the orbital margin that articulates with the prefrontal and curves ventrally. The dorsal margin of the frontal thins and projects caudodorsally, which makes the dorsal margin convex and where it reaches its highest point over the orbit. Between the frontal and the parietal, the suture is preserved as a short scarfjoint that is located behind the orbit. Caudally, the margin of the frontal is overlapped by the parietal.

Postorbital (FIGURE 2)— In lateral view, the postorbital is a triradiate-shaped bone that is composed of an infratemporal process that extends rostrodorsally, a jugal process that extends rostroventrally, and a squamosal process that extends caudally. The postorbital and the squamosal form the temporal bar, the former of which contributes more. The surface of IVPP V30936's postorbital is smooth and lacks ornaments on its surface while ornamentation on the postorbital is a synapomorphy of *Jeholosaurus* (Barrett and Han 2009), *Haya* (Barta and Norell 2021), and *Orodromeus* (Scheetz 1999). The orbital margin of the postorbital has no protuberance projecting into orbit as observed in *Haya* (Barta and Norell 2021) and *Orodromeus* (Scheetz 1999). Rostrally, the end of the infratemporal process is broken from the infratemporal process. However, based on the length of the infratemporal process and its broken rostral end, it is assumed that the length of the infratemporal process is about 11.4 mm while the length of the





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squamosal process is about 7.5mm. The infratemporal process is articulated with the caudoventral corner of the frontal. The squamosal process tapers into a narrow tip where it articulates with the squamosal ventrally. In IVPP V30936, the length of the squamosal process suggests that the rostrocadual length of infratemporal fenestrae is relatively short. The jugal process is overlapped by the quadrate, so its specific morphological characters and specific contact with the jugal remain unknown. However, based on the morphology of the jugal's dorsal process, the rostrocadual width of the jugal process of the postorbital decreases, ends on the orbital margin and it articulates with the jugal with a 'finger-in-recess' joint.

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Parietal (FIGURE 2)— In lateral view, the left parietal is a saddle-shaped bone. Due to 412 compression and displacement, the original contacts between the parietal and other cranial 413 elements are unknown, so the description is based on the observable contacts between cranial elements. The parietal forms the caudodorsal roof of the skull, the infratemporal fenestrae, and 414 part of the supratemporal fenestrae. Rostrally, the parietal contacts the frontal along with the postorbitals and squamosals laterally. The caudal margin of the parietal flares laterally to form 416 the nuchal crest as is A morphological feature is observed in *Haya* (Barta and Norell 2021), 418 Changmiania (Yang et al. 2020), Orodromeus (Scheetz 1999), and Oryctodromeus (where the nuchal crest is formed by the supraoccipital) (Krumenacker 2017) and considered as a adaptative 419 characteristic for digging. Along the supratemporal fenestrae, the parietal forms the caudal margin. It is unknown whether a sagittal crest or a median process inserts between the frontals.

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Squamosal (FIGURE 2)— In lateral view, although overlapped by the squamosal process of the postorbital, the squamosal is a small and triradiate bone; however, the quadrate process has been





taphonomically broken away. The dorsal surface of the squamosal is smooth and expands laterally. In rostrolateral view, the quadrate process is observable and its lateral surface forms a glenoid on the quadrate process where the squamosal articulates with the quadrate, as is observed in early-diverging ceratopsians such as *Yinlong* (You & Dodson 2004), *Hualianceratops* (You & Dodson 2004), and *Liaoceratops* (You & Dodson 2004). The contacting relationships between the squamosal and other bones remain uncertain due to the displacement of bones.

Quadrate (FIGURE 2)— The left quadrate is exposed in lateral view. Two-thirds of the upper quadrate shaft is bowed rostrally. The proximal head of the quadrate is bluntly round. Distally, the condyles are overlain by the displaced pterygoid while the quadrate articulates with the squamosal proximally. Due to displacement, the exact contact relationships between the quadrate and other cranial bones are unknown. The shaft of the quadrate flares laterally to form the rostrolateral-oriented jugal wing of the quadrate, which arises from the dorsal margin and terminates just above the distal condyle. The pterygoid wing and the distal condyle are overlapped, so their morphological characteristics are unknown. However, it is hypothesized that the quadrate condyle is ventrally offset from the level of the maxillary tooth row based on the position of the surangular retroarticular process. It could not be determined whether a quadrate foramen is present or not.

Quadratojugal (FIGURE 2, 3)— In lateral view, the left quadratojugal is a subtriangular and flat, thought displaced, bone. Due to displacement, the relationships between the quadratojugal and other cranial elements is unknown. Also, displacement has caused the quadratojugal to be partially overlapped by the jugal with its ventral margin covered by the ectopterygoid. The





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quadratojugal is comprised of an elongated process, a dorsoventrally thick process and a third process whose shape is uncertain. The estimated shape of the overlapped process is drawn with a dashed line. Due to displacement of the quadratojugal and overlapping of the quadrate, the exact lengths and orientations of the processes are unknown. It cannot be determined whether a quadratojugal foramen is present as is observed in neornithischians such as Hypsilophodon (Norman et al. 2004), Jeholosaurus (Barrett and Han 2009). Pterygoid (FIGURE 2, 3)— In lateral view, the medial side of the right pterygoid is exposed due to taphonomic displacement. Anatomically, the pterygoid consists of three processes: the quadrate process, the mandibular process, and the palatine process. The quadrate process is a thin, subtriangular sheet that projects caudodorsally and is comparably much smaller than the same process of *Thescelosaurus* (Boyd 2014). Medially, a caudomedially facing cup-shape that receives the basipterygoid process of the basisphenoid arises from the rostral corner of the quadrate process, where it joins with the other processes. The mandibular process is a thin, small, rostrocaudally elongate, and subtriangular process that projects dorsally arising from the caudal corner of the quadrate process. Finally, the palatine process originates from the caudomedially facing cup-shape. The palatine process extends and then tapers rostrally and it forms a lateromedially oriented tab that contacts the palatine. This tab is much shorter relative to the same tab found on the pterygoid's palatine process of *Thescelosaurus* (Boyd 2014). Ectopterygoid (FIGURE 2)— In lateral view, the ectopterygoid is a small and caudally bowed bar with a notch on the dorsal margin, similar to that of *Thescelosaurus* (Boyd 2014). The ectopterygoid extends caudoventrally where it is close to the caudal end of the maxilla. A notch





4/1	located along the ectopterygoid is interpreted as the postpalatine fenestra. Due to taphonomic
472	displacement, the exact contact relationships between the ectopterygoid and other bones remain
473	uncertain, but it is possible that the ectopterygoid contacted the palatine rostrally.
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475	Palatine (FIGURE 2)— In lateral view, the palatine is a sheet-like bone whose caudal end
476	extends laterally, forming a pyramidal boss that likely represents the point of articulation for the
477	pterygoid. Laterally, the palatine is overlapped by the lacrimal and the maxilla. Only the caudal
478	end of the palatine is observable within the orbit. Due to overlapping and mediolateral
479	compression, the overall morphology of the palatine and its contact relationships with other
480	adjacent bones remains uncertain.
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482	Laterosphenoid(?) (FIGURE 2)— In lateral view, a small, laterally extending, pyramidal
483	protuberance lies rostroroventrally to the orbital process of the postorbital. Based on its relative
484	location, it is possible that it could be part of the braincase and is interpreted as the
485	laterosphenoid. This protuberance is similar to the lateral articular head of the laterosphenoid that
486	would articulate with a socket on the ventral margin of the frontal. However, due to poor
487	preservation and overlapping, the true identification of this element, its morphology, and
488	therelationships between this element and other bones remain indeterminate.
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490	Basioccipital (FIGURE 2, 3)—The basioccipital of IVPP V30936 is disarticulated from the
491	supraoccipital and is preserved upside down relative to its life position next to the basisphenoid.
492	Rostroventrally, the basioccipital contacts the basisphenoid, but the suture between the two
493	elements is ambiguous. In lateral view, the basioccipital is an inverted saddle-shaped element, as





is the case in many neornithischians (Barta and Norell 2021; Jin et al. 2010; Peng 1992; Scheetz 1999). The basioccipital lies rostrodorsal to the axis and caudoventral to the parietal, with its rostrooventral margin overlapped by the atlas and its rostroodorsal margin. The caudal end of the basioccipital narrows dorsoventrally to form the caudally oriented occipital condyle. The shape of the occipital condyle is unknown as it is only visible in lateral view. It is also unknown how much the basioccipital contributes to the condyle or whether only the basioccipital contributes to the condyle. The rostral region of the basioccipital projects slightly ventrally and becomes dorsoventrally thicker, which then forms the caudal part of the basal tuber. As the ventral side of the basal tuber is not visible, the exact shape of the basal tuber cannot be confirmed. However, based on the ventral margin of the basioccipital, it is assumed that the basal tuber is undeveloped and the length of the basal tuber is about 2 cm.

Basisphenoid (FIGURE 2, 3)—As described above, the basisphenoid is preserved upside down and the suture between the caudal basisphenoid and the rostral basioccipital is ambiguous. In total, the length of the basisphenoid is subequal to the length of the basioccipital. The main body of the basisphenoid is visible in the lateral view, and it has the shape of a rostrodorsally oriented cubic forming the rostral part of the basal tuber. No foramina or openings are visible on the lateral side of the basisphenoid. Its rostral region tapers rostrally to support the bifurcated basipterygoid processes, which are broken away but are visible ventrally. Both basipterygoid processes are rostroventrally oriented and blunt at their ends, characteristics that are considered plesiomorphies in neornithischians (Barta and Norell 2021; Peng 1992; Scheetz 1999). The angle between the two basipterygoid processes and the main body is approximately 35° degrees.





Supraoccipital (FIGURE 2)— In lateral view, the supraoccipital is a sheet-like bone that is situated caudally to the parietal. Due to compression, there is little information about its morphological characters and contact relationships. It is unknown how much or even whether the supraoccipital contributes to the magnum foramen.

Exoccipital (FIGURE 2, 3)— Only a portion of the left exoccipital is observable in medial view. This partial exoccipital is preserved upside down with its paroccipital process taphonomically damaged. The exoccipital contacts the occipital condyle contacts the occipital condyle ventrally and forms the dorsolateral region of the foramen magnum. The rostrodorsal border of the exoccipital extends medially to form the articular surface for the supraoccipital, so it is hypothesized that the supraoccipital contributes to the dorsal margin of the magnum foramen. There is a canal ventral to the articular surface for the supraoccipital, but the function of the canal is difficult to determine due to poor preservation. The preserved portion of the paroccipital process is short and blunt. On its caudodorsal corner, the exoccipital extends moderately laterally.

Predentary (FIGURE 2)— The left predentary is in close association with the dentary and is only visible in lateral view as a mediolaterally and rostrocaudally compressed element. In lateral view, the predentary is triangular with a short ventral process. It is unknown whether the ventral process is forked or not. The rostrocaudal length of the predentary is short at approximately half the rostrocaudal length of the premaxilla, which is a plesiomorphy among neornithischians (Butler 2008). The dorsal margin of the predentary is longer than the ventral margin. The rostral





tip does not project above the main body, similar to *Haya* (Barta and Norell 2021) and *Changchunsaurus* (Jin et al. 2010). The ventral process is shorter than the lateral process.

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Dentary (FIGURE 2)—Both dentaries are preserved and observable. The left dentary is visible in lateral view and the right dentary is observable in medial view. Because the postdentary elements are displaced, the exact relationships between the dentary and other elements are uncertain. The dentary is an elongated and dorsoventrally thin element with a prominent, caudodorsally oriented process on the caudal end that contributes to the coronoid process. The dorsal and ventral margins of the dentary are subparallel to each other along most of the dentary length with the dorsal margin being greater in length than the ventral margin. Both margins converge rostrally to form the articulate surface for the predentary, which is located at the midpoint of the dentary in height, as observed in *Changchunsaurus* (Jin et al. 2010). Rostral on the dentary, a vascular from the anterior dentary foramen forms an impressed canal extending rostrally to the predentary. The lateral side of the dentary is smooth and unornamented. Four nutrient foramina are present on the lateral side and are located near the dorsal margin of the dentary (FIGURE 2). The caudoventral region of the dentary is concave and forms the articulation for the surangular and the angular. This articulation extends for one-third the length of the dentary along the dentary's caudal borderand extends to the midpoint of the dentary in height. The dentary reaches its maximal height on its caudal margin, forming a caudodorsally projecting process that contributes to the coronoid process. The medial surface of the dentary is relatively flat medialaterally when compared to the lateral side. The Meckelian sulcus is located near the ventral margin along the medial surface of the dentary. The Meckelian sulcus originates from the rostral end of the





dentary but it is unknown how far it extends caudally. The caudoventral margin of medial dentary surface is covered by the plate-like splenial.

The dentary tooth row is medially inset to form the buccal emargination (FIGURE 4), similar to the maxillary tooth row. Because the dentary tooth row is obscured by the maxillary tooth row, the exact number of dentary teeth is unknown. The toothrow stops at the base of the coronoid process. Only six dentary teeth are visible between the rostral and caudal ends of the left dentary. The shape of dentary teeth resembles the maxillary teeth. The first small dentary tooth is adjacent to the predentary, similar to many early-diverging neornithischians (Norman et al. 2004; Sereno 1991; He et al. 1984).

Splenial (FIGURE 2)—Both splenials are preserved although the rostroventral end is missing because of a taphonomical fracture of the dorsal border of the right splenial. Most of the left splenial is overlapped by the dentary, though the right splenial is observable in medial view. Although their exact contacting relationships with other bones are ambiguous due to displacement, the preservation state suggests that they cover the caudoventral margins of the dentary's medial margins. The splenial is a subtriangular sheet. Its caudal end bifurcates into a blunt dorsal process and an elongate, tapering caudoventral process. From rostral to caudal, the dorsoventral length of the right splenial thickens. The rostroroventral end of the right splenial is broken and lost, so it is uncertain whether there exists a foramen on the rostral end.

Surangular (FIGURE 2)—The left surangular is displaced and preserved upside down, thus exposing its medial side. It is a roughly flat, subtriangular element that is composed of three processes in medial view with no observable foramina along its surface. There are no foramina





observed in other neornithischians such as *Haya* (Barta and Norell 2021), *Thescelosaurus* (Boyd 2014), *Changchunsaurus* (Jin et al. 2010). Due to taphonomic displacement, the exact contacting relationships with other elements is unknown. The rostrodorsal process of the surangular is an elongate, arched, and slender process that possibly contacts the angular ventrally. At its base, the medial process of the surangular is flat. The rostroventral process is short and blunt-ended, forming the ventral border where the surangular meets the angular. The elongated, caudal-oriented process on the caudoventral corner is the retroarticular process, whose caudal end becomes concave to contact the articular.

Articular (FIGURE 2)— The articular is a subtrapezoidal element when viewed laterally. It is disarticulated from the retroarticular process and the quadrate, so its specific relationships with other elements is indiscernible. The articular surface for the quadrate is not visible. The ventral border of the articular extends laterally to form a ridge.

Angular (FIGURE 2)— The left angular is visible in lateral view and forms the caudoventral margin of the mandible. It is a subrectangular element, and, asother displaced cranial elements, the contact relationships of this bone with other elements remain uncertain. On its rostral border, the angular is mediolaterally concave to articulate with the dentary. The caudal border is mediolaterally convex and forms the possible articulating surface for the surangular.

Hyolaryngeal apparatus (FIGURE 5, Table 1)—Two arytenoids are elongate, flat, and leaf-shaped with a L-shaped cross-section and are located ventral to the mandible. The length of the arytenoids is about 80% of the dentary length. This pair of elements is similar to ossified





arytenoids observed in *Pinacosaurus* (IGM100/3186) (Yoshida et al. 2023). Each arytenoid is composed of a mediodorsal wing and a dorsolateral wing. The mediodorsal wing projects mediodorsally to form the arytenoid process which serves as an attachment site for the *M. dilator* (Yoshida et al. 2023). The dorsolateral wing widens caudally before tapering at its caudal end. The lateral edge, especially at its rostral end, is rugose, which is the possible articulation for the cricoid.

A pair of elongated, gracile rod-shaped elements (FIGURE 5) – which are interpreted here to be the ceratobranchials – are overlapped by the arytenoids and meet at a point that is caudooventral to the mandible. The length of ceratobranchials is less than 40% of the dentary length.

Postcranial elements

619 Axial skeleton

Cervical series (FIGURE 2, 6, Table 2)— A mostly complete cervical series is preserved in the specimen. Due to deformation and obscured elements, the exact number of cervical vertebrae is unknown. However, nine cervical vertebrae are assumed to be preserved in the specimen, since nine cervical vertebrae usually occur in early-diverging ornithopods (Han et al. 2012), while fewer occur in Thyreophora (Han et al. 2012) and more than nine cervical vertebrae appear in later-diverging ornithopods and ceratopsians (Han et al. 2012). The first three cervical vertebrae are visible in the lateral view with the following vertebrae being partially observable in dorsal view, though they are notably deformed and obscured. The 8th and 9th cervical vertebrae are disarticulated from the rest of the series. Most of the cervical ribs are disarticulated and scattered





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with the exception of the 5th and 6th cervical ribs. The sutures of the axis and 3rd cervical vertebra between the cervical neural arches and centra are unfused.

The atlas (FIGURE 2, 6) is visible in left lateral view. The proatlas and the atlantal ribs are not preserved. The atlantal intercentrum is disarticulated from the neural arches and is visible in rostroventral view, ventral to the axis. In ventral view, the atlantal intercentrum is subcrescentic in shape. The anterior side of the atlas intercentrum is excavated by a fossa that is the deepest on the midline and becomes shallower laterally. The lateral margins of the atlas intercentrum expand dorsoventrally to form the transverse processes to articulate with the atlantal ribs. A flange extends on the ventral margin that demarcates the anterior side of the atlas intercentrum from the ventral side. A groove is located along the ventral side of the atlas, similar to what is observed in Jeholosaurus (Han et al. 2012) and Hexinlusaurus (He et al. 1984). In lateral view, the neural arches are a pair of wing-shape elements that expand dorsolaterally and are jointed at the base. The transverse width of the base narrows from rostral to caudal. The base has two articulating surfaces, a caudodorsal one for the odontoid process and a ventral one for the intercentrum. Though taphonomically compressed, the odontoid process is caudal to the atlas. The prezygapophysis is not visible while the postzygapophysis is developed to articulate with the prezygapophysis of the axis.

The axis (FIGURE 2, 6) is well-preserved, is observable in lateral view, and is comprised of the intercentrum, the centrum and the neural arch. Dorsally, the neural spine forms a dorsocaudally oriented crest. Its relative rostrocaudal length is not as long as those of many neornithischians in which it extends beyond the caudal margin of the third cervical (Barta and Norell, 2021; Han et al. 2012; Butler et al. 2011). At the base of the neural spine, the prezygapophysis is located on the rostral end and a well-developed postzygapophysis on the



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caudal end that articulates with the prezygapophysis of the third cervical. A round diapophysis is located ventral to the prezygapophysis on the rostrolateral surface of the neural arch. The height of the neural arch is subequal to that of the centrum. The suture between the neural arch and the centrum is not fused. In lateral view, the centrum is opisthocoelous. The paraphysis is a small process present on the rostrodorsal surface of the centrum. A wedge-shaped intercentrum is rostralto the centrum. The axial rib is taphonomically disarticulated and missing. Based on the presence of both the diapophysis and the parapophysis, the axial rib was likely double-head. The 3rd cervical (FIGURE 6) is observable in lateral view though its caudal region is broken due to poor preservation. The overall shape and composition of the third cervical vertebra is similar to that of the axis except it lacks an intercentrum. Compared to the axis, the neural spine of the third cervical is much smaller and more caudally oriented. The prezygopaphysis is better developed and extends rostrally over the caudal margin of the axis centrum. The diapophysis of the 3rd cervical is smaller and more caudalventrally located. When compared with the parapophysis of the axis, the parapophysis of the 3rd cervical is larger. The following cervical series (FIGURE 6) are visible in dorsal view but are notably damaged and obscured. Their rostrocaudal length is longer than their transverse width. The 4th cervical vertebra is composed of a pair of visible, well-developed, laterocaudally oriented transverse processes, well-developed postzygapophyses, and a small, dorsocaudally oriented neural arch. The possible two-headed rib of the 4th cervical vertebra is ventral to the transverse process. The 5th cervical vertebra is comprised of a pair of well-developed prezygapophyses, a laterocaudally oriented transverse processes that articulates with a double-headed rib, and a small, dorsocaudally oriented neural arch. The 6th cervical vertebra is comprised of a small, dorsocaudally oriented neural arch, and a laterocaudally oriented transverse process that





articulates with a double-headed rib with more than twice the rostrocaudal length of the centrum of the 6th cervical vertebra. The 7th, 8th and 9th cervical vertebrae are so compressed that little morphological information can be meaningfully extracted from them.

Dorsal vertebrae and dorsal ribs (FIGURE 7)— Most dorsal vertebrae are obscured, deformed, or overlapped by the ribs and ossified tendons. Only four distal dorsal vertebrae close to the ilium can be observed in lateral view although these dorsal vertebrae are deformed. The dorsal vertebrae are amphiplatyan or moderately procoelous. Only one obvious diapophysis on one dorsal vertebrate centrum is visible. There is no unambiguous evidence of striated rims around the rostrolateral and caudolateral borders of the centra. Most dorsal ribs are obscured or disarticulated, and all are compressed. Six recognizable double-head and moderately bowed dorsal ribs are articulated with the proximal dorsal vertebrae (FIGURE 7). There is no evidence of the existence of intercostal processes.

Sacral vertebrae— All sacral vertebrae are overlapped by the ilium.

Caudal vertebrae— In lateral view, much of the caudal series of vertebrae is preserved. The preserved caudal vertebrae include two anterior caudal vertebrae (FIGURE 8 A), five middle caudal vertebrae (FIGURE 8 B), and 17 posterior caudal vertebrae (FIGURE 8 C). Most anterior caudals are damaged or missing while most of the posterior vertebrae are missing. All preserved caudal vertebrae are amphiplatyan. The anteroposteriorlength of the caudal vertebrae gradually increases while their dorsoventral height decreases. The neurocentral sutures of all caudal vertebrae are unfused.





The centra of the anterior caudais are dorsoventrany concave. The fatto between the
anteroposterior length and dorsoventral length of the vertebrae is approximately 1.5. Their neural
arches are low and all neurocentral sutures are unfused. Both prezygapophyses and
postzygapophyses extend a little beyond the anterior and posterior margins of the centrum,
respectively. The hatchet-shaped neural spines are flat and more elongate than those of the
posterior caudals, are positioned moderately anterior to the postzygapophyses extending
posterodorsally, and are dorsoventrally expanded. The chevrons are disarticulated and are an
elongated rod-shape in lateral view.
The middle caudal vertebrae are similar to the anterior caudals except that the dorsoventral
length of the middle caudal centrum is shorter than that of anterior ones. Small transverse
processes are present at the base of the neural arches. In lateral view, there are two hatchet-
shaped chevrons that are disarticulated with the centrum.
Proximodistally, the dorsoventral length of the posterior caudal vertebrae decreases and the
anteroposterior length increases to be more than twice the dorsoventral length. The
prezygapophyses extend beyond the anterior end of the centrum and the postzygapophyses do
not extend beyond the posterior end. Anteroposteriorly, the anteroposterior length of the
prezygapophyses and postzygapophyses gradually increases, the dorsoventral lengths decrease,
and both are fused into a single, unified structure. There are fourteen chevrons of the posterior
caudal vertebrae preserved, twelve of which are articulated with the anteroventral corner of the
centrum whose orientations gradually shift from vertical to horizontal in relation to the centrums.
In lateral view, the chevrons are flat and hatchet-shaped while in anterior view they are 'Y'-
shaped with two gracile proximal processes merging into one ventrally oriented tapering process.





Ossified tendons (FIGURE 7A)—Ossified tendons are present along with the vertebral column. They are found alongside the middle dorsal vertebrae and the sacral vertebrae. The ossified epaxial tendons present on caudal vertebrae are not visible in this specimen and it is possible that the ossified tendons are absent from the caudal vertebrae. In lateral view, the ossified tendons are preserved as compressed, black filaments that are arranged in a basket-like arrangement of fusiform tendons along the caudal region. They lack a lattice arrangement and are more similar to a linear bundle arrangement.

Appendicular skeleton

Scapula (FIGURE 9) — Both scapulae are heavily damaged and compressed. However, it is apparent that the scapulae formed an elongate and strap-like shape in life. Only part of the proximal plate of the right scapula, and fragments of the proximal and distal ends of the left scapula are observable in lateral view. As a result, the exact shape of either scapula remains indeterminate. The morphology and position of the glenoid are also unknown. Similar to most small ornithischians (Han et al. 2018), but different from *Koreanosaurus* (Min et al. 2011) and *Oryctodromeus* (Krumenacker 2017), the scapula is unfused with the coracoid. The acromion, positioned at the rostrodorsal corner of proximal plate, is blunt, not prominent, and does not develop into the scapular spine. Both the dorsal and the ventral margins of the proximal end of the scapula are concave. A laterally concave fragment of the distal end is located far from the proximal plate. Although the overall shape of the distal end of the scapula is unknown due to damage, the fragmentary remains suggest that the dorsoventral width of the scapula gradually decreases caudally.





Coracoid (FIGURE 9 and Table 3) — Both coracoids are preserved but displaced. The description of the orientation is based on the fully exposed right coracoid. In lateral view, the coracoid is a smooth, subquadrate plate with a concave surface. The dorsoventral length is subequal to the rostrocaudal length. The rostral border of the coracoid is relatively straight while the caudal border is strongly concave and forms an embayment that is observed in many neornithischian species (Barta and Norell, 2021; Min et al. 2011; He et al. 1984; Yang et al. 2020). The dorsal border, which is sutured with a scapula, is rugose while the ventral border is smooth. Unlike *Haya* (Barta and Norell, 2021), *Hexinlusaurus* (He et al. 1984), *Jeholosaurus* (Han et al. 2012), and many other taxa, no enclosed foramen is present laterally along the coracoid. Based on the rugose morphology of the dorsal border, the coracoid foramen could be on the dorsal border.

Sternum (FIGURE 9) — Only a proximal fragment of the left sternum is preserved. The degree of fracturing to the sternal elements has made the total shape indeterminable. The proximal fragment is mediolaterally thin. This fragment is fan-shaped along its proximal end and is dorsoventrally broad but gradually narrows from rostrocaudally, so it is likely that the sternum is a hatchet-shaped or a shafted element.

Humerus (FIGURE 10, Table 4) — Only the left humerus is preserved completely with its head partially overlapped by the coracoid. The left humerus is visible in lateral view. Generally, the humerus is similar to many small early-diverging neornithischians (Han et al. 2018), and it is a relatively gracile element compared to the humerus of *Koreanosaurus* (Min et al. 2011) and





Tenontosaurus (Winkler & Murray 1997). The length of the humerus is about half the length of the femur. The rostrocaudally thin proximal end of the humerus expands mediolaterally to form a centered head, which is rotated 37° relative to the shaft. The anterior surface of the head has a concave shape that forms the bicipital sulcus. The deltopectoral crest is damaged, so its total shape is poorly understood. The lateral tuberosity is obscured by the coracoid while the medial tuberosity is continuous with a ridge defining the proximal end concavity. The shaft is gracile and short with an elliptical cross-section. An intercondylar groove extends along the shaft to form an oval intercondylar fossa near the distal end of the humerus. The ulnar condyle is larger than the radial condyle and extends anterodistally.

Ulna and radius (FIGURE 11, Table 4)— The right and left ulnae, and right and left radii are all preserved in lateral view. Both ulnae and radii are straight, rod-like elements, both shorter than the humerus. The proximal end of the ulna is broad, and the shaft narrows distally with the distal end moderately expanded. The olecranon process of the ulna is low, which is a plesiomorphy usually observed in early-diverging neornithischians (Peng 1992; He et al. 1984). Laterally, the border of ulna's proximal end is keeled,, as is observed in *Orodromeus* (Scheetz 1999), *Changmiania* (Yang et al. 2020) and *Koreanosaurus* (Min et al. 2011). The proximal ulna has a concave and triangular articular facet for articulation with the radius. Distally, the ulna has a similar morphology and articular facet for articulation with the radius. However, the distal articular facet is smaller and shallower than the proximal one. Compared to the ulna, the radius is slender and gracile. The shaft of the radius is about 30% narrower than the ulna shaft (Table 4).





Carpals (FIGURE 11)— The left carpals are well-preserved and in loose articulation while the right carpals are preserved disarticulated with each other. Hence, our description is based on the left carpals. The carpals are composed of the ulnare, radiale, intermedium, and one distal carpal, and all are visible in dorsal view. The ulnare is a cranially convex, pyramidally-shaped block that articulates proximally with the ulna and distally with metacarpal V. The intermedium is located medial to the ulnare and is a square block with a proximally convex margin that articulates proximally with the ulna and distally with metacarpal II. Compared to the ulnare, the radiale is similar in shape but smaller in size. The radiale articulates proximally with the radius and distally both with metacarpal I and II. The distal carpal is pentagonal in outline, dorsally convex, and articulates with the third metacarpal ventrally.

Metacarpal (FIGURE 11)—All five of the left metacarpals are preserved in relative articulation while the right metacarpals are preserved but disarticulated with each other. The description is based on the left metacarpals that are visible in ventral view. Proximal ends of the metacarpals are expanded and flattened to meet with the carpals. Metacarpal 1 is mediolaterally compressed whereas metacarpals 2-5 are well-preserved. All metacarpals are observable in ventral view. The second metacarpal is the longest, followed by the third and the fourth. The fifth is the shortest. The proximal ends of the metacarpals are blunt but expanded. The shafts are constricted with the shaft of metacarpal 3 being constricted most abruptly. The distal ends of the metacarpals are moderately expanded with ginglymoid articulations. On the distal ends, the lateral condyles are larger than the medial condyles, making the metacarpals medially oriented. A shallow, oval fossa exists between the distal condyles on the ventral surfaces of each distal metacarpal.





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Carpal phalanges (FIGURE 11, Table 4)—The left carpal phalanges are preserved in articulation while the right metacarpals are preserved in disarticulation with each other. Hence, the description is based on the left carpal phalanges. The carpal phalanges are observable in lateral view. *Pulaosaurus*'s phalangeal formula is 2-3-4-3-2. The proximal-most phalanx of the digit 2 is the longest. Except for the unguals, the phalanges have ginglymoid articulations with each other. The unguals of manual digits 2 and 3 are claw-like, the ungual of manual digit 1 is subconical, and the unguals of manual digits 4 and 5 are blunt.

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Ilium (FIGURE 12A and 12B, Table 5)—Only the right ilium is visible in IVPP V30936, which is preserved upside down. The ilium is composed of an elongate and tapering preacetabular process, the main body, and a dorsoventrally deep, rostrocaudally short postacetabular process. The ischial peduncle is also compressed and damaged, so the exact shape of the ischial peduncle is unknown. Similar to *Dryosaurus* and other neornithischians (Galton 1981; Han et al. 2018), the preacetabular process is elongate, dorsoventrally narrow, and tapering. The preacetabular process is longer than the postacetabular process. The shape of the dorsal border is uncertain as it may be deformed. The acetabulum is rostrocaudally long and dorsoventrally narrow. On the rostroodorsal corner of the acetabulum, there is a supraacetabular crest that extends moderately laterally, which is a synapomorphy only seen in early-diverging ornithischians including Agilisaurus (Peng 1992) and Eocursor (Butler 2010) but absent in Heterodontosauridae (Butler 2010). The pubic peduncle is rostroventrally oriented and tapers into a stout and subrectangular process in lateral view. The ischial peduncle is also anteroventrally oriented with a sub-oval articulate surface. A sub-oval and shallow brevis fossa facing ventrolaterally is present on the ventral margin of the postacetabular process, the width of which is about 8mm.





Ischium (FIGURE 12C and 12D)— The left ischium is preserved with its proximal end overlapped by a rib and the left femur distal end overlapped by the left tibia. The ischium is a rod-like element that contacts the pubis rostroventrally. The proximal end is divided into the pubic peduncle and the iliac peduncle. In lateral view, the pubic peduncle is anterodorsally oriented while the iliac peduncle is dorsally oriented. The former is rostrocaudally broader and longer than the latter. The concavity between both peduncles is dorsoventrally shallow and rostrocaudally broad. The obturator process is small and subrectangular. Similar to the ischium of *Gilmoreosaurus mongoliensis* (AMNH FARB 30739) (Prieto-Márquez & Norell 2010), the obturator process is adjacent to the pubic peduncle and forms the obturator foramen with the pubis. The ischial shaft is in caudoventral orientation with a weakly expanded distal end.

Pubis (FIGURE 12C and 12D)— The left pubis is preserved with its proximal end overlapped by the right femur. It is a slender, mediolaterally thin, rod-like bone in caudoventral orientation. The length of the pubis shaft is subequal to the left ischium. It contacts the left ischium on the dorsal border of the shaft. Compared to the ischium, the pubic body is smaller. Due to overlapping, the prepubic process is not visible. A small bony sheet on the proximal pubic shaft indicates that the obturator opening in the pubis is a notch-shape and forms the obturator foramen with the ischium, as observed in *Haya* (Barta and Norell 2021), *Jeholosaurus* (Han et al. 2012), and *Thescelosaurus* (Brown 2011).

Femur (FIGURE 13, Table 6)— Both femora are preserved in medial view, so the greater and lesser trochanters are not visible. The distal end of the left femur is broken while the right femur is compressed with its fourth trochanter broken. The femur is robust with an elliptical cross-





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section and is longer than the humerus. Cranially, the femoral head is perpendicular to the shaft which is bowed rostrally. The neck between the head and the shaft is not visible. The fourth trochanter is pendant-shaped and is located entirely on proximal half of the shaft. Although the distal end is broken, eroded, or compressed, the medial condyle of the distal femur is larger than the lateral condyle based on the morphology of the intercondylar groove. However, due to incomplete or deformed preservation of the distal end, the exact shape of the intercondylar fossa is unknown.

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Tibia (FIGURE 14, Table 6)— Both tibiae are preserved. The left tibia is observable in lateral view while the right tibia is observable in medial view with both being elongated, rod-like bones. The tibia is longer than the femur with the ratio between the left tibia and the left femur being 1.10 while the ratio between the right tibia and femur is 1.22. Both ratios are similar to most early-diverging neornithischians (Barta and Norell 2021) and the ratio would become smaller in the more mature individuals (Barta and Norell 2021). Due to displacements of the left fibula and tarsals, the left tibia distal end overlapped by the fibula, thus obscuring the exact contacting relationship between the left tibia and the left fibula. In lateral view, the proximal end of the tibia extends rostrally. The cnemial crest is poorly developed and forms a short, anterolaterally extending ridge that is separated from the fibular condyle by the deep *incisura tibialis*. In lateral view, the fibular condyle is prominent and broader than the cnemial crest and extends lateral and proximally to form a proximodistally extending fibular crest. A tiny accessory condyle lies medial to the fibular condyle. The shaft of the tibia is elongated and the cross-section is elliptical in shape. The shaft expands moderately laterally to form a laterally convex ridge, the fibula eminence, to contact the fibula. In medial view, the distal end of the tibia expands





and the external malleolus. A shallow groove on the lateral side tibia separates the internal malleolus and the external malleolus. The external malleolus is obscured by the distal end of the left fibula while the medial malleolus on the distal end of the right tibia extends medially and anteriorly.

Fibula (FIGURE 14)— Both fibulae are preserved, but only the left one is observable in lateral view. The shaft of the observable fibula bows laterally. It lies lateral to the left tibia and is unfused with the left tibia. However, due to its displacement, the exact contact relationship between the fibula and the tibia is unknown. Compared to the left tibia, the length of the fibula is subequal to the length of the tibia while the fibula is much more gracile. In the lateral view, the proximal end of the fibula is narrow and extends anteromedially. From the midpoint to the distal third of the fibula, the shaft narrows. The distal end of the fibula gradually expands mediolaterally.

Astragalus and calcaneum (FIGURE 15, Table 7)—The astragalus and calcaneum of the right hindlimb are both preserved in anterior view. In anteroventral view, the right astragalus and calcaneum contact the distal end of the right tibia proximally and are tightly appressed without co-ossifying. The left astragalus is displaced, but it is visible in dorsal view. The astragalus and calcaneum are not fused to the distal end of tibia, similar to the condition seen in *Hypsilophodon* (Huxley 1870), *Orodromeus* (Scheetz 1999), *Haya* (Barta and Norell 2021), *Changchunsaurus* (Butler et al. 2011), *Jeholosaurus* (Han et al. 2012), and *Oryctodromeus* (Krumenacker 2017).





The length of right astragalus ventral margin is about three times the length of right calcaneum ventral margin.

The astragalus is sub-rectangular in dorsal view and subtriangular in anterior view. Only the dorsally extending anterior face and the concave dorsal astragalus surfaces are visible. The anterior face of the astragalus forms the ascending process, which is represented as a subtriangular flange that extends dorsolaterally. This character is also seen in *Gilmoreosaurus mongoliensis* (Ruiz-Omeñaca et al. 2012). The lateral margin of the astragalus is moderately convex to contact the calcaneum while the medial margin is concave and longer. A small oval articulating surface for the distal end of the fibula is located on the lateral margin, at the base of the ascending process. The proximal surface is dorsoventrally concave and forms an elliptical fossa that articulates with the tibia.

The calcaneum is square and is only visible in anterior view. Due to poor preservation, little information about the calcaneum is available.

Distal tarsals (FIGURE 15)— There are three distal tarsals preserved, which is the same number observed in *Heterodontosaurus* (Han et al. 2012). This is possibly an ornithischian plesiomorphy (Norman et al. 2011; Peng 1992). Both left and right distal tarsal 1 are preserved. The left distal tarsals 2 and 3 are preserved. Distal tarsal 1 is situated above metatarsal 1 and 2, distal tarsal 2 is situated above metatarsal 3 and 4, distal tarsal 3 is situated above metatarsal 4, and metatarsal 5 is dorsolaterally adjacent to distal tarsal 3. Distal tarsal 1 is a wedge-shaped element with a with a shallow fossa on its posterior surface. In posterodorsal view, distal tarsal 1 is an L-shaped element with a concave posterior surface. The dorsoventral height of distal tarsal 1 narrows from medial to lateral. In the posterior view, distal tarsal 2 is a block-like element, the mediolateral





width of which is much longer than its dorsoventral length. In the posterior view, distal tarsal 3 is a drop-shape element, the mediolateral width of which narrows from dorsal to ventral. A small foramen pierces the posterior surface of distal tarsal 3, a feature also observed in the posterior surface of distal tarsal 3 in *Jeholosaurus* (Han et al. 2012).

Metatarsals (FIGURE 15, Table 8)— The metatarsals are preserved as mostly complete elements. Right metatarsals I, II, III, and IV are visible in dorsal view while left metatarsals II, III, IV, and V are visible in ventral view. The proximal end of right metatarsal I is obscured by right metatarsal II. The midpoint and distal end of right metatarsal III are damaged and the distal half of right metatarsal IV is not preserved. Metatarsal V is broken. Only the proximal end is preserved and it is dorsolaterally appressed to distal tarsal 3. Metatarsals are almost in the same plane and appressed to each other throughout most of their lengths. As in most early-diverging neornithischians, metatarsal III is the longest and the stoutest with its length being about twice the length of metatarsal I. Metatarsal I is splint-like with its proximal end mediolaterally compressed and its shaft gradually widening proximodistally. In dorsal view, metatarsal II, III and IV are elongated, rod-like elements with the shafts proximodistally narrow and their distal ends moderately expanded. Shallow grooves, ginglymoid distal articular surfaces, and collateral ligament pits are present on the dorsal sides of distal ends. In the ventral view, medial condyles on the distal ends of metatarsal II, III and IV are larger than the lateral condyles.

Pedal phalanges (FIGURE 15)— Right pedal phalanx I, II, III, IV, and left phalanx II, III, IV are preserved. Due to poor preservation, it is uncertain whether phalanx V exists in this taxon. Based on the preserved digits, the phalange formula of *Pulaosaurus* is 2-3-4-5-?, which is similar to



most early-diverging ornithopods and early-diverging ceratopsians (Norman et al. 2004; You & Dodson 2004). Most of the left digits are observable in the dorsal view except for the ungual digits of phalanx II and IV which are visible in the lateral view. Most of the right digits are observable in the ventral view except for the ungual digits of phalanx II, III, and IV which are visible in the lateral view, and phalanx I which is visible in the dorsal view. Right phalanx I and IV, and left phalanx III are disarticulated from their metatarsals. Except for ungual digits, the proximal and distal ends are expanded while the shafts are constricted. In dorsal view, except for III-1 and III-2, all of the left digits have dorsal lips on proximal ends that articulate with the extensor grooves of the preceding phalange distal ends. Deep oval collateral ligament pits on distal ends of the dorsal surface of III-1 and III-2.On the ventral sides of the right phalanges, lateral and medial condyles on distal ends are subequal. The intercondylar grooves are deep. All ungual digits preserved are claw-like. The ungual of phalanx III is longest, followed by phalanx II, IV and I (Table 8). The ungual of phalanx II has the greatest curvature among the pedal unguals while others are subequal.

Gut contents (FIGURE 16)— On the posteroventral corner of the specimen's thoracic cavity, variegated impressions and flat pebbles can be found. The shape of these impressions is not the same as each other and the diameter of these impressions ranges from 3mm to 8mm, obviously larger than the diameter of the surrounding matrix. The exact number of these impressions is unknown as it is hard to distinguish them from the surrounding matrix and their outlines are ambiguous. Different morphological characters and scattered distribution among these impressions make it impossible to be ovarian follicles as ovarian follicles that are preserved in fossil enantiornithine specimens are circular and uniform in size (Bailleul 2020; O'Connor et al.





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2014; Wang et al. 2016). These oval or elliptical marks and flat peddles are similar to impressions and cavities preserved in *Minmi* which are considered to be left by the displaced plant seeds (Molnar 2000).

Therefore, it is possible that the gut contents, which are likely plant seeds, are preserved in this specimen. However, the actual identity of this thoracic anomaly needs further study that is beyond the scope of this project.

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Discussion

Phylogenetic Analysis

The first analysis produced 26901 most parsimonious trees of 1225 steps, a consistency index (CI) of 0.36, and a retention index (RI) of 0.71. The resolution of the strict consensus tree (FIGURE 17) is lower than the strict consensus tree by Han et al. (2018). Most clades supported by the previous strict consensus tree by Han et al. (2018), except for Heterodontosauria, Pachycephalosauria, Iguanodontia, Neoceratopsia, Thyreophora, are not supported by this strict consensus tree. Most early-diverging ornithischian taxa and early-diverging neornithischian taxa form a polytomy which makes it difficult to recover the phylogenetic relationships between early-diverging ornithischian and early-diverging neornithischian taxa, especially the phylogenetic position of *Pulaosaurus*. This is possibly because of the incompleteness of morphological characters found in unstable taxa, as the results and resolution of different phylogenetic analyses also vary with the sampling taxa, the sampling characters and the completeness of specimens (Brown et al. 2022). The reduced consensus tree (FIGURE 18) is from 572 of the most parsimonious trees of 1210

steps, with a consistency index (CI) of 0.37 and a retention index (RI) of 0.71. The resolution of





995	the reduced consensus tree is much higher compared to the strict consensus tree. It supports most
996	clades established by previous analyses (Han 2017). The reduced consensus tree recovers
997	Pulaosaurus as one of the most early-diverging taxa of neornithischians together with
998	Agilisaurus. This result agrees on the topology of previous analyses (Boyd 2015; Han et al.
999	2018; Dieudonné et al. 2020; Butler 2008).
1000	The monophyly of Neornithischia is supported by the combination of following
1001	synapomorphies: buccal emargination on the maxilla; both of the quadrate condyles are subequal
1002	in size; the frontal does not participate in the supratemporal fenestra; a longitudinal ridge is
1003	present along the basioccipital; a well-developed coronoid process is present on the mandible;
1004	the external mandible fenestra is absent; the prepubic process is rod-like or dorsoventrally
1005	compressed. The combination of following synapomorphies support the monophyly of
1006	Cerapoda: enamel distribution on cheek teeth is asymmetric; the fossa trochanteris is modified
1007	into a distinct constriction that separates the head and the greater trochanter of the femur; the
1008	anterior trochanter of the femur is closely appressed to the greater trochanter. Based on these
1009	phylogenetic analysis results, as the anterior side of <i>Pulaosaurus</i> femur and the sacrum are not
1010	visible in this specimen, it is possible that <i>Pulaosaurus</i> could be a member of Cerapoda.
1011	The second phylogenetic analysis based on Fonseca et al. (2024) dataset produces 200 most
1012	parsimonious trees of 7085 steps, a consistency index (CI) of 0.16, and a retention index (RI) of
1013	0.62. The analysis recovers <i>Pulaosaurus</i> as the earliest-diverging neornithischian (FIGURE
1014	19), a result similar to that produced by the first analysis.
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1016	Compared to later-diverging neornithischians, many plesiomorphies of Neornithischia are
1017	maintained in <i>Pulaosaurus</i> : there are five premaxillary teeth; the first maxillary tooth is close to





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the posterior margin of the premaxilla; the maxillary teeth are subtriangular without ridges on the labial surfaces; the forelimbs are relatively shorter compared to the hindlimbs; there are five digits on the manus; the manual ungual digits are claw-like in shape; a supra-acetabular crest is located on the ilium; there are three unfused distal tarsals. The above listed plesiomorphies suggest *Pulaosaurus* is one of the earliest-diverging neornithischians. However, there are also derived synapomorphies also present in *Pulaosaurus*. For example, the posterolateral process of the *Pulaosaurus* premaxilla contacts the lacrimal, which is a synapomorphy of iguanodontian taxa such as Dryosaurus, Iguanodon, and Ouranosaurus (Norman et al. 2004); the frontals of Pulaosaurus are elongated and narrow which is more similar to early-diverging Ornithopoda taxa (Norman et al. 2004); the dorsal surface of the squamosal is flat and expands laterally, which is a synapomorphy of Pachycephalosauria and certain early-diverging ceratopsians such as Yinlong and Huayangceratops (You & Dodson 2004). These features suggest that mosaic acquisition of traits has occurred in the course of neornithischian evolution and some synapomorphies of later-diverging neornithischian taxa have appeared early in basal taxa. However, it should be noted that the results described herein are unstable due to the incompleteness of specimens and the lack of certain morphological characteristics. To solve the problems about early-diverging neornithischian phylogeny and recover the actual systematical position of *Pulaosaurus*, more complete specimens are required in future research. New information of distribution of early-diverging neornithischian taxa Taxa of early-diverging Neornithischia in China are mainly found in Middle Jurassic strata throughout southwestern China: Agilisaurus (Peng 1992), Hexinlusaurus (Barrett et al. 2005), Yandusaurus (He et al. 1984) are found in Lower Shaximiao Formation of Province Sichuan and Sanxiasaurus (Li et al. 2019) are found in Xintiangou Formation of Chongqing. Agilisaurus and



Hexinlusaurus are considered as the most early-diverging neornithischian taxa by most analyses
(Boyd 2015; Dieudonné et al. 2020; Butler 2008) while Sanxiasaurus is the earliest record of
Neornithischia in Asia (Li et al. 2019). The fossil record of Neornithischia in northern China is
limited but includes: Jeholosaurus (Han et al. 2012) and Changmiania (Yang et al. 2020) found
in Province Liaoning, Changchunsaurus (Jin et al. 2010) found in Province Jilin, all of which are
from Lower Cretaceous strata (Yang et al. 2020; Liyong Jin et al. 2010; Han et al. 2012). There
is a temporal and geographical gap between early-diverging taxa found in the Middle Jurassic
strata of southwestern China and late-diverging taxa found in the Lower Cretaceous strata of
northeastern China. The missing evolutionary link of Neornithischia is an early-diverging
neornithischians in the Late Jurassic strata between southwestern and northeastern China.
Pulaosaurus helps to serve as this 'missing link' as it is found in the Late Jurassic strata found in
Province Hebei, which is between southwestern and northeastern China, helping fulfill the
temporal and geographical gap. Moreover, ornithischian taxa that are present in Jurassic fauna,
including the Shishugou and Shaximiao faunas, are missing in the Yanliao Biota (Liu 2022),
which usually play the role of small and middle-bodied herbivores in the Mesozoic ecosystems.
Pulaosaurus is found in the Tiaojishan Formation of Province Hebei where it is geographically
situated between southwestern China and northeastern China. The most recent dating of the
bottom of the Tiaojishan Formation in the Xuanhua District—Zhuolu County area of
Zhangjiakou City, produces an age of approximately 164. 3±2. 6 Ma (Bai et al. 2024).
Alternatively, Wu et al. (2024) proposed an age range of 153 Ma to 162 Ma based on zircon U-
Pb dating of the samples from borehole Yang D1 in Western Liaoning within the Tiaojishan
Formation (Wu et al. 2024). Regardless of the dating study, the age of Tiaojishan Formation is
reliably recovered as being deposited between the Late Middle Jurassic to the Early Late





Jurassic. As a result, the discovery of *Pulaosaurus* fills the temporal and geographic gap of neornithischian fossil record in China, and provides new information on the biodiversity of the Yanliao Biota. Huang (2015) proposed that there was geographical isolation between northeastern China and other regions during the Middle and Late Jurassic, which inhibited the dispersal of organisms between faunas and made the species composition of the Yanliao Biota unique from other Jurassic faunas (Liu 2022). However, the discovery of *Pulaosaurus* in Upper Jurassic Tiaojishan Formation also suggests that Asian neornithischian taxa originated near or within the Middle Jurassic-aged Sichuan Basin (Li et al. 2019) and spread to northern China during the Late Jurassic and the Early Cretaceous (Yang et al. 2020; Jin et al. 2010; Han et al. 2012). This indicates that the geographical isolation of the Yanliao Biota was not as large of a preventative faunal barrier between other regions as hypothesized and the biodiversity of the Yanliao Biota has been severely underestimated.

Hyolaryngeal apparatus and acoustic function of *Pulaosaurus*

The hyolaryngeal apparatus of Archosauria comprises the following elements: one basihyal, a pair of ceratohyals, one pair of ceratobranchials, one pair of cricoids, and one pair of arytenoids (Yoshida et al. 2023; Friedman et al. 2018; Hill et al. 2015) (Fig 20). Additionally, one procricoid, one pair of epibranchials, and one paraglossal are only present in Aves (Yoshida et al. 2023; Friedman et al. 2018; Hill et al. 2015). In extant reptiles, the hyolaryngeal apparatus elements are cartilaginous with the exception of the ossified ceratobranchials (Yoshida et al. 2023). In extant Aves, the ceratobranchials, the epibranchials, and the larynx are ossified (Yoshida et al. 2023). The hyolaryngeal apparatus plays a significant role in acoustic function, airway protection, respiratory modification, and circulation assistance in tetrapods (Kirchner 1993). However, few fossilized larynx elements have been found in non-avian reptile fossils



1089	when compared to their significance in Archosauria evolution and ecology. In the case of the
1090	Dinosauria, most non-avian dinosaur specimens only preserve the first pair of rod-like
1091	ceratobranchials. Other hyolaryngeal elements preserved areas following. In Carnotaurus,
1092	Microraptor, Confuciusornis, the basihyal has been recovered (Bonaparte et al. 1990; Yoshida et
1093	al. 2023); the basihyal and ceratohyals are preserved in Saichania chulsanensis (Bonaparte et al.
1094	1990; Yoshida et al. 2023); a pair of ceratobranchials and the epihyal are preserved in
1095	Pinacosaurus granger (Morschhauser 2013); the plate-like second pair of ceratobranchials is
1096	preserved in Psittacosaurus mongoliensis (Morschhauser 2013); the splint-like second pair of
1097	ceratobranchials is preserved in Leptoceratops gracilis (Morschhauser 2013); the tetraradiate
1098	first pair of ceratobranchials and the plate-like second pair of ceratobranchials are preserved in
1099	Protoceratops andrewsi (Morschhauser 2013); and the sigmoid ceratohyal and the basihyal are
1100	preserved in Triceratops horridus (Morschhauser 2013). Of note, Pinacosaurus granger was the
1101	first dinosaur whose larynx element has been reported (Yoshida et al. 2023). Here, <i>Pulaosaurus</i>
1102	is the second reported non-avian dinosaur specimen with a preserved larynx apparatus since
1103	Pinacosaurus granger, which demonstrates that an ossified hyolaryngeal apparatus has existed
1104	more taxonomically broadly among non-avian dinosaurs rather than just in ankylosaurids.
1105	Pulaosaurus also possesses a pair of rod-like ceratobranchials, which is also preserved in
1106	Jeholosaurus and many other dinosaur species (Friedman et al. 2018).
1107	The arytenoids of <i>Pulaosaurus</i> are elongated with arytenoid processes. The length of the
1108	arytenoids is about 80% of the dentary length (Table 1). This structure is similar to the
1109	arytenoids preserved Pinacosaurus, but the arytenoid processes of Pulaosaurus are less
1110	prominent. In the case of acoustic function, the larynx functions differently between extant Aves
1111	and non-avian reptiles (Yoshida et al. 2023). In extant non-avian reptiles – such as turtles and



1112	crocodiles - the larynx functions as the vocal source (Yoshida et al. 2023; Sacchi et al. 2004;
1113	Riede et al. 2015). During phonation, the glottis is almost closed by the larynx and its
1114	surrounding muscles and ligaments and then air pressure forces the glottis to open, making vocal
1115	folds to vibrate and phonate, thus producing sounds (Yoshida et al. 2023; Sacchi et al. 2004;
1116	Riede et al. 2015). In extant Aves, the vocal source is the syrinx, which is located at the inferior
1117	end of the trachea (Kingsley et al. 2018; Yoshida et al. 2023; Sober et al. 2019), which has also
1118	been found in the Mesozoic bird Vegavis (Clarke et al. 2016). The larynx serves as part of the
1119	vocal resonator tract to improve vocal efficiency and sounds are emitted through it, which
1120	requires control over the glottal opening (Kingsley et al. 2018; Yoshida et al. 2023; Sober et al.
1121	2019). A longer arytenoid provides more attachment area for the dilator muscle, thus making its
1122	lever arm longer, which assists the arytenoid with the opening of the glottis (Yoshida et al.
1123	2023). The arytenoids, with their prominent arytenoid processes and firm cricoid-arytenoid
1124	joints, allow for the horizontal rotation of the arytenoid to open and close the glottis (Yoshida et
1125	al. 2023). Such structures allow extant birds to communicate with more complicated sounds in
1126	broader vocal ranges and with greater efficiency (Yoshida et al. 2023; Sober et al. 2019).
1127	Yoshida et al (2023) proposed that the arytenoid length is positively correlated to the mandible
1128	width and there is a distinction in the relative arytenoid size compared to the mandible between
1129	the group of the larynx vocal source and the group of the larynx vocal modifier (Yoshida et al.
1130	2023). Due to the compression of the <i>Pulaosaurus</i> mandible, the exact width of the mandible is
1131	unknown, so acoustic calculations of Pulaosaurus cannot be made. However, based on the
1132	cranial morphology, the <i>Pulaosaurus</i> mandible width is less than 8cm, which is the length of its
1133	skull. Therefore, the mandible width is shorter than the mandible width of <i>Pinacosaurus</i> , which
1134	is 10cm (Hill et al. 2015). Pulaosaurus arytenoids are subequal to those of Pinacosaurus in



1135	length (Yoshida et al. 2023). Therefore, the relative arytenoid length of <i>Pulaosaurus</i> is larger
1136	than Pinacosaurus. Pulaosaurus is likely to possess a non-laryngeal vocal source similar to
1137	Pinacosaurus although the acoustic function is more primitive as its arytenoid processes are less
1138	prominent. The larynx of <i>Pulaosaurus</i> possibly functions to modulate and enhance sounds, thus
1139	allowing <i>Pulaosaurus</i> to communicate with more complicated sounds, similar to extant birds
1140	(Yoshida et al. 2023). This suggests that a non-laryngeal vocal source was present among the
1141	Dinosauria during at least the Late Jurassic, regardless of whether non-laryngeal vocalization is a
1142	plesiomorphy of Dinosauria or it is convergent in different dinosaur taxa (Yoshida et al. 2023).
1143	As the fossilized syrinx could be found in the Mesozoic bird Vegavis (Clarke et al. 2016), it is
1144	possible that a fossilized syrinx could be found in non-avian dinosaur specimens in the future.
1145	Additionally, the preservation of ossified arytenoid in Pulaosaurus strongly suggests that
1146	ossification of the laryngeal apparatus has occurred not only in Ankylosauria and Aves (Yoshida
1147	et al. 2023) but also in Neornithischia (Fig 21). This indicates that ossified laryngeal
1148	apparatuses should have been phylogenetically widespread among non-avian dinosaurs.
1149	However, except for <i>Pulaosaurus</i> and <i>Pinacosaurus</i> , there are no other reports of ossified
1150	laryngeal apparatus preserved in non-avian dinosaur fossils. There are two possible explanations
1151	for this paucity of laryngeal anatomy within the non-avian dinosaur fossil record. Firstly,
1152	laryngeal elements are gracile elements that rarely preserve or are taphonomically destroyed
1153	prior to discovery and description. Secondly, it is possible that other ossified laryngeal elements
1154	have been preserved, but have been misidentified (Yoshida et al. 2023). For example, the
1155	cricoids and arytenoids of <i>Pinacosaurus</i> were originally incorrectly identified as the paraglossals
1156	and the first pair of ceratobranchials (Hill et al. 2015). As described above, there have been many
1157	reports that there are two pairs of ceratobranchials in many dinosaur taxa which are defined as



1158	plate-like, splint-like, or tetraradiate (Morschhauser 2013). However, the second pair of
1159	ceratobranchials are lost in extant archosaurs. It is possible that hyolaryngeal elements preserved
1160	in many non-avian dinosaur specimens that are currently identified as ceratobranchials are, in
1161	fact, ossified laryngeal elements. Reanalysis of vocal anatomy within non-avian dinosaurs needs
1162	to be carried out to assess the accuracy of identification among curated specimens.
1163	Rod-like ceratobranchials are observed in Pulaosaurus. The length of Pulaosaurus
1164	ceratobranchials is less than 50% of the dentary length and their relative length is subequal to
1165	those of Jeholosaurus (Barrett and Han 2009) but shorter than those of Paraves and quadrupedal
1166	ornithischians (Friedman et al. 2018). Elaborate ossified hyoid elements are typically observed in
1167	Aves, pterosaurs, and quadrupedal ornithischians such as ankylosaurids and hadrosauroids,
1168	which increases the mobility of the tongue and makes up for the diminished utility of forelimbs
1169	(Friedman et al. 2018). In pterosaurs, the ceratobranchials are elongated and fused (Friedman et
1170	al. 2018). In Aves, epibranchials arise to increase the overall length of the hyoid element. In
1171	quadrupedal ornithischians, the frequency of ossification of hyoid elements increases (Friedman
1172	et al. 2018). The elongation of the ceratobranchials supports the mobility of the avian tongue
1173	(Friedman et al. 2018), which is closely associated with the feeding and ecological radiation of
1174	Aves.
1175	The short relative length of Pulaosaurus' ceratobranchials suggests that the tongue mobility of
1176	Pulaosaurus would have been limited. This may have been because Pulaosaurus was an obligate
1177	biped with its forelimbs used for food acquisition and processing. The limited tongue mobility
1178	and primitive tooth morphology of <i>Pulaosaurus</i> also indicate that the food intraoral processing
1179	of <i>Pulaosaurus</i> is less prominent than later-diverging ornithischian taxa (Friedman et al. 2018)
1180	and it could only feed on softer food.



118111821183	Conclusions Pulaosaurus qinglong gen.et sp. nov. is an early-diverging neornithischian species found in
1184	the Upper Jurassic Tiaojishan Formation of Province Hebei, China. A phylogenetic analysis
1185	places Pulaosaurus at the base of Neornithischia close to Agilisaurus, which is the earliest-
1186	diverging neornithischian. Pulaosaurus represents the first neornithischian found in the Yanliao
1187	Biota, and helps to fill the temporal and geographical gap in the distribution of Neornithischia
1188	within China. A pair of arytenoids are preserved in the <i>Pulaosaurus</i> holotype and represents the
1189	second case of an ossified laryngeal apparatus among non-avian dinosaurs. The arytenoids of
1190	Pulaosaurus indicates that ossified laryngeal apparatuses were present in Neornithischia, thus
1191	suggesting that the ossified laryngeal apparatus could be widespread across Dinosauria. As the
1192	morphology of <i>Pulaosaurus</i> arytenoids resembles the arytenoids of extant birds, it is possible for
1193	Pulaosaurus to have an avian-like vocalization.
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1372	

The photograph of the whole skeleton of Pulaosaurus qinglongin left lateral view (IVPP V30936)

Photograph provided by Hailong Zhang.



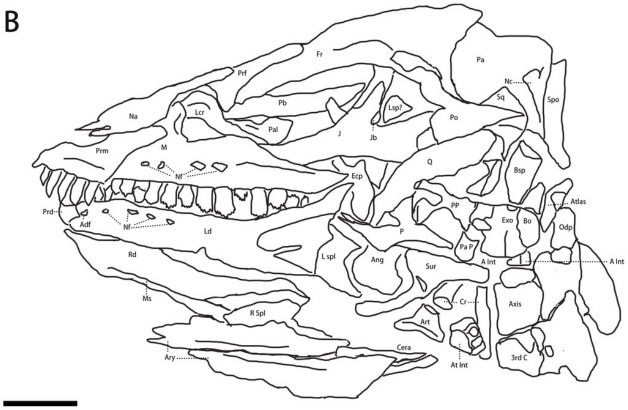


The skull of Pulaosaurus qinglongin left lateral view (IVPP V30936).

(A) Photograph. (B) Outline drawing Abbreviations: Adf-Anterior dentary foramen; Ang-Angular; Art-articular; Ary-arytenoid; A Int-Axial intercentrum; At Int-Atlantal intercentrum; Bo-Basioccipital; Bsp-Basisphenoid; Cera-Ceratobranchial; Cr-Cervical rib; Ecp-Ectopterygoid; Exo-Exoccipital; Fr-Frontal; J-Jugal; Jb-Jugal boss; Lcr-Lacrimal; Ld-Left dentary; Lsp?-Laterosphenoid?; L spl-Left splenial; M-maxilla; Ms-the Meckelian sulcus; Na-Nasal; Nc-Nuchal crest; Nf-Nutrient foramina; Odp-Odontoid process; P-Pterygoid; Pa-parietal; Pal-Palatine; Pa p-Paraoccipital process; Pb-Palpebral; Po-Postorbital; PP-Pterygoid process of basisphenoid; Prd-Predentary; Prf-Prefrontal; Prm-Premaxilla; Sq-Squamosal; Q-Quadrate; Qj-Quadratojugal; Rd-Right dentary; R spl-right splenial; Spo-Supraoccipital; Sur-surangular; 3 rd C-the 3 rd cervical vertebrate.



1 cm



1 cm

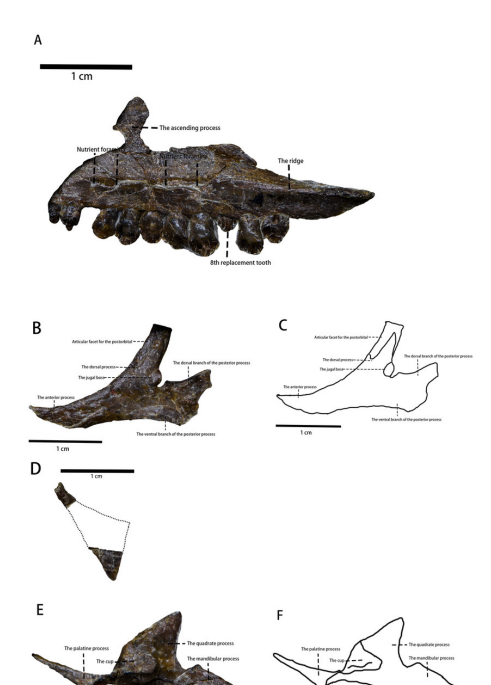


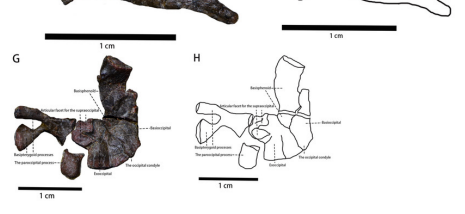
The photo of several skull elements in lateral view.

(A) The photograph of the visible part of the maxilla. (B) The photograph of the jugal. (C) The line drawing of the jugal. (D) The photograph of the quadratojugal. (E) The photograph of the pterygoid. (F) The line drawing of the pterygoid. (G) The photograph of partial braincase. (H) The line drawing of partial braincase.

Photograph provided by Hailong Zhang.

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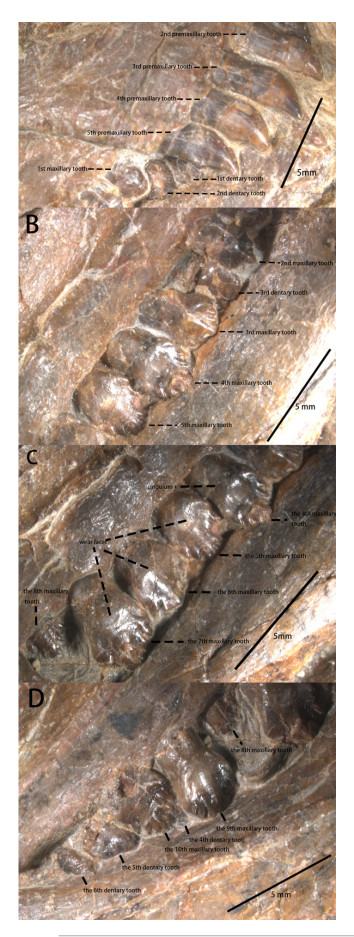


The teeth of IVPP V30936 under the microscope.

(A) The premaxillary teeth and the most anterior cheek teeth. (B) The anterior cheek teeth. (C) The middle cheek teeth. (D) The posterior cheek teeth.

Photograph provided by Yunfeng Yang.





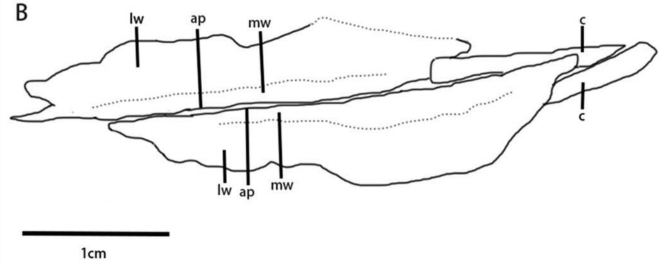
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Arytenoids and ceratobranchials of *Pulaosaurus qinglong*in laterodorsal view (IVPP V30936).

(A) The photograph of ceratobranchials and arytenoids. (B) The line drawing of ceratobranchials and arytenoids. Abbreviations: ap-arytenoid process; c-ceratobranchial; lw-laterodorsal wings of the arytenoids; mw-mediodorsal wings of the arytenoids.

Photograph provided by Hailong Zhang.

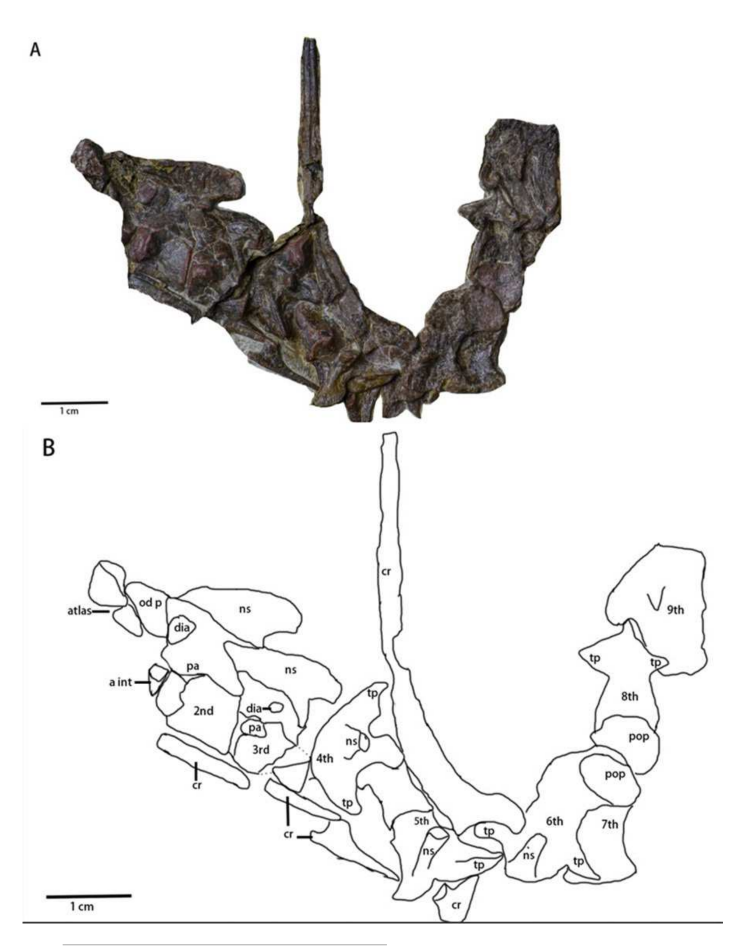






Cervical Series of *Pulaosaurus qinglong*: (IVPP V30936). Atlas, axis and 3rd cervical visible in left lateral view; from 4th cervical to 9th cervical visible in dorsal view.

(A) The photograph of the cervical series. (B) The line drawing of the cervical series. Abbreviation: a int-axial intercentrum; cr-cervical rib; dia-diapophysis; ns-neural spine; od p-odontoid process; pa-parapophysis; poppostzygapophysis; tp-transverse process.



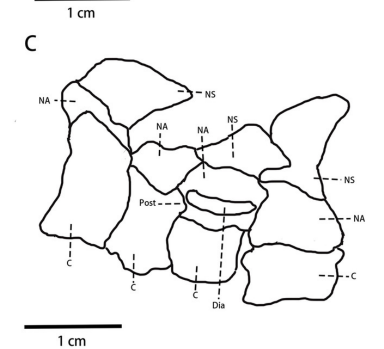


The visible dorsal vertebrae and the ossified tendons in lateral view.

(A) Dorsal ribs and ossified tendons in lateral view. (B) The photograph of four visible dorsal vertebrae. (C) The line drawing of four visible dorsal vertebrae. Abbreviations: C-Centrum; Dia-Diapophysis; NA- the Neural Arch; NS- the Neural Spine; Post- Postzygapophysis.



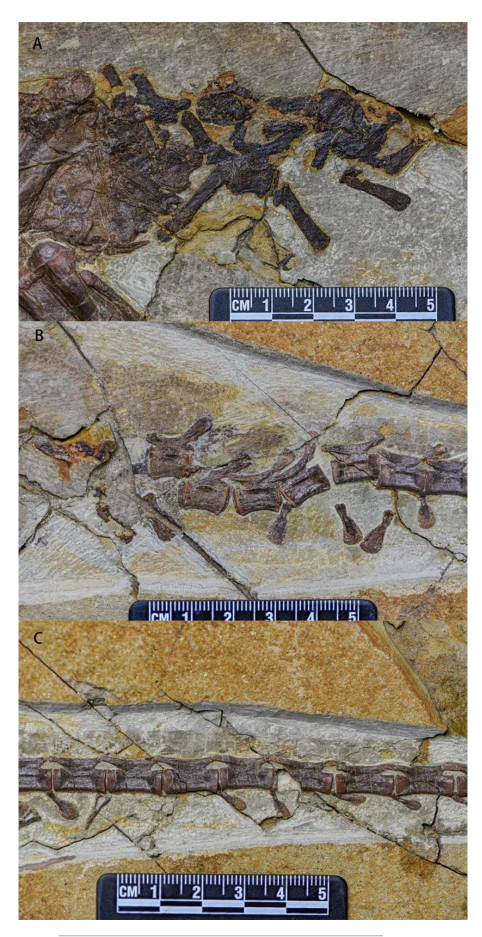






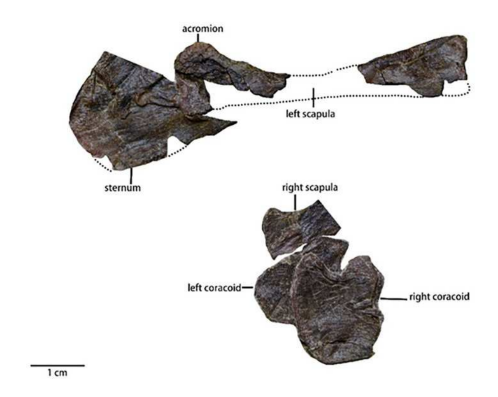
Caudal vertebrae series of *Pulaosaurus qinglong*in left lateral view (IVPP V30936).

(A) The proximal caudal vertebrae. (B) The middle caudal vertebrae. (C) The distal caudal vertebrae. Photograph provided by Hailong Zhang.

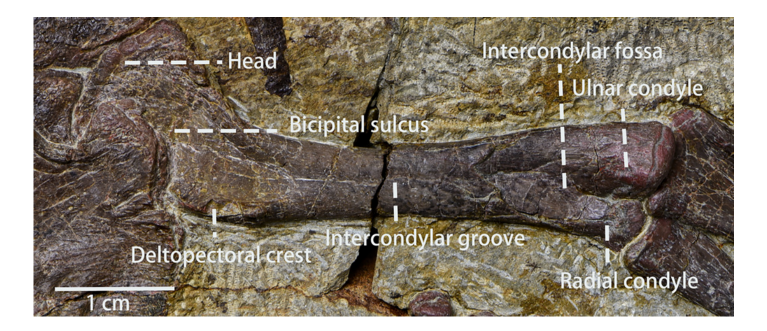


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The pectoral girdle and the sternum of *Pulaosaurus qinglong*(IVPP V30936) in lateral view.



The left humerus in lateral view.



The left ulna, the left radius and the left manus in lateral view.

Abbreviations: DC-Distal Carpal, Int-Intermedium, R-Radiale, U-Ulnare, Mc I-Metacarpal I, Mc II-Metacarpal II, Mc IV-Metacarpal IV, Mc V-Metacarpal V, I-1-Phalanx I-1, I-2-Phalanx I-2, II-1-Phalanx II-1, II-2-Phalanx II-1, II-3-Phalanx III-3, III-1-Phalanx III-1, III-2-Phalanx III-2, III-3-Phalanx III-3, III-4-Phalanx III-4, IV-1-Phalanx IV-1, IV-2-Phalanx IV-2, IV-3-Phalanx IV-3, V-1-Phalanx V-1, V-2-Phalanx V-2.



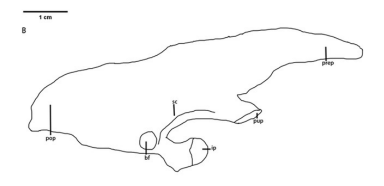


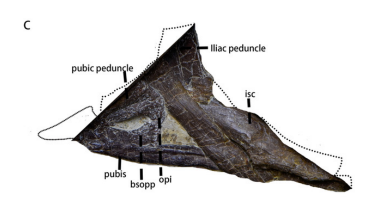
The pelvic girdle in lateral view.

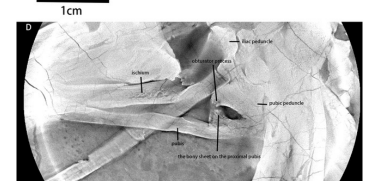
(A) The photograph of the left ilium in lateral view. (B) The line drawing of the left ilium in lateral view. (C) The photograph of the left pubis and the left ischium in lateral view, the missing part supplemented by the line drawing. (D) The CT scaning image of the left pubis and the left ischium. Abbreviations: bf-brevis fossa, bsopp-the bony sheet on the proximal pubis, ip-the ischial peduncle, isc-ischium, opi-the obturator process of the ischium, pop-the postacetabular process, prep-the preacetabular process, pup-the pubic peduncle, sc-the supra-acetabular crest. Photograph provided by Hailong Zhang.

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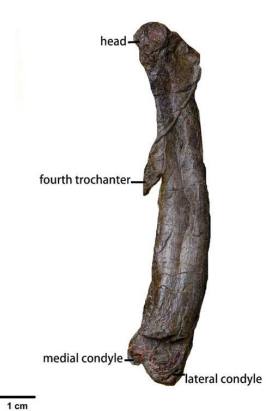
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The femora of *Pulaosaurus qinglong*(IVPP V30936).

(A) The left femur in medial view. (B) The right femur in medial view.

Α





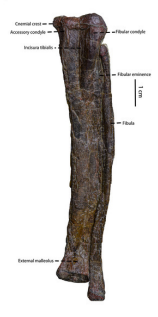
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The tibiae and the fibulae.

(A) The left tibia and the left fibula in lateral view. (B) The right tibia and the right fibula in medial view. Photograph provided by Hailong Zhang.





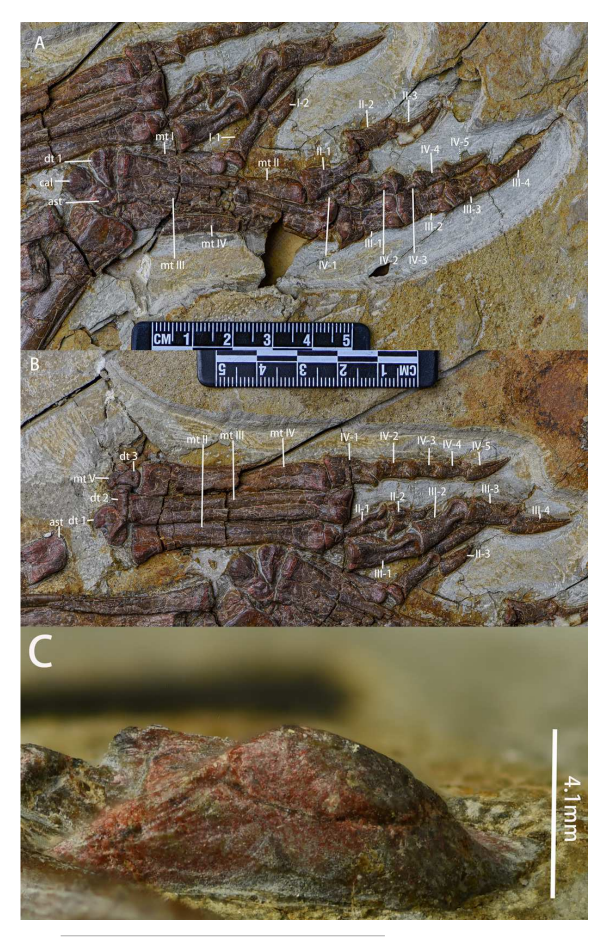


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The right and left astragali, the right and left pedes and the right calcaneum.

(A) The right astragalus and calcaneum in anterior view, right metatarsals in dorsal view and the phalanges in lateroventral view. (B) The left astragalus in dorsal view, the right metatarsals in ventral view and the right phalanges in lateral view. (C) The left calcaneum in anterior view. Abbreviations: ast-astragalus, cal-calcaneum, dt-distal tarsal, mt-metatarsal, I-1-phalanx I-1, I-2-phalanx I-2, II-1-phalanx II-1, II-2-phalanx II-2, II-3-phalanx II-3, III-1-phalanx III-1, III-2-phalanx III-2, III-3-phalanx III-3, III-4-phalanx III-4, IV-1-phalanx IV-1, IV-2-phalanx IV-2, IV-3-phalanx IV-3, IV-4-phalanx IV-4, IV-5-phalanx IV-5. Photograph provided by Hailong Zhang.



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The gut contents preserved in IVPP V30936.

P-Pebbles; I-Impressions of possible plant seeds.

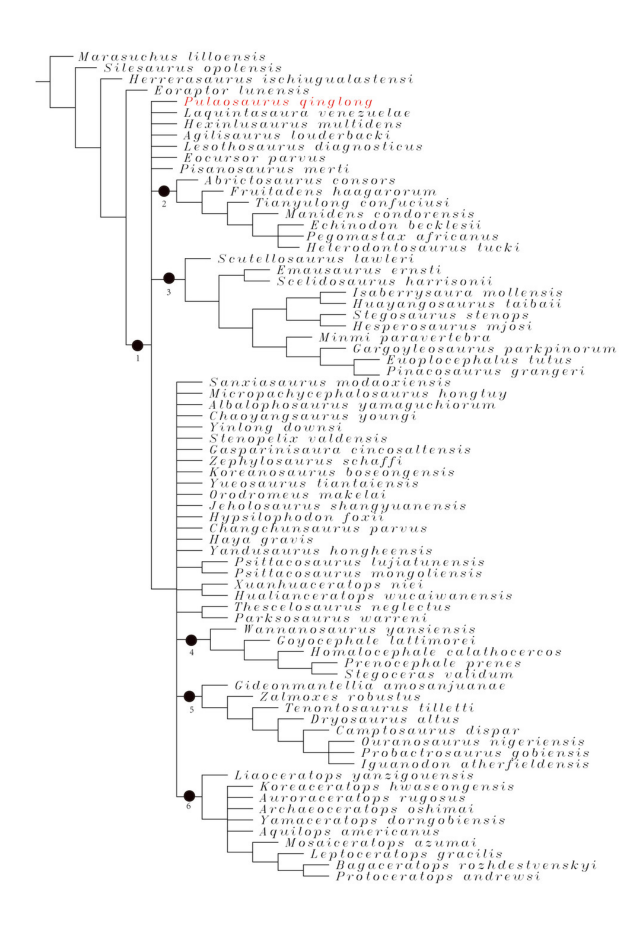


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The strict consensus tree from 26901 most parsimonious trees including 74 taxa and 380 characters generated by the analysis.

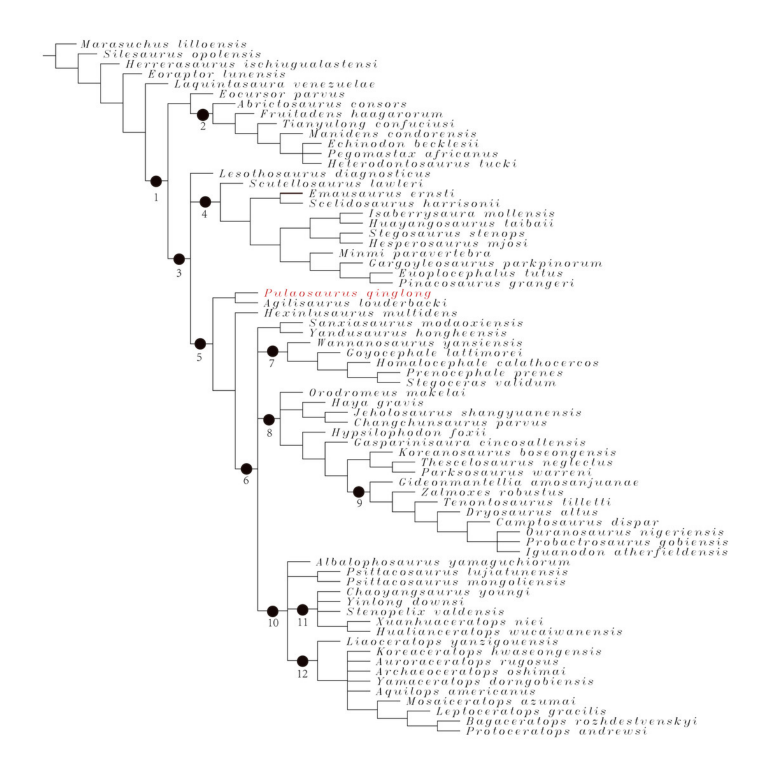
Nodes: 1-Ornithischia, 2-Heterodontosauria, 3-Thyreophora, 4-Pachycephalosauria, 5-Iguanodontia, 6-Neoceratopsia.





The reduced consensus tree from 572 most parsimonious trees including 70 taxa and 380 characters generated by the analysis.

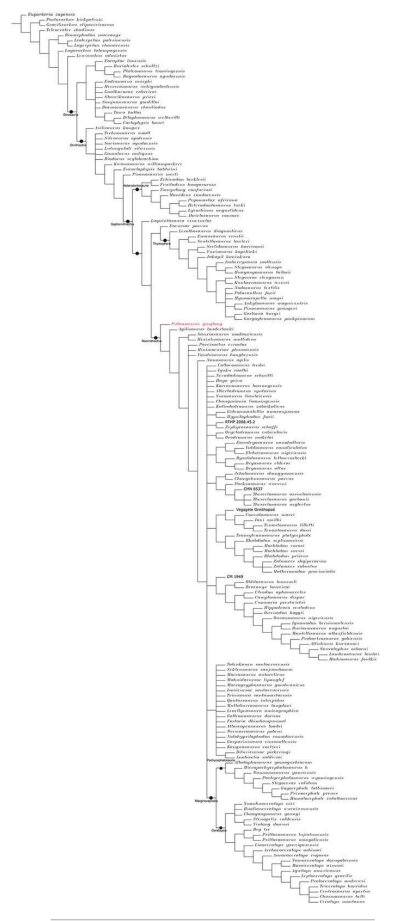
Nodes: 1-Ornithischia, 2-Heterodontosauridae, 3-Genasauria, 4-Thyreophora, 5-Neornithischia, 6-Cerapoda, 7- Pachycephalosauria, 8-Ornithopoda, 9-Iguanodontia, 10-Ceratopsia, 11-Chaoyangsauridae, 12-Neoceratopsia.





The strict consensus tree generated by analysis 2 based on the character matrix dataset provided by Fonseca et al. (2024).





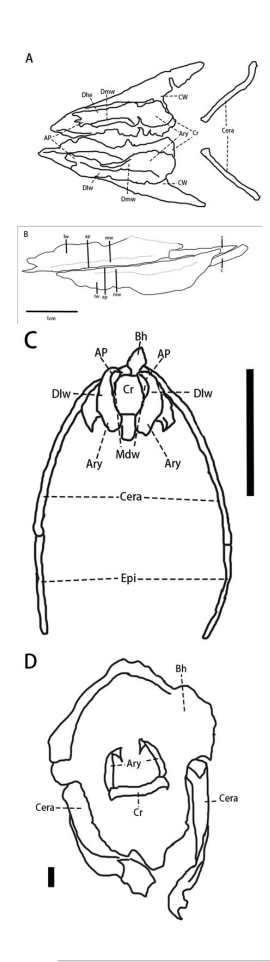


The hyoaryngeal apparatuses in different taxa of Archosauria.

☐ A ☐ The line drawing of *Pinacosaurus grangeri*(IGM100/3186) hyolaryngeal apparatus based on the 3D reconstruction. (B) The line drawing of *Pulaosaurus qinglong*(IVPP V30936) hyolaryngeal apparatus. (C) The line drawing of *Nothoproctasp*.(AMNH6502) hyolaryngeal apparatus. (D) The line drawing of *Tomistoma schlegelii*(AMNH R113078) hyolaryngeal apparatus. All the specimens involved except for IVPP V30936 are from the supplementary materials provided by Yoshida et al. (2023) Abbreviations: AP-Arytenoid Process, Ary-Arytenoid, Bh-Basihyal, Cera-Ceratobranchial, Cr-Cricoid, Dlw-Dorsolateral wing, Dmw-Dorsomedial wing, Epi-Epibranchial, lw-the laterodorsal wing, mw-the mediodorsal wing.

Illustration credit: Junki Yoshida, Ph.D.





Evolution of hyolaryngeal elements in Archosauria. The figure is adapted from illustration by Tatsuya Shinmura (Yoshida et al. 2023).

The figure is adapted from illustration by Tatsuya Shinmura (Yoshida et al. 2023). Yellow-arytenoid, green-first pair of ceratobranchials, black-second pair of ceratobranchials, red-cricoid, white-basihyal, blue-procricoid, orange-paraglossal, grey-epibranchials. Numbers represent the ancestral state of characteristics: 1-Laryngeal vocal source, 2-Loss of second ceratobranchials, 3-Procrocoid, 4-Arytenoid process, 5- Arytenoid process, 6-Ossified larynx, 7-Immobile lungs, 8-Procrocoid, 9-Paraglossal.

Illustration credit: Junki Yoshida, Ph.D.

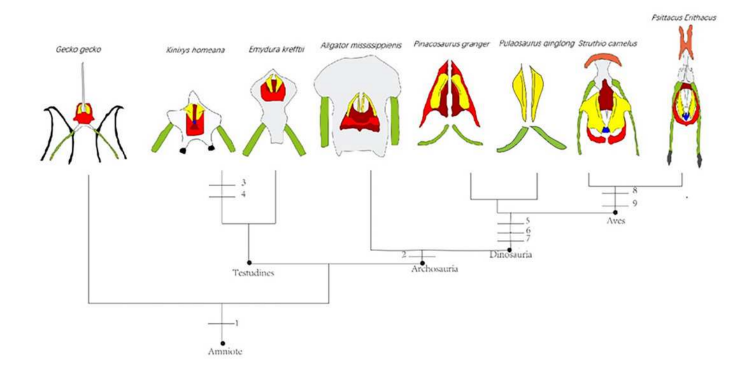




Table 1(on next page)

Measurements of *Pulaosaurus qinglong* specimen V30936 skull elements.

Skull length	82mm
Skull height	42mm
Orbit maximal diameter	30.14mm
Preorbital skull length	30.84mm
Supratemporal fenestra length	15.24mm
Frontal length	36.92mm
Frontal minimal width	14.60mm
Dentary length	39.48mm
Right arytenoid length	31.9mm
Right arytenoid maximal width	5.8mm
Left arytenoid length	34.3mm
Left arytenoid maximal width	6.2mm
Right ceratobranchial length(visible	13.8mm
part)	
Left ceratobranchial length (visible part)	10.0mm

Table 1 Measurements of *Pulaosaurus qinglong* specimen V30936 skull elements



Table 2(on next page)

Measurements of *Pulaosaurus qinglong* specimen V30936 coracoids.



Right coracoid length in medial view	19.86mm
Right coracoid maximal width in medial view	16.68mm
Left coracoid length in lateral view	18.52mm
Left coracoid maximal width in lateral view	?



Table 3(on next page)

Measurements of *Pulaosaurus qinglong* specimen V30936 cervical series.



Axial length	8.50mm
3 rd cervical centrum length	9.04mm
4 th cervical centrum length	12.76mm
5 th cervical centrum length	?
6 th cervical centrum length	15.62mm
7 th cervical centrum length	?
8 th cervical centrum length	?
9th cervical centrum length	?



Table 4(on next page)

Measurements of *Pulaosaurus qinglong* V30936 forelimbs.



50.16mm
10.86mm
?
6.04mm
40.86mm
7.30mm
10.12mm
4.92mm
39.24mm
6.76mm
6.24mm
3.40mm
43.24mm
5.4mm
9.24mm
4.04mm
41.8mm
6.7mm
5.48mm



Right radius shaft minimal width in posterior view	3.58mm
Left manual portion length	32.5mm
Longest left manual digit(2 nd digit) length	29.76mm
Left metacarpal I length	7.62mm
Left metacarpal II length	13.08mm
Left metacarpal III length	12.19mm
Left metacarpal IV length	8mm
Left metacarpal V length	5.36mm



Table 5(on next page)

Measurements of *Pulaosaurus qinglong* specimen V30936 pelvic girdle.



Total length of the ilium	69.84mm
Depth of the blade above the acetabulum	18.88mm
The width between the rostral margin of the pubic peduncle and the caudal margin of the ischial peduncle at the base	23.62mm
Preacetabular process length	28.08mm
Postacetabular process length	21.90mm
Postacetabular process depth	21.16mm



Table 6(on next page)

Measurements of *Pulaosaurus qinglong* V30936 femora, tibiae and fibula.



Left femur length in medial view	88.52mm
D : 1 1 :11 cl c c	14.04
Proximal end width of left femur in medial	14.04mm
view	14.60
Distal end width of left femur in medial	14.68mm
view	11.00
Shaft minimal width of left femur in	11.80mm
medial view	00.04
Right femur length in medial view	80.94mm
Proximal end width of right femur in	?
medial view	·
Distal end width of right femur in medial	16.24mm
view	
Shaft minimal width of right femur in	?
medial view	
Left tibia length in anterior view	97.84mm
Proximal end width of left tibia in anterior	20.02mm
view	
Distal end width of left tibia in anterior	?
view	
Shaft minimal width of left tibia in	10.94mm
anterior view	
Right tibia length in medial view	98.76mm
Proximal end width of right tibia in medial	20.44mm
view	
Distal end width of right tibia in medial	16.98mm
view	
Shaft minimal width of right tibia in	9mm
medial view	
Left fibula length in anterior view	94.90mm
Proximal end width of left fibula in	2.68mm
anterior view	
Distal end width of left fibula in anterior	7.3mm
view	



Shaft minimal	width	of le	ft	fibula	in	1mm
anterior view						



Table 7(on next page)

Measurements of the right astragalus, the right calcaneum and the left astragalus



The length of right astragalus ventral margin	18.42 mm
The length of right calcaneum ventral	6.40 mm
margin	1.50
The height of the ascending process on the anterior side of the right astragalus	4.58 mm
The length of left astragalus ventral	11.10 mm
side's anterior margin	
The length of left astragalus ventral	11.83 mm
side's posterior margin	
The height of the ascending process on	4.10 mm
the anterior side of the left astragalus	
The anteroposterior width of the left	9.16 mm
atragalus	



Table 8(on next page)

Measurements of *Pulaosaurus qinglong* V30936 metatarsals and pedal digits.



Left/Right	digit	perspective	Length/mm	Proximal end width/mm	Distal end width/ mm
Right	Metatarsal I	anterior	27.68	?	5.64
Right	I-1	anterior	16.23	5.36	3.61
Right	I-2	anterior	10.06	4.66	/
Right	Metatarsal II	anterior	47.10	6.58	4.86
Right	II-1	anterior	16.82	7.44	4.96
Right	II-2	lateral	14.5	6.46	4.7
Right	II-3	lateral	12.9	5.28	/
Right	Metatarsal III	anterior	53.86	5.6	11.6
Right	III-1	posterior	15.68	11.42	8.04
Right	III-2	posterior	12.28	8	6.78
Right	III-3	posterior	12.34	8.12	7.22
Right	III-4	posterior	15.66	6.22	/
Right	Metatarsal IV	anterior	?	6.42	?
Right	IV-1	anterior	?	?	5.48
Right	IV-2	lateral	8.86	6.44	4.22
Right	IV-3	lateral	7.96	5.6	5.02
Right	IV-4	lateral	6.6	5.16	3.74
Right	IV-5	lateral	11	4.32	/
Left	Metatarsal II	posterior	47.28	8.32	7.42
Left	II-1	posterior	16.46	5.72	5.32
Left	II-2	anterior	?	8.2	?
Left	II-3	anterior	12.8	3.22	/
Left	Metatarsal III	posterior	54.36	6.92	9.52
Left	III-1	anterior	18.06	8.32	7.62
Left	III-2	anterior	14.71	8.28	7.36
Left	III-3	lateral	12.72	6.5	5.5
Left	III-4	lateral	16.36	5.41	/
Left	Metatarsal IV	posterior	66.04	8.52	5.56
Left	IV-1	anterior	11.53	7.52	6.08
Left	IV-2	anterior	8.9	5.53	5.38
Left	IV-3	anterior	8.6	4.92	4.26





Left	IV-4	lateral	7.32	4.48	4.28
Left	IV-5	lateral	12.04	5.90	/