

A new plesiacerathere (*Perissodactyla*, *Rhinocerotidae*) from the late Early Miocene of northern China (#87877)

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A new plesiacerathere (Perissodactyla, Rhinocerotidae) from the late Early Miocene of northern China

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Aceratheriinae is a group of rhinoceroses widely distributed in Eurasia and North America during the Miocene. As a member of Aceratheriinae, the genus *Plesiaceratherium* in Europe is widely distributed and highly diverse, but only one species of *Plesiaceratherium* (i.e. *P. gracile*) exists in China with a discontinuous distribution range. Recently, we have discovered new materials of *Plesiaceratherium* in the lower layers of the Zhang'enbao Formation exposed in Miaolerling in Tongxin County, China. The new materials are well-preserved with a series of diagnostic features: the long and generally flat skull, with a narrow distance between the sides of the parietal crests; the deep nasal notch positioned at the level of P4; the high supraorbital margin, with its anterior margin positioned at the level of the M1/M2 boundary; the short distance between the posterior edge of the nasal notch and the orbit; the ventrally closed pseudomeatus external auditory; the medium-sized upper incisor I1, with an oval abraded surface; the semi-molarized upper premolars with the protocone and hypocone joined by a lingual bridge; the strong constrictions of protocone on the upper molars; the absent buccal cingulum on upper cheek teeth. Based on the aforementioned combination of characteristics and phylogenetic analysis, we herein establish the new species as *Plesiaceratherium tongxinense* sp. nov. which survived in the late Early Miocene corresponding to the European Land Mammal Age MN 5. Phylogenetic analysis reveals that *P. tongxinense* sp. nov. is located in the basal position of the genus *Plesiaceratherium*, not only providing more detailed morphological characteristics of the plesiaceratheres but also serving as a linkage to the discontinuous distribution of the genus *Plesiaceratherium*.

A new plesiacerathere (*Perissodactyla*, *Rhinocerotidae*) from the late Early Miocene of northern China

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Abstract

Aceratheriinae is a group of rhinoceroses widely distributed in Eurasia and North America during the Miocene. As a member of Aceratheriinae, the genus *Plesiaceratherium* in Europe is widely distributed and highly diverse, but only one species of *Plesiaceratherium* (i.e. *P. gracile*) exists in China with a discontinuous distribution range. Recently, we have discovered new materials of *Plesiaceratherium* in the lower layers of the Zhang'enbao Formation exposed in Miaolerling in Tongxin County, China. The new materials are well-preserved with a series of diagnostic features: the long and generally flat skull, with a narrow distance between the sides of the parietal crests; the deep nasal notch positioned at the level of P4; the high supraorbital margin, with its anterior margin positioned at the level of the M1/M2 boundary; the short distance between the posterior edge of the nasal notch and the orbit; the ventrally closed pseudomeatus external auditory; the medium-sized upper incisor I1, with an oval abraded surface; the semi-molarized upper premolars with the protocone and hypocone joined by a lingual bridge; the strong constrictions of protocone on the upper molars; the absent buccal cingulum on upper cheek teeth. Based on the aforementioned combination of characteristics and phylogenetic analysis, we herein establish the new species as *Plesiaceratherium tongxinense* sp. nov. which survived in the late Early Miocene corresponding to the European Land Mammal Age MN 5. Phylogenetic analysis reveals that *P. tongxinense* sp. nov. is located in the basal position of the genus *Plesiaceratherium*, not only providing more detailed morphological characteristics of the plesiaceratheres but also serving as a linkage to the discontinuous distribution of the genus *Plesiaceratherium*.

Keywords: *Plesiaceratherium*; osteology; phylogeny; late Early Miocene; northern China.

Introduction

~~Add your introduction here.~~ Aceratheriinae is a group of rhinoceroses that lived in Eurasia and North America during the Miocene. As a member of Aceratheriinae, the genus *Plesiaceratherium* is well-documented throughout Eurasia (Young, 1937; Yan & Heissig, 1986; Antoine, Bulot & Ginsburg, 2000; Peter, 2002; Antoine et al., 2010). *Plesiaceratherium* is a primitive acerathere rhinoceros with elongated nasals, long but robust limb bones, and a four-toed manus (Yan & Heissig, 1986; Antoine, 2002). Young (1937) established the genus *Plesiaceratherium* based on some isolated teeth and limb bones discovered in the Early Miocene of Shanwang in Linqu, Shandong Province, China, with *P. gracile* serving as the type species. Later, Chen & Wu (1976) described some dental materials from the Miocene of Jiulongkou in Cixian, Hebei Province, China, as belonging to *P. gracile*. And then, more well-preserved materials of *Plesiaceratherium* were discovered from Shanwang, China, including many skeletons, complete skulls, and many teeth and limb bones, giving more detailed characteristics of *Plesiaceratherium* (Yan, 1983; Yan & Heissig, 1986). In addition to the discovery of *Plesiaceratherium* in eastern China, *Plesiaceratherium* also has been found in the Early Miocene of Lunbori, Baingoin County, northern Tibet, China, including a humeral material (Deng et al., 2012).

In Europe, five species have been attributed to the genus *Plesiaceratherium*. Heissig (1972) established the species *Aceratherium fahlbuschi* based on a nearly complete, uncrushed skull and numerous isolated bones, teeth and mandibular fragments (BSP 1959 II) as a holotype discovered in the locality Sandelzhausen in Bavaria, which was later classified as *P. fahlbuschi* by Yan (1983). According to the depressed skull with mandibular, Merier (1895) established the species *Aceratherium platyodon*, afterwards referred to *P. platyodon* by Yan (1983). Antunes & Ginsburg (1983) established the species *P. lumiarensis* based on right maxilla with P1-M3 as holotype from Portugal which was previously identified as *Aceratherium lumiarensis* by Antunes & Ginsburg (1983). Antoine & Becker (2013) referred a species established by Répelin (1917) to *P. aquitanicum*. Becker & Tissier (2020) established a new species *P. balkanicum* based on a left premolars row as holotype from Bugojno Basin, Bosnia-Herzegovina. There was a species from the lower Miocene of Can Julia, Barcelona, Spain, referred to '*P. mirallesi*' by Yan (1983); however, Lu et al. (2016) considered that '*P. mirallesi*' should be eliminated from the genus *Plesiaceratherium*, and the initial genus name *Dromoceratherium* should be revived.

Until now, the genus *Plesiaceratherium* in Europe is widely distributed and highly diverse, but only one species of *Plesiaceratherium* (i.e. *P. gracile*) exists in China with discontinuous distribution range. Fortunately, we have recently discovered new materials of *Plesiaceratherium* in Tongxin County, Ningxia Hui Autonomous Region, China. The Tongxin region, which was impacted by tectonic activity, contains an abundance and continuous deposit of Cenozoic sediments (Wang et al., 2011, 2016). Our new plesiacerathere materials reported here were found in the lower layers of the Zhang'enbao Formation exposed in Miaerling, which dates to the European Land Mammal Age MN5 (Wang et al., 2016). The studied materials allow describing a new species of plesiacerathere, *Plesiaceratherium tongxinense* sp. nov., not only providing more

detailed multiple characters of *Plesiaceratherium*, but also serving as a linkage to the discontinuous distribution of the genus.

Materials & Methods

The fossils for this study are an adult skull and mandibular discovered in Tongxin, Ningxia Hui Autonomous Region, China, stored in the collection of the Institute of Vertebrate Paleontology and Paleoanthropology (IVPP), Chinese Academy of Sciences, Beijing, China. The fossils are described and identified through anatomical descriptions, comparative anatomy as well as biometrical measurements. Rhinocerotid terminology and taxonomy follow Heissig (1972, 1999), Guérin (1980), and Antoine (2002). Anatomical features described follow basically the same sequence as in Antoine (2002), and Antoine *et al.* (2010). The specimens were measured in accordance with the procedures described in Guérin (1980).

The phylogenetic analysis in this paper is performed using a modified data matrix from Antoine (2002, 2003) in order to assess the phylogenetic position of the new specimen (Appendix). The matrix analyzed in the present study contains 282 morphological characters including 52 cranial characters, 10 mandibular characters, 120 dental characters, and 100 postcranial characters. The phylogenetic analysis was performed using a traditional search under TNT (Version 1.1) (Goloboff, Farris & Nixon, 2008), with all multistate characters treated as additive, except for the characters 72, 94, 102, 140, and 187 (non-additive). The current matrix consists of 30 taxa coded at the species level.

Institutional abbreviations

I/i, upper/lower incisor, **M/m**, upper/lower molar, and **P/p**, upper/lower premolar. **L**, length, and **W**, width. **BSP**, Bayerische Staatssammlung für Paläontologie und Geologie, München, Germany; **MNHN**, Muséum National d'Histoire Naturelle, Paris, France; and **IVPP**, Institute of Vertebrate Paleontology and Paleoanthropology, Chinese Academy of Sciences, Beijing, China.

Results

Systematic paleontology

Order **Perissodactyla** Owen, 1848

Family **Rhinocerotidae** Gray, 1821

Subfamily **Aceratheriinae** Dollo, 1885

Tribe **Aceratheriini** Dollo, 1885

Genus ***Plesiaceratherium*** Young, 1937

Type species. *Plesiaceratherium gracile* Young, 1937

Other species. *P. platyodon* (Mermier, 1895), *P. aquitanicum* (Répelin, 1917), *P. fahlbuschi* (Heissig, 1972), *P. lumiarense* (Antunes & Ginsburg, 1983), *P. naricum* (Pilgrim, 1910), *P. balkanicum* Becker & Tissier, 2020, *P. tongxinense* sp. nov.

Revised Diagnosis. Medium to large-sized primitive aceratheriine; limb bones more slender than in other Miocene aceratheriine genera; the nasal bones are elongated and straight, with a deep nasal notch at the level of P4; I1 is medium-sized, i2 is large and slightly curved; the upper cheek

teeth have low crowns; the upper premolars are semi-molarized; the lower premolars are narrow and long, with relatively shallow ectoflexid (Yan & Heissig, 1986; Lu et al., 2016).

Distribution. Early Miocene (MN 1-5), Eurasia.

Plesiaceratherium tongxinense sp. nov.

(Figs 1-4; Tables 1-3)

Holotype. IVPP V23959, a well-preserved and complete skull and mandible (Fig. 1-3)

representing a full adult individual, which are preserved at the Institute of Vertebrate Paleontology and Paleoanthropology, Chinese Academy of Sciences, Beijing, China.

Derivation of name. The specific name, tongxinense, refers to the geographical location of the discovery.

Type locality and horizon. Miaoerling in Shishi Township, Tongxin County, Ningxia Hui Autonomous Region, China; late Early Miocene.

Diagnosis. The skull is long and relatively flat, with a distance between the sides of the parietal crests; the nasal notch is deep and located at the level of P4; the supraorbital margin is high and its anterior margin is located at the level of the M1/M2 boundary; the distance between the posterior edge of the nasal notch and the orbit is short; the pseudomeatus external auditory is closed ventrally; the upper incisor I1 is developed and specialized, medium in size, with an oval abraded surface; the upper cheek teeth are low crowned; the upper premolars are semi-molarized with the protocone and hypocone connected with each other by a lingual bridge; the protocone on the upper molars has developed anterior and posterior constrictions; the buccal cingulum is absent on upper cheek teeth.

Description

Skull. The skull of IVPP V 23559 is complete and well-preserved with the upper cheek teeth moderately worn. The skull was slightly deformed by lateral compression, with the frontal and posterior part of the nasal bones collapsed downward at the middle suture, the basioccipital and basal pterygoid parts narrowed, and the palatal bones deeply sunken while keeping the tooth rows close together.

In the lateral view, the dorsal skull profile is flat and long. The occipital part is slightly raised, and its profile is almost vertical. The occipital condyle is low and small. The posttympanic process is short and fused with the paraoccipital process, and anteriorly touches the postglenoid process. The pseudomeatus external auditory is closed ventrally, and its upper edge is short located in the lower half of the occipital crest. The area between the temporal and occipital crests is depressed. The zygomatic arch is fairly thin (particularly the middle part), the anterior end of which is located at the level of M1 and close to the cheek teeth row, and the posterior end of the dorsal edge has a short process. The temporal condyle that is articulated with the mandible is protruding from the ventral edge of the zygomatic arch. The postglenoid process is laterally flattened. The position of the dorsal margin of the orbit is high, and the anterior margin of the orbit is located at the level of the M1/M2 boundary. The supraorbital edge of the frontal bone has a coarse area but lacks any process or tubercle. The posterior orbital border is formed by the zygomatic bone, and presents a coarse area, without any tubercles. The nasal bone is thin and flat

without lateral apophyses on both sides. The nasal notch has a U-shaped outline, and its posterior edge is at the level of the middle part of P4. The distance between the posterior edge of the nasal notch and the orbit is short, about 67 mm. The infraorbital foramen is located dorsally to the level of P4 and posteriorly to the nasal notch. The premaxillary bones are fairly well-saved with deeply abraded I1s.

In the dorsal view, the parietal crests are not fused to a sagittal crest, and the smallest width between parietal crests is located anterior to the nuchal crest, about 35 mm. The frontals are constricted at the middle of the temporal fossa. The ratio of zygomatic width to frontal width is more than 1.5. The postorbital process is present. The widest position of the dorsal surface is located between the supraorbital processes, about 139 mm. The nasal bone becomes narrow gradually before the orbits (i.e., the nasal base does not have a constriction). The nasal bone is narrow, flat, and long with a nasal suture.

In the ventral view, the skull is long with a length (from premaxilla to occipital condyle) of 584 mm. The ventral and occipital surfaces of the occipital condyle are rounded, without a median ridge. The hypoglossal foramen is laterally positioned, at the basement of the paraoccipital process. The posttympanic process is and fused with the paraoccipital process, and anteriorly touches the postglenoid process. The alar foramen is opened on the lateral wall of the posterior nares, anteroposteriorly at the level of the temporal condyle. The tympanic bulla has been crushed, exposing the inner bones. The temporal condyle is high, and its transverse axis is straight. The posterior margins of pterygoid are nearly vertical. The anterior edge of the posterior nares is V-shaped in outline, at the level of M3. The posterior edge of the lateral wall of the posterior nares with a steep part is continuous, extending to the foramen lacerum anterius that is at the back of the level of the temporal condyle. The medial and lateral edges of the cheek tooth row are nearly straight. The slender and straight premaxillary bones are two separated and faintly paralleled plates with a length of 66.7 mm. I1 is deeply abraded and oval-shaped.

In the posterior view, the occipital face is bell-shaped in outline, and the upper part is narrower than the lower part. The occipital crest is rounded above and gradually inclined laterally. The nuchal tuberosity is developed. The foramen magnum is small, rounded in shape, and its width is 38 mm. The occipital condyles are relatively small, and its lateral margin has a short and steep upper part, and a long and curved lower part. The width between exterior edges of occipital condyles is 97 mm.

Upper teeth. The upper teeth are moderately worn. The upper cheek tooth row is almost in a straight line posteriorly positioned relative to the orbit. The ratio of the length of the upper premolars (P3-4) to the molars (M1-3) is low, about 45%. On the upper cheek teeth, the buccal wall undulates; the expansion of the lingual cusps is developed; the protocone is constricted anteroposteriorly; and the enamel foldings are absent. On the premolars, the lingual cingulum is present and continuous; the medisinus is closed; there is a lingual bridge between the protocone and hypocone; the protocone is constricted, and its lingual margin is curved; and the postfossette is narrow and closed. On the molars, the antecrochet is developed; the crochet is developed but

the crista is absent; the buccal cingulum is absent, but the lingual cingulum is present; and the protocone and hypocone are strongly constricted.

I1 is oval-shaped in the middle size with a longitudinal length of 26.5 mm in the cross-section. DP1 is not saved.

P2 is nearly quadrangular in occlusal view with a relatively flat buccal wall. The parastyle fold and the paracone rib are weak. The protocone and hypocone connect with each other by a lingual bridge. The hypocone is marginally larger than the protocone. The hypocone is at the same level as that of the metacone. The protoloph is as buccally narrow as the metaloph, and joins with the ectoloph. The crochet and crista are connected forming a medifossette. Both the medisinus and the postfossette are closed. The anterior and the posterior cingula are developed. The lingual cingulum is V-shaped at the entrance of the medisinus. The buccal cingulum is absent.

P3 has a weak parastyle fold and paracone rib with a shallowly undulating buccal wall. The hypocone has an anterior constriction. The protocone and hypocone connect with each other by a lingual bridge. The protocone is slightly larger than the hypocone. The crochet and crista are developed. The medisinus is narrow and closed. The postfossette is small in a round shape. The lingual cingulum is developed and continuous and the buccal cingulum is absent.

P4 is similar to P3, but much larger. The hypocone is expanded, with slight anterior constriction. The protocone is slightly smaller than the hypocone. The lingual margin of the protocone is curved. The protoloph is longer than the metaloph. The lingual cingulum is developed and continuous. The buccal cingulum is absent.

M1 has a projecting parastyle with an undulating buccal wall. The strongly constricted protocone has a flat lingual margin, and the hypocone has a strong anterior constriction. The antecrochet is strong and elongates to the entrance of the medisinus. The medisinus leans to the rear which is narrow. The postfossette is round in shape and closed. The anterior cingulum is developed, and the lingual cingulum is reduced, forming a pillar around the entrance of the medisinus. The buccal cingulum is absent.

M2 has a long parastyle and a developed parastyle fold, and a paracone rib with an undulating buccal wall. The protocone has developed anterior and posterior constrictions. The hypocone has a strong anterior constriction. The crochet is well-developed. The antecrochet is strongly developed, and the stout end extends to the entrance of the medisinus. The antecrochet and hypocone are separated. M2 has an open medisinus, an oval-shaped and closed postfossette. M2 has well-developed anterior and cingulum, and the lingual cingulum is reduced, forming a pillar around the entrance of the medisinus. The buccal cingulum is absent.

M3 has a quadrangular outline in occlusal view. It has a short and sharp parastyle, a wide protoloph, and an ectometaloph. The protoloph is transverse on the antero-lingual side. The crochet is well-developed but does not form a medifossette. The protocone has anterior and posterior constrictions. The anterior cingulum is well developed and continuous, and the posterior and lingual cingula are reduced, forming a pillar respectively.

Mandible: The mandible of IVPP V 23559 is well-preserved with the lower cheek teeth moderately worn. The mandible was slightly deformed by extrusion from the right to left

direction. The right i2 is broken, and the left p2 is lost. The horizontal ramus is long and raised. The lower margin is fairly concave under the cheek teeth, with an upturned mandibular symphysis. The length of the mandibular symphysis along median plane is long, about 153 mm. The posterior border of the mandibular symphysis is situated at the level of the p3. The oval mental foramen is small and located in the lower half of the horizontal ramus at the level before p2. The ascending ramus is relatively high with a height of 278 mm at the coronoid process, and 238 mm at the condyloid process. The mandibular condyle is transversely extended with a width of 87 mm, corresponding to the length of the glenoid fossa of the skull. The medial end of the condyloid process is curved posteriorly. The lateral half of the condyle is slightly inclined anteriorly. The mandibular notch between the coronoid and condyloid processes is narrow and deep. The lower part of the coronoid process is wide antero-posteriorly, and the upper part above the condyloid process tapers gradually as it curves posteriorly, with a flat anterior margin and rounded posterior margin. The posterior margin of the ascending ramus is slightly posteriorly inclined. The mandibular angle is rounded forming an obtuse angle. On the lateral surface of the ascending ramus, the masseter fossa is very deep under the coronoid process. The medial surface of the ascending ramus is depressed. The mandibular foramen is very large and situated anteriorly, with its bottom above the alveolar level. The groove behind the mandibular foramen is deep and wide, extending upward.

Lower teeth: The lower teeth are moderately worn. The row of the lower cheek teeth is aligned with the longitudinal axis of the horizontal ramus.

The i2 is medium-sized, dagger-shaped, and extending obliquely forward and upward almost parallel, with a root thicker than the crown. Its transverse section is a round triangle with an interior sharp angle, and the cross section of the root is oval.

The p2 is small in a triangular shape. It has a short and wide protolephid and a shallow ectoflexid. The trigonid basin is small and open, and the talonid basin is rounded and nearly disappeared. The buccal cingulum is developed under the hypolephid but absent under the protolephid.

The p3 is trapezoid in the occlusal view, with a slightly shorter anterior margin than the posterior one. The postero-buccal corner of the protoconid is rounded. The ectoflexid is shallow. The metalophid is robust, much wider than the entolophid. The trigonid basin is small and shallowly V-shaped, and the talonid basin is deeply V-shaped. The lingual cingulum is developed, and the buccal cingulum is developed under the hypolephid but absent under the protolephid.

The p4 is similar to p3 in morphology, but bigger in size: The occlusal surface is nearly rectangular. The postero-buccal corner of the protoconid is more angular than that of p3. The ectoflexid is wider and deeper than that of p3. The trigonid basin is nearly disappeared, and the talonid basin is deeply V-shaped.

The m1 is deeply worn. The occlusal surface is nearly rectangular. The postero-buccal corner of the protoconid is nearly right-angled. The ectoflexid is wide and shallow. The trigonid basin is nearly disappeared, and the talonid basin is deeply V-shaped. The lingual cingulum is reduced, and the buccal cingulum is absent.

The m2 has a rectangular occlusal surface. The postero-buccal corner of the protoconid is right-angled. The ectoflexid is shallowly V-shaped. The metalophid is robust and wider than the entolophid. The trigonid basin is U-shaped and the talonid basin is deeply V-shaped. Both the protolophid and hypolophid are slightly lingually oblique. The buccal cingulum is absent and the lingual cingulum is reduced.

The m3 has a trapezoid occlusal surface, with slightly shorter anterior margin than the posterior one. The postero-buccal corner of the protoconid is right-angled. The ectoflexid is wide and shallow. The metalophid is robust and wider than the entolophid. The trigonid basin is nearly disappeared, and the talonid basin is deeply V-shaped. Both the protolophid and hypolophid are slightly lingually oblique. The buccal cingulum is absent and the lingual cingulum is reduced.

Comparison and Discussion

The well-preserved new materials (IVPP V23959) from Tongxin, Ningxia have typical features easily recognizable as typical of aceratheriine, including a flat and long nasal bone with a retracted nasal notch; the posttympanic process fused with the paraoccipital process; the upturned mandibular symphysis with large and straight i2s; the constricted lingual cusps on the upper cheek teeth.

Compared with the aceratheriines, the Tongxin specimen differs in the morphology of the skull and mandibular, as well as the degree of specialization of the incisors and cheek teeth. The Tongxin specimen differs from *Mesaceratherium* living in Eurasia from the upper Oligocene to lower Miocene by the relatively smaller I1s, and more complex occlusal patterns of the upper cheek teeth (Heissig, 1969; Blanchon et al., 2018). The Tongxin specimen also differs from *Alicornops* by the relatively smaller I1s, shorter distance from the nasal notch to the orbit, relatively low nuchal crest above the parietal and frontal surfaces, the reduction of buccal and lingual cingulum of lower molars, and the presence of a medifossette on upper premolars and a longer crochet on upper molars (Cerdeño & Sanchez, 2000; Deng, 2004; Heissig, 2012). The Tongxin specimen with developed I1s is clearly different from the derived Eurasian aceratheriines, such as *Hoploaceratherium*, *Chilotherium*, *Acerorhinus*, *Subchilotherium*, and *Shansirhinus* (Borissiak, 1915; Ringström, 1924; Colbert, 1935; Deng, 2005; Heissig, 2012). However, the Tongxin specimen shares diagnostic characters with the genus *Plesiaceratherium*, such as the narrow and slightly raised nuchal crest; the posttympanic process anteriorly touches the postglenoid process; the ventrally closed pseudomeatus external auditory; the nasal notch retracted to the level of P4; the upturned mandibular symphysis with large and straight i2s; the medium-sized I1s; the constricted lingual cusps on the upper cheek teeth; the narrow and long lower premolars with relatively shallow ectoflexid. Therefore, based on the combination of characters, we refer the Tongxin specimen to the genus *Plesiaceratherium* (Young, 1937; Yan, 1983; Yan & Heissig, 1986).

Compared with *P. gracile* (Young, 1937; Lu et al., 2016), the Tongxin specimen differs by the relatively wide parietal crests, but those of *P. gracile* are fused to form a single sagittal crest. The anterior margin of the orbit of the Tongxin specimen is retracted at the level of the M1/M2

boundary, while that of *P. gracile* is located at the level of M1. The constrictions of the protocone and hypocone on the upper molars of the Tongxin specimen are stronger than those of *P. gracile*. The antecrochet and crochet on the upper molars of the Tongxin specimen are developed and stout, whereas *P. gracile* has slightly developed antecrochet and slender crochet on the upper molars. The crista on the upper molars of the Tongxin specimen is absent, but that of *P. gracile* is present. The cheek teeth of the Tongxin specimen are covered by cement on the buccal walls different from that of *P. gracile* without cement.

Different from *P. fahlbuschi* (Heissig, 1972), the Tongxin specimen is larger in size than *P. fahlbuschi*. The distance between the posterior edge of the nasal notch and the orbit the Tongxin specimen is longer than that of *P. fahlbuschi*, which respectively are about 67 mm and 43-35 mm. The parietal crests of the Tongxin specimen are relatively wide, but those of *P. fahlbuschi* are fused to form a single sagittal crest. The anterior margin of the orbit of the Tongxin specimen is retracted at the level of the M1/M2 boundary, and that of *P. fahlbuschi* is located at the level of middle part of M1. The Tongxin specimen has semi-molarized upper premolars with the protocone and hypocone connected with each other by a lingual bridge, while *P. fahlbuschi* has molarized upper premolars with the protocone and hypocone separated.

Compared with *P. platyodon*, the Tongxin specimen is larger in size than *P. platyodon*. The distance between the posterior edge of the nasal notch and the orbit the Tongxin specimen is about 67 mm longer than that of *P. platyodon* about 58mm. The anterior margin of the orbit of the Tongxin specimen is retracted at the level of the M1/M2 boundary, and that of *P. platyodon* is located at the level of middle part of M1. The crochet on the upper molars of the Tongxin specimen is developed and stout, whereas that of *P. platyodon* is weak or absent. The M3 of the Tongxin specimen has a quadrangular outline in occlusal view, while that of *P. platyodon* has a triangular outline.

In addition, there are other species within the genus *Plesiaceratherium* and the preserved materials are scarce. Compared with *P. lumiarense*, the Tongxin specimen has semi-molarized upper premolars with the protocone and hypocone connected with each other by a lingual bridge, while *P. lumiarense* has upper premolars with the protocone and hypocone mostly separated.

The Tongxin specimen differs from *P. aquitanicum* by the features as follows. The protoloph of the Tongxin specimen joins with the ectoloph on P2 but that of *P. aquitanicum* is separated from the ectoloph. The crista on the upper molars of the Tongxin specimen is absent, but that of *P. aquitanicum* is present. The M3 of the Tongxin specimen has a quadrangular outline in occlusal view, while that of *P. aquitanicum* has a triangular outline. Different from *P. balkanicum*, the Tongxin specimen has semi-molarized upper premolars with the protocone and hypocone connected with each other by a lingual bridge, while *P. balkanicum* has molarized upper premolars with the protocone and hypocone separated. The protoloph of the Tongxin specimen joins with the ectoloph on P2 but that of *P. balkanicum* is separated from the ectoloph.

Therefore, the Tongxin specimen is obviously distinguished from all known species of the genus *Plesiaceratherium* by a unique combination of characters: the skull is long and relatively flat with separated parietal crests; the supraorbital margin is high and its anterior margin is located at

the level of the M1/M2 boundary; the upper incisor I1 is developed and specialized, medium in size, with an oval abraded surface; the upper premolars are semi-molarized with the protocone and hypocone connected with each other by a lingual bridge; the protocone on the upper molars has developed anterior and posterior constrictions; the buccal cingulum is absent on upper cheek teeth; the M3 has a quadrangular outline in occlusal view. Based on those, we attribute the Tongxin specimen to a new species, *P. tongxinense* sp. nov.

Although retaining some primitive characters, *Plesiaceratherium* is already a rather specialized genus, such as the complex occlusal surface of the upper cheek teeth, the rather deep nasal incision, and the ventrally closed pseudomeatus external auditory, representing an earlier specialized taxon within Aceratheriinae (Yan & Heissig, 1986). As far as the dentition is concerned, *Aceratherium* and *Plesiaceratherium* are almost indistinguishable, and the skull characters are also fairly similar (Yan, 1983). Therefore, the study of postcranials will be necessary to further understand the relationship between *Aceratherium* and *Plesiaceratherium* and to establish the phylogenetic position of *Plesiaceratherium*.

The genus *Plesiaceratherium* is widely distributed in Eurasia with various occurrences in China, South Asia, and Europe (Young, 1937; Yan & Heissig, 1986; Antoine, Bulot & Ginsburg, 2000; Peter, 2002; Antoine et al., 2010). The earliest representative of this genus is *P. naricum*, from the earliest Miocene of Pakistan (MN1-MN2) (Antoine et al., 2010, 2013; Antoine & Becker, 2013). In Europe, *Plesiaceratherium* was previously discovered in six localities, Sandelzhausen and Voggersberg in Germany, Pont du Manne as well as Estrepouy in France, Charneca de Lumiar in Portugal, and Can Julia in Spain (Heissig 1999; Antoine & Becker 2013). In China, *Plesiaceratherium* was found in three localities, namely, Shanwang in Linqiu, Shandong Province (Young 1937; Yan & Heissig 1986), Jiulongkou in Cixian, Hebei Province (Chen & Wu, 1976), and Lunbori in Baingoin, northern Tibet (Deng et al., 2012). According to the published data, the age of the Shanwang Fauna was about 18 Ma (Deng, Wang & Yue, 2003; Deng et al., 2012), and the fossil locality of the Jiulongkou Fauna should be the latest Shanwangian Age at about 16 Ma (Deng et al., 2012). In addition, the upper part of the Dingqing Formation at the Lunbori locality bearing *Plesiaceratherium* fossil is characteristic of the Early Miocene (Deng et al., 2012). Moreover, the localities yielding the more progressed *Plesiaceratherium* in Europe belong to MN 4 or 5 of the mammalian ages at 18-15 Ma (Steininger, 1999; Deng et al., 2012). Thus, the localities in *Plesiaceratherium*'s distribution in Eurasia are very close in age, i.e. the late Early Miocene.

The new fossil materials of *Plesiaceratherium* reported here were discovered in the lower part of the Zhang'enbao Formation exposed in Miaoerling, corresponding to the European Land Mammal Age MN5 (Wang et al., 2016). Besides, according to Wang (2021), the magnetostratigraphic analyses currently implied an age MN5 (late Early Miocene) for the locality Miaoerling, which is consistent with the observed rhinoceroses association. Therefore, *P. tongxinense* sp. nov. survived in the late Early Miocene.

Phylogenetic analysis

In order to explore the phylogeny of the Tongxin specimen, we performed a phylogenetic analysis of the Rhinocerotidae (based on the data matrix of Antoine (2002, 2003), with the addition of *Plesiaceratherium tongxinense* sp. nov., *P. fahlbuschi*, *P. gracile*, *P. lumiaense*, and *P. platyodon*, resulting in four equally most parsimonious trees. The tree length is 989 steps, with a consistency index of 0.371 and a retention index of 0.602.

The members of Teleoceratina are located in a stable monophyletic clade (Node A) (Fig. 4). They share twenty-four equivocal synapomorphies including the concave dorsal profile of skull (15^{0-1}), the distant processus posttympanicus and processus paraoccipitalis (46^{0-1}), the subtriangular foramen magnum (49^{0-1}), the constricted metaconid on lower cheekteeth (144^{0-1}), the usually absent d1/p1 (in adults) (151^{1-2}), the usually open posterior valley on d2 (180^{0-1}), the usually fused proximal ulna-facets of radius (199^{0-2}), the weak gutter for the m. extensor carpi of radius (202^{0-1}), the closed angle between diaphysis and olecranon of ulna (205^{0-1}), the present anterior tubercle on the distal end of ulna (206^{0-1}), the short posterior tuberosity of magnum (220^{1-0}), the straight magnum-facet of McII (224^{0-1}), the invisible magnum-facet in anterior view of McIII (229^{0-1}), the vestigial McV (231^{0-1}), the salient insertion of the m. extensor carpalis of metacarpals (232^{0-1}), the straight proximal border of the patellar trochlea of femur (241^{0-1}), the usually present medio-distal gutter (tendon m. tibialis posterior) of tibia (243^{0-1}), the ratio of transverse diameter/height of astragalus ≥ 1.2 (252^{1-2}), the absent posterior stop on the cuboid-facet of astragalus (257^{0-1}), the nearly flat calcaneus-facet of astragalus (262^{0-1}), the usually present fibula-facet of calcaneus (264^{1-2}), the present cuboid-facet of MtIII (275^{0-1}), the robust limbs (279^{0-1}), the present postero-distal tubercle on the diaphysis of central metapodials (281^{0-1}). The clade (Node B) includes the plesiaceratheres (Fig. 4) supporting by twelve equivocal synapomorphies including the very long nasal bones (26^{0-2}), the absent i1 (76^{0-1}), the always simple crochet on P2-4 (85^{1-0}), the postfossette on P2-4 with posterior wall (89^{0-2}), the protoloph on P2 joined to the ectoloph (99^{1-0}), the usually absent constriction of the protocone on P3-4 (101^{0-1}), the always present crista on P3 ($105^{1.2-3}$), the absent metacone fold on M1-2 (119^{0-1}), the low and reduced posterior cingulum on M1-2 (124^{0-1}), the always present constriction of the protocone on M3 (135^{0-2}), the present vertical external rugosities on p2-3 (139^{0-1}), and the reduced labial cingulum on lower premolars (150^{0-1}). The new species *Plesiaceratherium tongxinense* sp. nov. established here is located in the basal position of the genus *Plesiaceratherium* supporting by nineteen equivocal synapomorphies including the close frontoparietal crests (35^{0-1}), the high articular tubercle of squamosal (39^{0-1}), the dihedron processus postglenoidalis of squamosal (42^{1-2}), the absent sagittal crest on the basilar process (44^{1-0}), the inclined backward ramus ($60^{0.1-2}$), the foramen mandibulare above the teeth neck (62^{0-1}), the ratio of compared length of the premolars/molars rows between 42% to 50% (63^{0-1}), the present cement on cheekteeth (65^{0-1}), the divergent orientation of i2 (80^{0-1}), the present metaloph constriction on P2-4 (86^{0-1}), the always present antecrochet on P2-3 (90^{0-3}), the protocone on P2 less strong than the hypocone (97^{0-1}), the always present medifossette on P3-4 (100^{0-3}), the always present constriction of the protocone on P3-4 (101^{1-3}), the the lingual bridge existing between protocone and hypocone on P3-4 (102^{2-1}), the strong constriction of the

protocone on M1-2 (116^{0-1}), the continuous metaloph on M2 (129^{1-0}), the external groove developed until the neck on lower cheekteeth (141^{0-1}), the rounded trigonid on lower cheekteeth (142^{0-1}). The scarcity of fossil material of *P. tongxinense* sp. nov. suggests that it may be at an early stage, while the later flourishing of *P. gracile* and the diversity of European *Plesiaceratherium* indicate that the genus has reached an adaptive radiation stage. The clade (Node C) containing elasmotheres (Fig. 4) is supported by thirty-two equivocal synapomorphies including the ratio of compared length of the premolars/molars rows between 42% to 50% (63^{0-1}), the always simple crochet on P2-4 (85^{1-0}), the always absent lingual cingulum on P2-4 (87^{0-3}), the wide postfossette on P2-4 (89^{0-1}), the lingual wall existing between protocone and hypocone on P2 (94^{1-3}), the transverse metaloph on P2 (95^{0-1}), the lingual wall existing between protocone and hypocone on P3-4 (102^{2-3}), the usually present lingual cingulum on upper molars (114^{0-1}), the always present constriction of the protocone on M3 (135^{0-2}), the usually absent on lingual cingulum lower premolars (147^{0-2}), the absent labial cingulum on lower premolars (149^{0-1}), the usually present lingual cingulum on lower molars (157^{0-1}), the deep latero-distal gutter (tendon peroneus muscles) of fibula (250^{0-1}), the median position of the latero-distal gutter of fibula (251^{0-1}), the pad-shaped and continuous postero-proximal tuberosity of MtIV (277^{0-1}). *Diceratherium armatum* and *Menoceras arikareense* are the sister-groups of Elasmotheriina (i.e. elasmotheriines sensu stricto) which is consistent with the results of Antoine's (2003) phylogenetic analysis.

Conclusions

The particularly complete skull and mandible from (IVPP V23959) from Tongxin, Ningxia described here does not quite match any known aceratheres. Although the skull and mandible conform to the generic characters of *Plesiaceratherium*, it also differs from the known species of this genus. The Tongxin specimen is well-preserved with a series of identification features. The skull is long and generally flat, with a distance between the sides of the parietal crests; the nasal notch is deep positioned at the level of P4; the supraorbital margin is high, with its anterior margin positioned at the level of the M1/M2 boundary; there is a short distance between the posterior edge of the nasal notch and the orbit; the pseudomeatus external auditory is closed ventrally. The upper incisor I1 is developed and specialized, medium in size, with an oval abraded surface; the upper cheek teeth have low crowns; the upper premolars are semi-molarized with the protocone and hypocone joined by a lingual bridge; the protocone on the upper molars has developed anterior and posterior constrictions; the buccal cingulum is absent on upper cheek teeth. As a result, based on the aforementioned combination of characteristics and phylogenetic analysis, we herein establish the new species as *Plesiaceratherium tongxinense* sp. nov. surviving in the late Early Miocene corresponding to the European Land Mammal Age MN 5. Phylogenetic analysis reveals that *P. tongxinense* sp. nov. is located in the basal position of the genus *Plesiaceratherium* supporting by nineteen equivocal synapomorphies including the close frontoparietal crests, the high articular tubercle of squamosal, the dihedron processus postglenoidalis of squamosal, the absent sagittal crest on the basilar process, the inclined

backward ramus, the foramen mandibulare above the teeth neck, the ratio of compared length of the premolars/molars rows between 42% to 50%, the present cement on cheekteeth, the divergent orientation of i2, the present metaloph constriction on P2-4, the always present antecrochet on P2-3, the protocone on P2 less strong than the hypocone, the always present medifossette on P3-4, the always present constriction of the protocone on P3-4, the the lingual bridge existing between protocone and hypocone on P3-4, the strong constriction of the protocone on M1-2, the continuous metaloph on M2, the external groove developed until the neck on lower cheekteeth, the rounded trigonid on lower cheekteeth. The discovery of new specimen not only improves the morphological characteristics but also increases the diversity of the plesiaceratheres, serving as a linkage to the discontinuous distribution of the genus *Plesiaceratherium*.

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Figure 1

Photographs and sketches of the skull of *Plesiaceratherium tongxinense* sp. nov., holotype (IVPP V23959) a. dorsal view; b. lateral view; c. ventral view.

~~a. dorsal view; b. lateral view; c. ventral view.~~

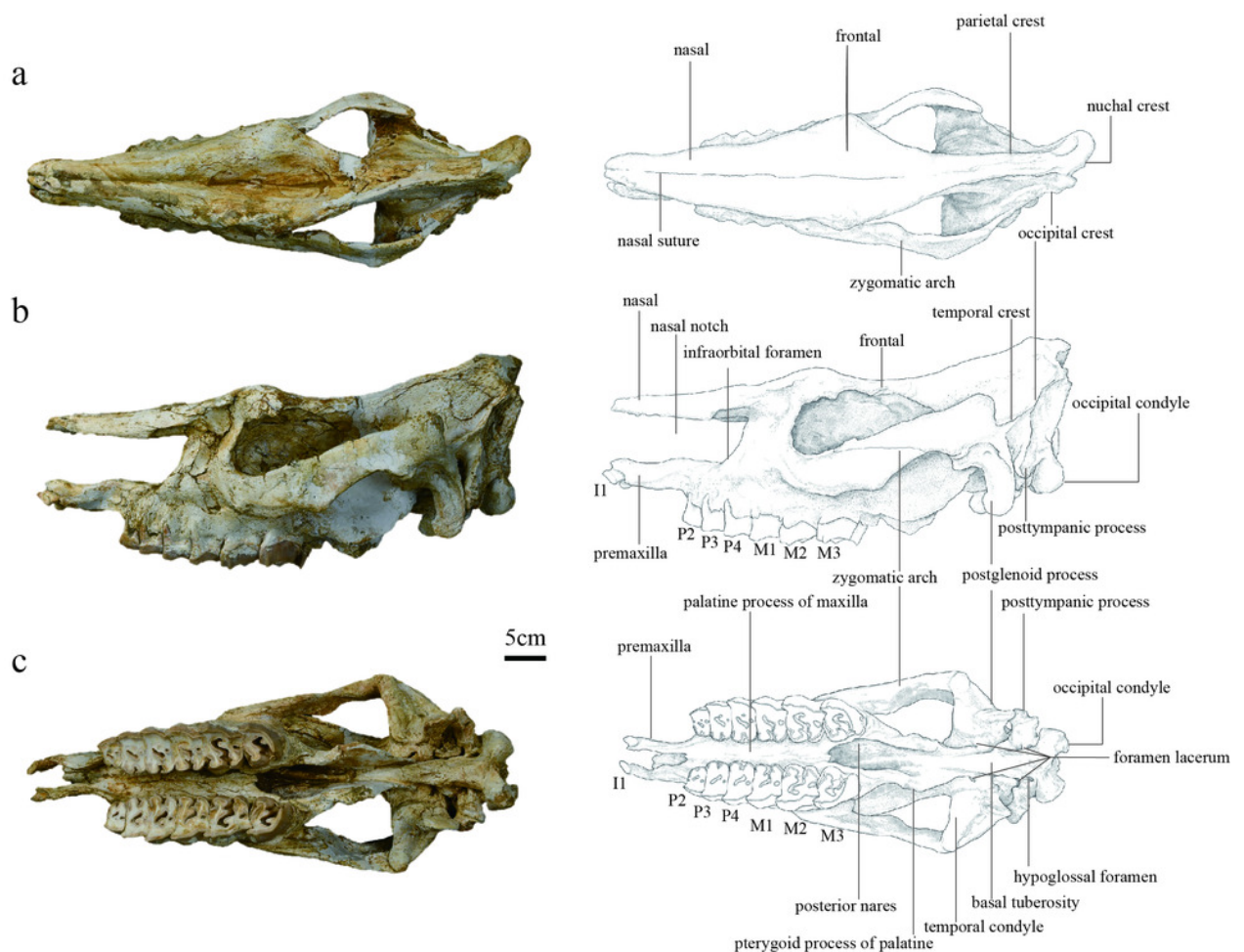


Figure 2

Photograph of the posterior view of the skull of *Plesiaceratherium tongxinense* sp. nov., holotype (IVPP V23959).



5cm

Figure 3

Photographs and sketches of the skull of *Plesiaceratherium tongxinense* sp. nov., holotype (IVPP V23959)

~~a. lateral view; b. occlusal view.~~

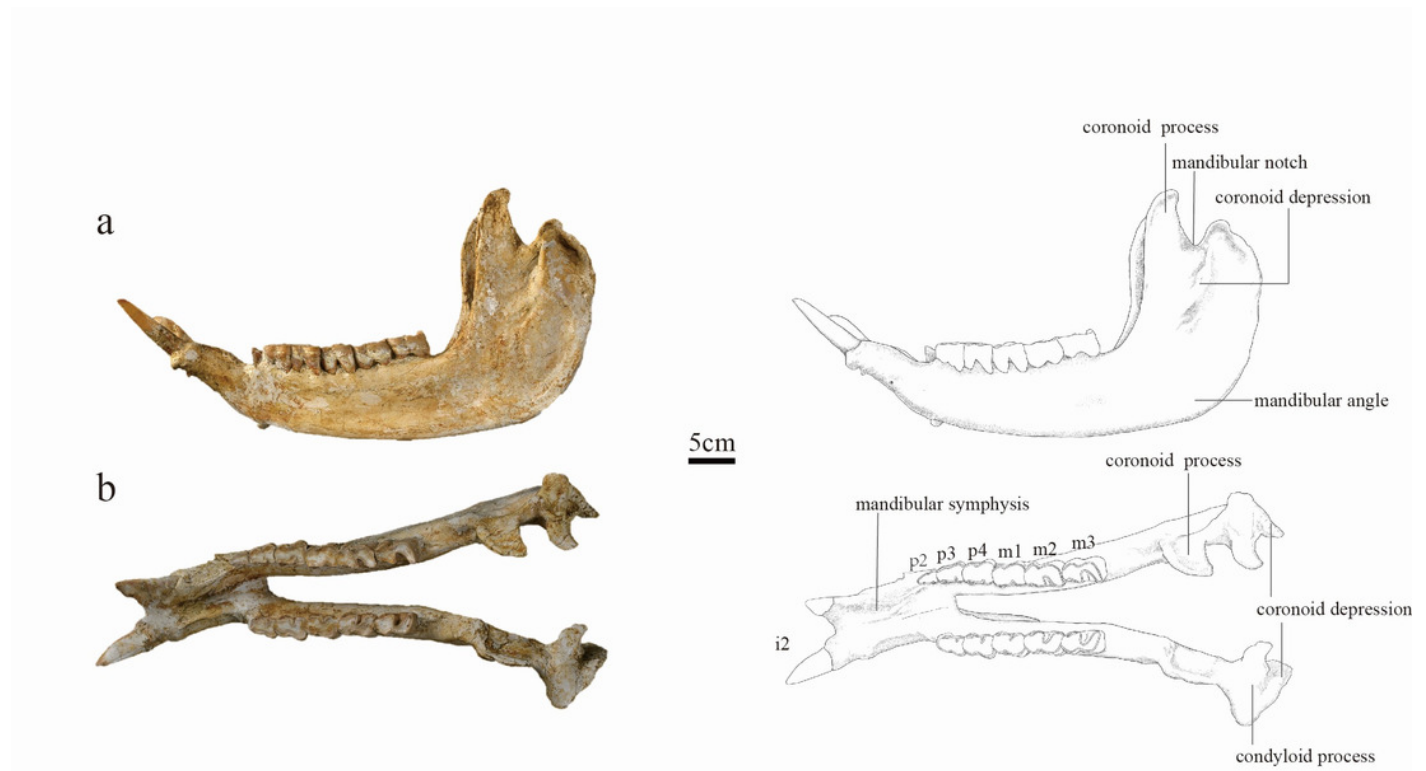


Figure 4

Strict consensus of four most parsimonious trees showing phylogenetic positions of *Plesiaceratherium tongxinense* sp. nov.

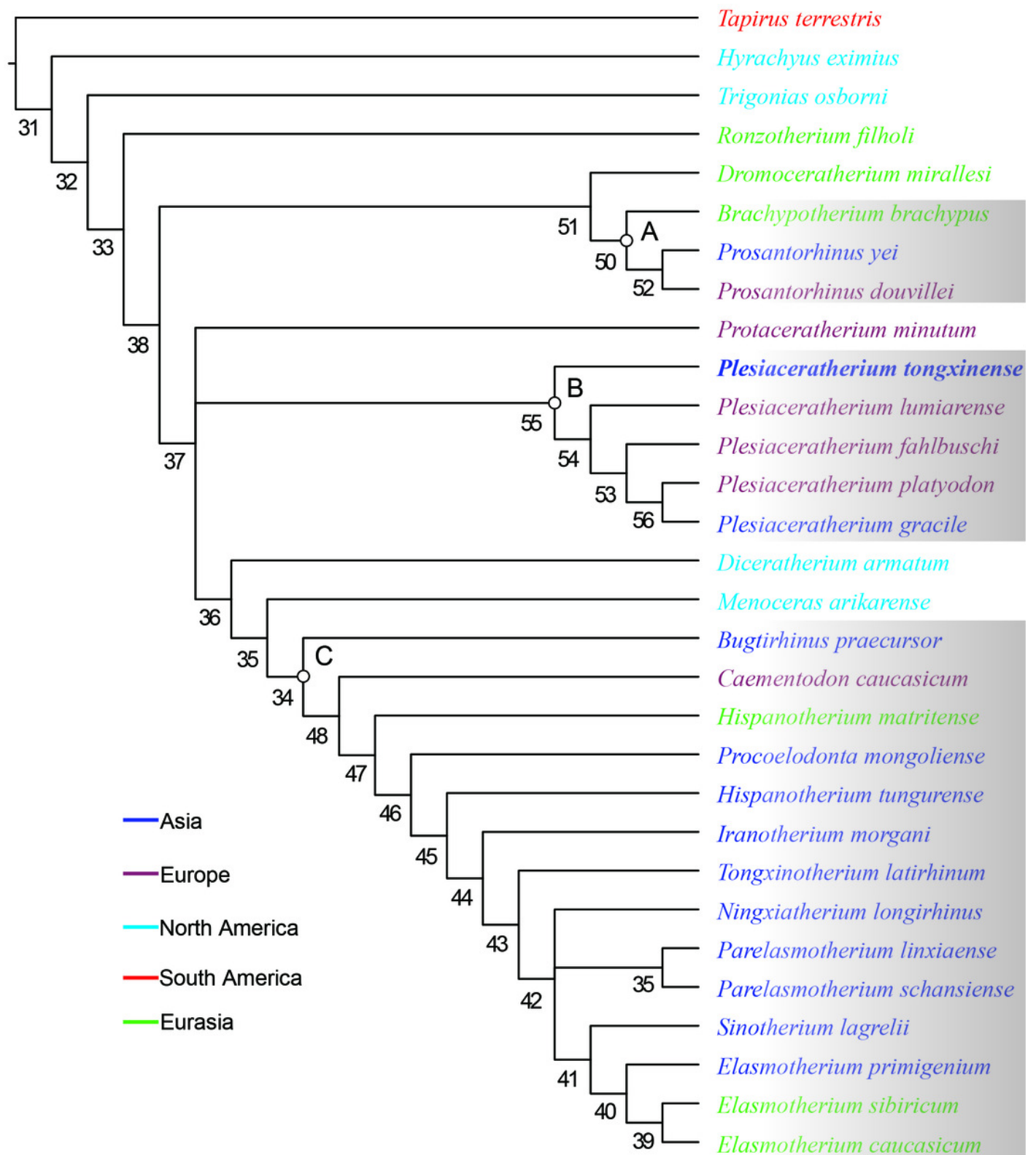


Table 1(on next page)

Measurements of the cheek teeth of *Plesiaceratherium tongxinense* sp. nov. and comparison with other *Plesiaceratherium* species. (mm)

Table 1 Measurements of the cheek teeth of *Plesiaceratherium tongxinense* sp. nov. and comparison with other *Plesiaceratherium* species. (mm)(Measurements of *P. gracile*, *P. fahlbuschi*, *P. platyodon*, and *P. mirallesi* are from Yan and Heissig (1986); *P. lumiaense* is from Antunes and Ginsburg (1983) and Ginsburg and Bulot (1984))

Teeth		<i>P. tongxinense</i> sp. nov.	<i>P. gracile</i>	<i>P. fahlbuschi</i>	<i>P. platyodon</i>	<i>P. mirallesi</i>	<i>P. lumiaense</i>	<i>P. aquitanicum</i>	<i>P. balkanicum</i>
DP1	L	-	-	-	20	23	23	-	21
	W	-	-	-	18	20	19	-	20
P2	L	28	31	24	30	29	29	29	29
	W	36	36	33	35	36	35	36	37
P3	L	34	34	29	28	35	35	34	34
	W	45	42	39	43	45	44	46	43
P4	L	39	37	32	34	38	38	36	35
	W	51	43	43	45	49	49	51	47
M1	L	46	46	32	35	42	45	-	34
	W	53	48	44	43	47	49	-	41
M2	L	54	48	36	38	46	45	47	38
	W	56	50	47	46	51	50	53	45
M3	L	44	43	41	37	42	38	42	33-36
	W	52	45	42	42	45	49	49	37-40
p2	L	23	28	26	27	32	-	-	-

p3	W	16	19	16	17	14	-	-	-
	L	29	31	30	33	35	-	-	-
p4	W	23	20	19	23	21	-	-	-
	L	37	35	32	33	39	-	-	-
m1	W	27	26	22	24	25	-	-	-
	L	38	38	32	35	43	-	-	-
m2	W	28	28	22	24	23	-	-	-
	L	44	40	37	38	46	42	-	36
m3	W	29	26	22	24	24	29	-	24
	L	44	43	-	40	47	43	-	39
	W	27	27	-	23	25	27	-	24

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6 **Table 2** Measurements of the skull of *Plesiaceratherium tongxinense* sp. nov. (IVPP V 23959). (mm) (Measurements of *P. gracile*, *P.*
7 *fahlbuschi*, *P. platyodon*, and *P. mirallesi* are from Yan and Heissig (1986))

numbers	measures	<i>P. tongxinense</i> sp. nov.	<i>P. gracile</i>	<i>P. fahlbuschi</i>	<i>P. platyodon</i>
1	Distance between occipital condyle and premaxilla	584	-	-	-
2	Distance between occipital condyle and nasal tip	598	590-598	516	550

3	Distance between nasal tip and occipital crest	582	-	-	-
4	Distance between nasal tip and bottom of nasal notch	186	184-191	185	215
5	Minimal width of braincase	56	-	-	-
6	Distance between occipital crest and postorbital process	297	-	-	-
7	Distance between occipital crest and supraorbital process	329	-	-	-
8	Distance between occipital crest and lacrimal tubercle	376	-	-	-
9	Distance between nasal notch and orbit	67	66-67	43-35	58
13	Distance between occipital condyle and M3	268			
14	Distance between nasal tip and orbit	243	-	-	-
15	Width of occipital crest	68	-	-	-
16	Width of paramastoid process	134	-	-	-
17	Minimal width between parietal crests	35	-	-	-
18	Width between postorbital processes	124	-	-	-
19	Width between supraorbital processes	139	182-186	175	178
20	Width between lacrimal tubercles	126	-	-	-
21	Maximal width between zygomatic arches	134	258-265	270	205
22	Width of nasal base	87	85-87	90-100	100
23	Height of occipital surface	149	-	-	-
25	Cranial height in front of P2	145	-	-	-
26	Cranial height in front of M1	176	-	-	-

27	Cranial height in front of M3	189	-	-	-
28	Width of palate in front of P2	40	-	-	-
29	Width of palate in front of M1	35	-	-	-
30	Width of palate in front of M3	41	-	-	-
31	Width of foramen magnum	38	-	-	-
32	Width between exterior edges of occipital condyle	97	-	-	-

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numbers	measures	<i>P. tongxinense</i> sp. nov.	<i>P. gracile</i>	<i>P. fahlbuschi</i>	<i>P. platyodon</i>
1	Distance between anterior end of symphysis and mandible angle	470	433-435	435	455
2	Distance between posterior end of symphysis and mandible angle	317	-	-	-
3	Height of horizontal part of ramus at posterior edge of p2	61	-	-	-
4	Height of horizontal part of ramus at posterior edge of p3	65	-	-	-
5	Height of horizontal part of ramus at posterior edge of p4	68	-	-	-
6	Height of horizontal part of ramus at posterior edge of m1	80	-	-	-
7	Height of horizontal part of ramus at posterior edge of m2	89	-	-	-
8	Height of horizontal part of ramus at posterior edge of m3	98	79-88	85-77	85
9	Transverse diameter of lingual edge of horizontal part of ramus at p4-m1	52	-	-	-
10	Transverse diameter of lingual edge of horizontal part of ramus at anterior edge of m3	61	19-21	35	34

11	Length of symphysis along median plane	153	89-96	80-108	96
13	Antero-posterior diameter of vertical part of ramus	131	-	-	-
14	Transverse diameter of condyle	88	-	-	-
15	Height of condyle	238	-	-	-
16	Height of coronoid process	279	-	-	-

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11 **Table 3** Measurements of the mandible of *Plesiaceratherium tongxinense* sp. nov. (IVPP V 23959). (mm) (Measurements of *P.*
 12 *gracile*, *P. fahlbuschi*, *P. platyodon*, and *P. mirallesi* are from Yan and Heissig (1986))

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