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
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Cranial anatomy of the Early Triassic trematosaurine *Angusaurus* (Temnospondyli: Stereospondyli): 3D endocranial insights and phylogenetic implications

Meritxell Fernández-Coll¹ · Thomas Arbez² · Federico Bernardini^{3,4} · Josep Fortuny^{2,5} 

Abstract

Background Trematosaurines are a widespread group of early tetrapods (Temnospondyli, Stereospondyli) known from all continents except South America and Antarctica. They radiated rapidly during the Early Triassic just after the End Permian mass extinction and are of interest to understand the recovery of the ecosystems just after extinction. Trematosaurines disappeared during the Late Triassic.

Objective Herein, a re-description of the genus *Angusaurus* is presented based on a new specimen. This genus is known from the Early Olenekian (Early Triassic) of Russia and comprises four valid species, although the diagnostic characters that define some of them are vague and controversial.

Methods The new specimen described, using MicroCT scanner and 3D digital modeling, sheds light on the anatomical details of the external and inner cranial structure, and provides new details of the neurocranium as well as the ontogeny of this genus.

Results and Discussion A cladistic analysis of trematosaurines (including most trematosauroids) confirms the problematic nature of some *Angusaurus* species and provides a basis for detailed discussion about the phylogeny of trematosaurines.

Keywords Trematosauria · Trematosauroidea · Trematosauridae · Trematosaurinae · Stereospondyli

Resumen

Antecedentes Los trematosaurinos son un grupo ampliamente distribuido de tetrápodos basales (Temnospondyli, Stereospondyli) conocidos de todos los continentes excepto Sud América y la Antártida. Este grupo se diversificó rápidamente durante el Triásico Inferior justo después de la extinción en masa del Pérmico terminal y son de interés para entender la recuperación de los ecosistemas justo después de la extinción. Los trematosaurinos desaparecieron durante el final del Triásico.

Objetivo En el presente trabajo se presenta una re-descripción del género *Angusaurus* basada en un nuevo espécimen. Este género es conocido del Olenekiense Inferior (Triásico Inferior) de Rusia y comprende cuatro especies validas, aunque los caracteres diagnósticos que definen algunos de ellos son vagos y controvertidos.

Métodos El nuevo espécimen descrito, usando un escáner de MicroTomografía y modelado digital 3D, aporta nueva luz sobre detalles craneales anatómicos externos y de la estructura craneana interna y aporta nuevos detalles del neurocráneo así como de la ontogenia de este género.

Resultados y discusión Un análisis cladístico de los trematosaurinos (incluyendo la mayoría de trematosauroides) confirma la problemática naturaleza de las especies de *Angusaurus* y aporta las bases para una profunda discusión sobre la filogenia de los trematosaurinos.

Palabras clave Trematosauria · Trematosauroidea · Trematosauridae · Trematosaurinae · Stereospondyli

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1 Introduction

Temnospondyls are a large group of tetrapods ranging from the Early Carboniferous to the late Early Cretaceous (Schoch 2013). They survived the end-Permian mass extinction and

during the Early Triassic (Induan) their diversity rapidly increased, although it decreased gradually after the Late Triassic (Milner 1990). One of the most successful clades of temnospondyls are stereospondyls. Their members made their first appearance during the Late Permian and survived until the Early Cretaceous, reaching their maximum diversity during the Early Triassic (Stayton and Ruta 2006), presenting a worldwide distribution. They were predominantly, but not exclusively, aquatic–semiaquatic tetrapods living within a range of habitats from freshwater to coastal and even marine environments. During the beginning of the Early Triassic two stereospondyl groups, trematosauroids and capitosauroids, dominated these ecosystems, some acquiring gigantic sizes (Fortuny et al. 2016). Trematosauroids lived from the Early to the Late Triassic (Hellrung 1987), expanded over Pangaea, and are known from all continents except South America and Antarctica. The phylogenetic relationships within trematosauroids are still debated, but they include few basal trematosauroids (such as *Benthosuchus*) and a huge clade of trematosaurids divided into two subclades: Trematosaurinae (short-snouted members) and Lonchorhynchinae (long-snouted members; see Welles 1993; Steyer 2002; Schoch 2006; Fortuny et al. 2018 for further discussion).

Most trematosaurids belong to Trematosaurinae. Several trematosaurine taxa have been described, mainly from Central European basins and the Russian Eastern Platform, but also from other European basins (e.g. Spitsbergen), Africa (e.g. Madagascar, South Africa) and Australia. The radiation of trematosaurines during the Early Triassic was rapid, demonstrating the importance of this group in understanding the recovery of the ecosystems just after the End Permian mass extinction.

Herein, the Early Triassic trematosaurine *Angusaurus* is analyzed and its phylogenetic position addressed. This genus is known from the Russian East Platform and four species have been erected to date based on cranial and mandible specimens (Kuzmin 1935; Getmanov 1982, 1989; Novikov 1990). Based on the description of an undescribed small-sized specimen our aims are: (a) to describe the external anatomy; (b) to describe in detail its neurocranium using MicroCT scanning; (c) the taxonomic study of the analyzed skull to evaluate its taxonomical assessment; and (d) to perform a cladistic analysis of trematosaurids, with particular emphasis on trematosaurines in order to evaluate the phylogenetic position of *Angusaurus* and shed new light on the phylogeny of trematosaurines and its paleobiogeographical implications.

Institutional Abbreviations—MNHN: Museum National d’Histoire Naturelle, Paris, France; PIN: Paleontological Institute, Moscow, Russian Federation; SMNS, Staatliches Museum für Naturkunde in Stuttgart, Germany.

2 Materials and methods

The analyzed specimen (SMNS 81782) is stored at the Staatliches Museum für Naturkunde in Stuttgart, Germany and corresponds to an almost complete skull. It has been reported (not figured nor described) by Schoch and Milner (2000) and referred to *A. succedaneus*. The specimen was recovered in the region of Rybinsk, Russia. This area is well-known geologically and paleontologically, located in the upper part of the Volga River. The sediments consists of lacustrine marls dated as Olenekian, Early Triassic (Grauvogel-Stamm and Ash 2005). The region of the upper Vetluga River, one of the Volga effluents, is well known for the presence of temnospondyl remains: capitosaurids such as *Wetlugasaurus*, trematosauroids such as *Benthosuchus* and trematosaurids as *Thoosuchus*, *Angusaurus* and *Trematotegmen*, have all been found in the Upper Vetluga formations (Sennikov 1996).

The specimen was scanned using X-ray computed microtomography (microCT) at the Multidisciplinary Lab (MLAB) of the “Abdus Salam” International Centre for Theoretical Physics (ICTP) in Trieste, Italy. See Tuniz et al. (2013) for further details about the equipment. Due to the size of the sample and the reduced size of the microCT scan detector, it was required to scan it in two parts: one from the anterior end of the snout to the posterior edge of the prefrontal, and the second one from the posterior edge of the prefrontal to the occipital condyles. The parameters for each part of the skull were the same: 149 kV voltage, 50 μ A current with 0.1 mm filter of copper. A total of 1800 X-ray slices were obtained with a voxel resolution of 35.71 μ m.

Raw data from each scan was imported (as stack of TIFF 8-bit files) to Avizo 7.0 to generate a 3D surface from the microCT images. Data produced by segmentation in Avizo were exported to the programs Rhinoceros 5.0 and 3matic 9.0 for PDF 3-D creation (see supplementary information).

The direct study of the specimen complements the digital part of its analysis. Measurements of some areas of the skull were taken with a digital calliper (Electro DH, model 60.205, resolution of 0.01 mm).

Anatomical abbreviations ans, anterior sulcus; apv, anterior palatal vacuity; br, basiptyergoid ramus; ch, choana; cls, cristae laterosphenoidales; co, crista obliqua; cp, cultriform process; ct, chorda tympani; cv, *columna verticalis*; f, foramen; fm, foramen magnum; is, infraorbital sulcus; iv, interptyergoid vacuity; js, jugal sulcus; jf, jugular foramen; la, lamina ascendens; ld, lamina descendens; fl, flexura lacrimalis; lp, lamellar process; mf, Meckelian foramen; na, naris; oc, occipital condyle; ocs, occipital sulcus; pap, paroccipital process; pf, pineal foramen; pl, *processus lamellosus*; plf, palatoquadrate fissure; pr, palatine ramus; psm, *processus*

submedullaris; pst, *processus subtympanicus*; ptf, posttemporal fenestra; qr, quadrate ramus; sos, supraorbital sulcus; sv, supratemporal vacuity; ts, temporal sulcus. Bones: AN, angular; COR, coronoid; D, dentary; ECPT, ectopterygoid; EO, exoccipital; F, frontal; ICOR, intercoronoid; J, jugal; L, lacrimal; MX, maxilla; N, nasal; P, parietal; PAR, prearticular; PCOR, precoronoid; PL, palatine; PMX, premaxilla; PO, postorbital; POF, postfrontal; PP, postparietal; PRF, prefrontal; PS, parasphenoid; PSP, postsphenial; PT, pterygoid; Q, quadrate; QJ, quadratojugal; SA, surangular; SP, splenial; SQ, squamosal; ST, supratemporal; T, tabular; VO, vomer.

3 Systematic paleontology

Temnospondyli Zittel 1887–1890

Stereospondyli Zittel 1887–1890

Trematosauria Romer, 1947 sensu Yates and Warren 2000

Trematosaurioidea Säve-Söderbergh 1935 sensu Schoch 2013

Trematosauridae Watson 1919

Trematosaurinae Watson 1919, sensu Fortuny et al. 2018

Angusaurus Getmanov 1989

Angusaurus cf. *tsylmensis* Getmanov 1989

Type species.—*Angusaurus dentatus*

Valid species.—*Angusaurus dentatus*, *Angusaurus weidenbaumi*, *Angusaurus succedaneus*, *Angusaurus tsylmensis*.

Definition All taxa sharing a more recent common ancestor with *Angusaurus dentatus* than with *Thoosuchus yakovlevi*.

Emended diagnosis Skull up to 200 mm long. Oval orbits, laterally emplaced. Elongated postorbitals and postfrontals. Interchoanal tooth row strongly reduced. Ventral opening of the orbit in the mid part. Postglenoid process of the mandible elongated. Medial process of supraangular well developed.

Type horizon Vetlugian Horizon with two subhorizons: Sludkinsk Subhorizon, early Early Olenekian (Early Triassic) for *Angusaurus dentatus*, *Angusaurus succedaneus*, *Angusaurus weidenbaumi*. Ustmylskian Subhorizon, late Early Olenekian (Early Triassic) for *Angusaurus tsylmensis*.

Locus typicus Left shore of Samara river at Dolgiy Yar, Buzulukskij Rajon, Orenburgskaya Oblast (*A. dentatus*); Chapayevsk, Alexeyevka District, Kuybyshevskaja Oblast (*A. succedaneus*); Pleas near Kineshma, Furmanovskij Rajon, Ivanovskaya Oblast (*A. weidenbaumi*); Mala locality, Tsył'ma River region, Komi Republic (*A. tsylmensis*). All of them eastern European Russia.

4 Description

4.1 Preservation

The specimen (SMNS 81782) has a narrow, elongated and triangular-shaped skull (Fig. 1). Its maximum length is 112 mm and it has a maximum width of 74 mm

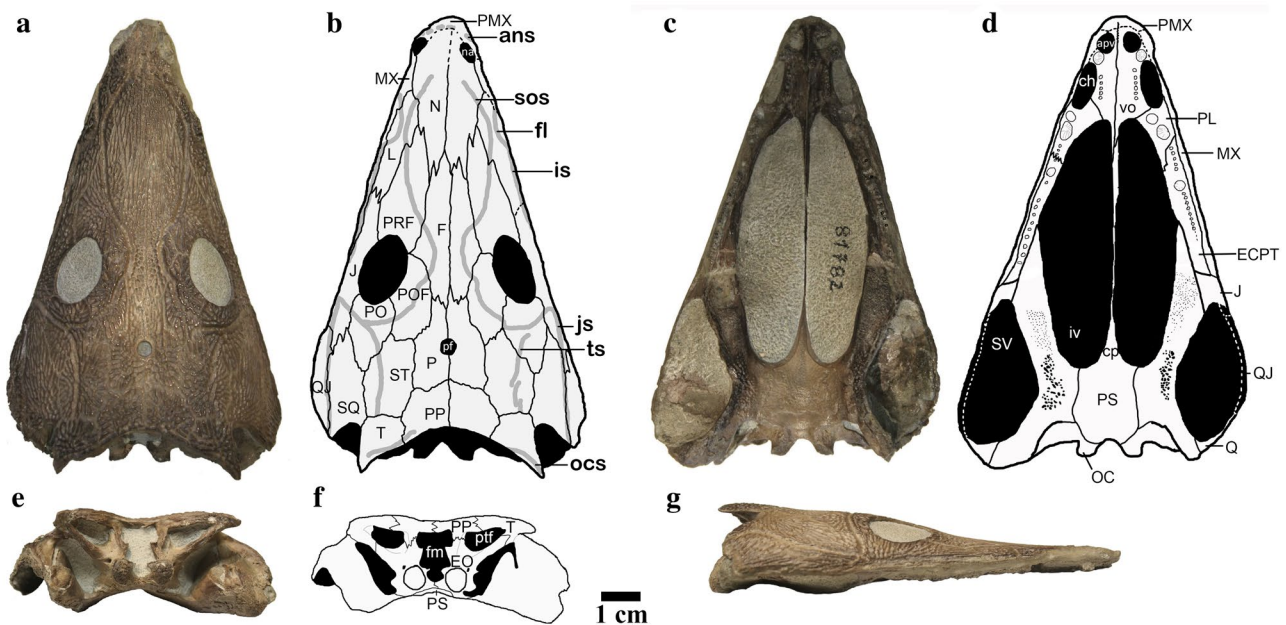
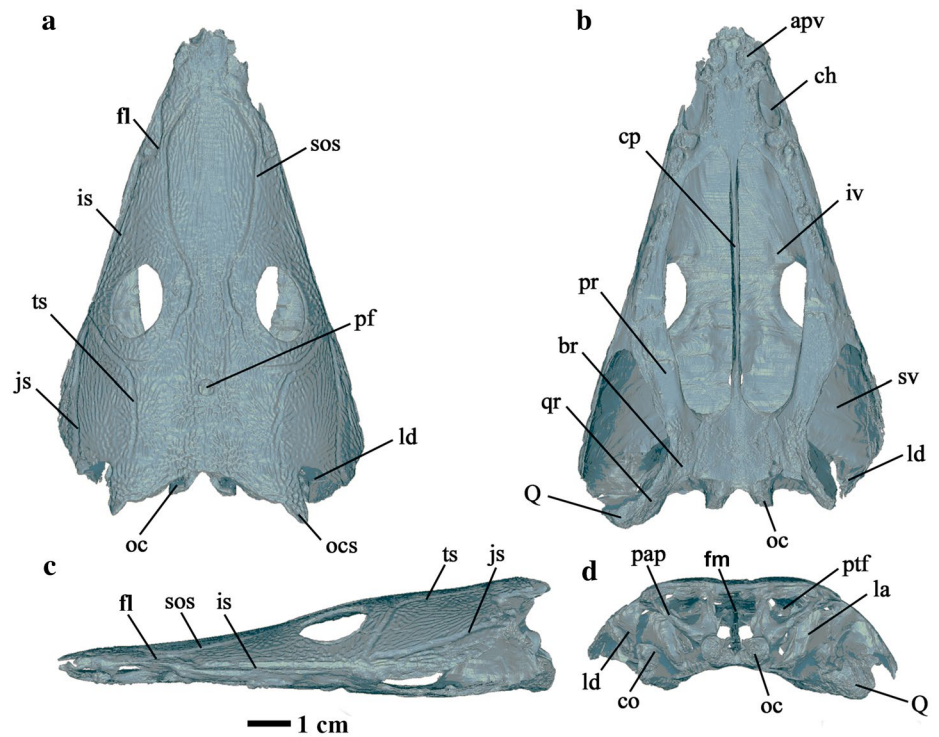


Fig. 1 *Angusaurus* cf. *tsylmensis*. Early Olenekian (Lower Triassic) of Russia (SMNS 81782) in dorsal (a, b), ventral (c, d), occipital (e, f) and lateral view (g)

Fig. 2 3D model of *Angusaurus* cf. *tsylmensis* (SMNS 81782). Matrix removed in dorsal (a), ventral (b), lateral (c) and occipital (d) views



(supplementary material—Table). The specimen is a nearly complete and well-preserved skull (Figs. 1, 2). The anteriormost part of the snout, formed by the maxilla, premaxilla and part of the nasal bone, as well as the cheek region, comprising the jugal and the quadratojugal, is the only slightly damaged part of the specimen that precludes proper observation. There is no evident damage due to mechanical transport in the skull. The posterior margin of the skull, formed by the tabulars and the postparietals, is slightly concave (Figs. 1, 2).

4.2 Lateral line system and ornamentation

Between most of the bones the sutures are clearly visible, while on the anterior part of the snout—as previously mentioned—and on the cheek region they are less well defined due to damage. Also, on some bones (nasal, premaxilla, maxilla) the ornamentation and the sensorial sulci are difficult to discern with confidence. Sensorial sulci are generally continuous and strongly marked (Figs. 1, 2). The anterior sulcus, generally found at the anterior end of the snout, close to the naris, is faintly present. The supraorbital sulcus is continuous with an S-shaped morphology and passes through the nasal, lacrimal, prefrontal, frontal, postfrontal, and postorbital bones, where it connects with the jugal and the temporal sulci, the latter found on the left side (but not present on the right side). The infraorbital sulcus goes mainly through the maxilla, and in the lacrimal bone it forms a flexure, the flexura lacrimalis, that is faintly marked and

rounded. The jugal sulcus is continuous and goes through the jugal, following the quadratojugal–squamosal suture in posterior direction (Fig. 2). The temporal sulcus contacts the jugal sulcus in the jugal and goes through the postorbitals and the supratemporals, contacting the supraorbital sulcus in the postorbital bone (clearly on the left side and almost contacting it on the right side). The occipital sulcus is faintly present on the left side, from the tabular horn up to one-third of the postparietal. On the right side, the occipital sulcus is only visible on the tabular bone. On the right side of the skull roof, part of the supraorbital, temporal and occipital sulci are slightly marked. On this side, the supraorbital sulcus is less evident through the postfrontal while the temporal sulcus is less evident through the supratemporal. On the other hand, the occipital sulcus is less evident on the postparietal bone.

The skull roof has a prominent ornamentation with some areas presenting ridge-groove ornamentation, indicating intensive growth on the skull (Damiani and Yates 2003). The bones on the skull roof are characterized by dermal bone with polygonal sculpture in the ossification centre and radial sculpture in the peripheral region of the bone. The polygonal sculpture consists of polygonal grooves separated by thin ridges, while the radial sculpture is characterized by elongated furrows and ridges that radiate in opposite direction from the ossification centre of the bone (Witzmann et al. 2010). The sculpture is not homogenous along the entire skull roof. Suture morphology was identified thanks to micro-CT images. Most sutures are groove-like (sensu

Kathe 1999), but nonetheless some sutures present on the parietal and supratemporal bones are shelf-like.

4.3 Skull roof

The premaxilla and the maxilla are anteroposteriorly elongated but badly preserved anteriorly, precluding assessment of the morphology of the anteriormost part of the snout. The nasal bones are more elongated than the premaxilla. The premaxilla–nasal suture morphology cannot be discerned due to damage. The nostrils, situated at the tip of the snout, are oval, laterally positioned, and have a length between 8 and 9 mm. This measurement is approximate because the left one is badly preserved. The nostril borders are formed by the premaxilla, the nasal and the maxilla. There is no trace of a septomaxilla, although its presence cannot be completely discarded due to bad preservation. The lacrimal is present, with the supraorbital and infraorbital sulci passing through it. The latter sulcus includes the flexura lacrimalis. The prefrontal connects with the postfrontal, frontal, nasal, lacrimal and jugal bones. The prefrontal is longer than wide, being wider in its posterior part. The jugal is long, forming part of the orbital rim, and representing most of the orbital margin (35%). The rest of bones that form the orbital margin (postorbital, postfrontal, prefrontal, and jugal) occupy a slightly smaller part of the contour, around 20% each (Fig. 1). The frontal is one of the longest bones of the skull roof with a maximum length of 40.1 mm (total length of the skull is 112 mm). It is followed by the nasal, which is approximately 38.3 mm in length. The frontal does not contact the orbital margin, a characteristic found in other stereospondyls.

The orbits are oval and large in size, 19.1 mm long (right) and almost 18 mm long (left), around 16.6% of the total preserved dorsal midline length. The orbits are placed in the posterior half of the skull and are lateralized, located near the edge of the skull. The interorbital distance is 23.3 mm (supplementary material—Table). There is a pronounced depression between the two orbits, formed by part of the postfrontal and frontal bones. The ventral opening of the orbits is placed on the posterior half of the interpterygoid vacuities. The shape of the postorbital bone is raindrop-like, being posteriorly elongated. The supratemporal is rectangular and contacts the tabular, postparietal, parietal, postfrontal, postorbital and squamosal bones. The postfrontal is also rectangular. The quadratojugal is elongate and narrower than its adjacent bones. The quadrate condyles are anterior to the exoccipital condyles. The squamosal is longer than wide and articulates with the supratemporal, postorbital and quadratojugal bones. The parietal is elongate and contacts the postparietal, supratemporal, postfrontal and frontal. The pineal foramen is circular and large, with a diameter of almost 4 mm. It is situated in a posterior position between both parietals, 8.7 mm posterior to the orbits (supplementary

material—Table). The tabular is triangular, with a short, pointed and poorly-developed horn (Figs. 1, 2). The tabular and squamosal bones conform an open and deep otic notch. The postparietal is slightly wider than long.

4.4 Palate

In ventral view, the palatal region of *Angusaurus* is similar to *Thoosuchus* (Damiani and Yates 2003), although some differences deserve special attention (Figs. 1, 2). The premaxilla–vomere suture possibly reaches the anterior margin of the anteropalatal vacuity (or fenestra sensu Schoch and Milner 2000), or ends really close to it. These vacuities (Fig. 1) are rounded and have a diameter of 5 mm (right) and 4.9 mm (left). The vomere–palatine suture is located on the posterior part of the choana. The choanae are oval, both 11.9 mm on the long axis, and placed on the lateral margin of the snout. The maxilla is very elongated and contacts the vomere, palatine, ectopterygoid and jugal bones. The palatine contacts the vomere, ectopterygoid, and maxilla. The edge of the palatine forms part of the interpterygoid vacuity and the choana (Figs. 1, 2). The palatine is adjacent to the ectopterygoid posteriorly. The ectopterygoid bears some dentition, mainly anteriorly and on the central region. The ectopterygoid contacts the maxilla laterally, the pterygoid posteromedially, and the jugal dorsolaterally. The ectopterygoid forms the central part of the margin of the interpterygoid vacuities.

The pterygoid is tri-radiate, with a palatine, basipterygoid, and quadrate rami. The pterygoid has no contact with the exoccipital. From the anterior part of the quadrate ramus to the posterior part of the palatine ramus, the lateral border of the pterygoid bears a strip of polygonal ornamentation. Almost the entire ventral surface of the palatine ramus presents a shagreen field except for its most posterolateral part, as previously mentioned. The palatine ramus forms a long, wide and thick sheet of bone. It is anterolaterally directed, with the anterior extremity contacting the ectopterygoid and the jugal. On the anterior tip of both rami a part of the ventral side is missing, creating an empty space. On the left ramus, a small patch of dot-shaped ornamentation anterior to the above-mentioned empty space indicates the anterior limit of the pterygoid. In anterior direction, the rami become slightly wider until their contact with the jugal. Then, their lateral borders become straight and anteriorly directed. The medial edge of each ramus forms the posterior half of the lateral border of the interpterygoid vacuity. It is slightly curved near the basipterygoid ramus and then become nearly straight anteriorly.

The basipterygoid ramus is very short and laterally wide compared to the other rami. On the medial side it has a complex, tight and dorsoventrally elongated suture with the parasphenoid bone, only visible from the ventral side. A deep and well-delimited subconical depression, the conical recess,

is visible on the dorsal portion of this branch. This depression is laterally delimited by the anterior base of the lamina ascendens and posteriorly by the crista parapterygoidea of the parasphenoid. This is a similar condition to that found in *Lyrocephaliscus euri* (Mazin and Janvier 1983 and personal observations of T.A. on MNHN F SVT 520), both in terms of shape and location on the basiptyergoid ramus.

On the ventral surface of each basiptyergoid ramus, a pair of grooves is visible, with an anterior one reaching the interptyergoid fenestra and a posterior one reaching the posterior border of the basiptyergoid ramus (Fig. 1, 2). These grooves are nearly straight, with an antero-posterior orientation, and being between 3 and 8 mm wide. In addition, based on microCT data, the anterior one is connected with the posterior one by a canal inside the bone. Similar grooves are found in several species of temnospondyls, running through the whole length of the basiptyergoid rami (e.g. *Lyrocephaliscus euri*, Mazin and Janvier 1983; *Thoosuchus yakovlevi*; personal observations of T.A. on specimen SMNS 58880; *Trematolestes hagdorni*, Schoch 2006).

The quadrate ramus of the pterygoid forms a bony blade, about two times shorter than the palatine ramus. It shares a tight suture with the quadrate (but the left quadrate is missing) and it is dorsally in contact with the lamina ascendens. The quadrate ramus is slightly inclined medially and transversally oriented. The lateral edge of this ramus is concave and forms the posteromedial part of the subtemporal vacuity.

The lamina ascendens of the pterygoid is a slender, subvertical bony lamella, slightly thicker anteriorly. It medially delimits the columellar cavity in extending the lamina descendens of the squamosal, but these two laminae remain separated by a space, the palatoquadrate fissure. This structure lies on the quadrate and the basiptyergoid rami with a transversal orientation except for its anterior tip, which points medially. At the level where the lamina ascendens contacts the quadrate ramus, a crest is present on the medial side, called ‘crista obliqua’ by Bystrow and Efremov (1940: fig. 5). It delimits a large, long and shallow canal on the medial side of the lamina ascendens.

The ventral surface of the pterygoid presents three kinds of ornamentation. In ventral view, the medial part of the basiptyergoid ramus and the margin delimiting the interptyergoid vacuities are smoothed. This ornamentation pattern on the ventral surface of the pterygoid is also found in several trematosaurids (e.g. *Thoosuchus yakovlevi*, Damiani and Yates 2003; *Lyrocephaliscus euri*, Mazin and Janvier 1983 and personal observations of T.A. on MNHN F SVT 520; *Trematosaurus galae*, Novikov 2010), presenting variation in the relative extension of the ornamentation.

The parasphenoid is divided in two parts: the cultriform process anteriorly and the basal plate posteriorly. The cultriform process is a long and narrow rod, laterally compressed, connecting the anterior part of the basal plate to the vomers.

It makes the separation between the two interptyergoid vacuities.

On the dorsal side of the cultriform process, two crests, the ‘cristae laterosphenoidales’ sensu Schoch (1999) are visible. They are more marked in the posterior part of the cultriform process. The ventral side of the cultriform process is convex in almost its entire length, with a strong lateral compression that gives a “knife-edge” shape to the cultriform process.

The basal plate of the parasphenoid is almost rectangular (length of 18.1 mm and width of 15.3 mm) with a pointed anterior side and a posterior projection (only visible in ventral view because it is covered by the exoccipital in dorsal view). Laterally, the basal plate shares a complex suture with the pterygoids, a typical stereospondyl character (e.g. Yates and Warren 2000), and posteriorly with the exoccipital.

The anterior margin of the basal plate of the parasphenoid is slightly concave and forms the posterior limit of the interptyergoid vacuities and the beginning of the cultriform process.

On the dorsal surface of the basal plate of the parasphenoid, two crests are posterolaterally located, named ‘cristae parapterygoideae’ sensu Bystrow and Efremov (1940: fig. 9A). They have a transverse orientation, like in *Lyrocephaliscus euri* (Mazin and Janvier 1983: fig. 7; personal observations of T.A. on MNHN F SVT 520).

Lateral to the ventral side of the posterior projection there is a pair of well-separated muscular pockets, as in *Thoosuchus yakovlevi* (Damiani and Yates 2003): they show a curved anterior border, weakly marked, and open posteriorly.

Regarding the interptyergoid vacuities, they are placed in middle palatal position and occupy more than half of the palate. Each vacuity forms an elongated oval with a length of 63 mm (right one) and 64.1 mm (left one) and a width of 16.2 and 15.3 mm respectively. The anterior margin of the interptyergoid vacuities is more pointed than the posterior one.

The subtemporal vacuity is oval with an elongated anterior projection, more pointed than the posterior end of the vacuity. Its total length is 37.6 mm (right), and 34.6 mm (left) and maximum width of 19.3 mm (right) and 18.6 mm (left). The margins of this vacuity are formed by the pterygoid, which forms the major part of it, the jugal, the quadratojugal and the quadrate. The jugal, the quadratojugal and the quadrate are difficult to observe from this view because they are broken.

4.5 Dentition

The dentition is characterized by small teeth, uniform in morphology and size, and six fangs that are bigger than the other teeth. These fangs are usually aligned with rows of small teeth. Both of the fangs and the smaller teeth are

labyrinthodont. There is one fang of about 5 mm in diameter between each anterior palatal vacuity and the choana. This fang, placed on the vomer, precedes a row of between 5 and 7 small teeth (Figs. 1, 2) that surrounds the internal border of each choana. The exact number of teeth that forms the row is difficult to determinate due to poor preservation. Between the two vomers, closely placed on the upper part of the choana, there is a slight bone elevation with a V-shaped morphology. The presence of teeth in this area is doubtful but cannot be discarded. There are two additional fangs on the palatines. One palatine fang is lost. The other palatine fang, with a diameter of about 4 mm, is followed by a row of 4–6 small teeth. After this row of teeth, on the ectopterygoid, there is another badly-preserved fang. This fang is closely placed to a last row of small teeth that goes through the ectopterygoid. This row is longer than the previous ones but the exact number of teeth cannot be assessed.

4.6 Occipital region

The exoccipital is formed by the columna verticalis, bearing the articular condyle, and four processes: the processus submedullaris, the processus lamellosus, the processus paroticus and the processus subtympanicus (Figs. 1, 2, 3). The exoccipital delimits two openings on the skull roof: the foramen magnum has a keyhole shape (14 mm × 9 mm at maximum size) as in *Thoosuchus yakovlevi*, and a posttemporal fenestra with an isosceles subtriangular shape with the tip laterally oriented (6 mm in height × 11 mm in width for the right one and 6 mm × 10 mm for the left one). For this posttemporal fenestra, the shape is the same as in *Thoosuchus yakovlevi*, although it is larger in *Angusaurus*. The size of the posttemporal fenestra corresponds to 33% of the skull height and 26% of the skull length in *Angusaurus*, against 20 and 22% respectively in *Thoosuchus yakovlevi* (personal observations of T.A. on specimen SMNS 58880). Skull height is measured as the distance between the skull roof and the base of the parasphenoid at the level of the suture between the two postparietals, whereas skull length corresponds to the distance between the two tabular horns (since a part of the left cheek is missing in *Angusaurus*, the distance between the two cheeks was not used). Once again, *Angusaurus* and *Thoosuchus yakovlevi* present similar skull shape.

The columna verticalis contacts a posterior column-like projection of the postparietal dorsally. Beginning from the ventral side, the suture between the exoccipital and the postparietal is located at three quarters of the occipital height (Fig. 1e, f, 3c).

The processus submedullaris is located medially on the columna verticalis, at one-third of the height of the exoccipital (Fig. 3a). The dorsal surface of the processus submedullaris is concave with a truncated oval shape but the anterior

border is straight, similar to *Benthosuchus sushkini* (Bystrow and Efremov 1940: fig. 11).

The processus lamellosus is located above the processus submedullaris and medial to the columna verticalis, at around mid-height of the foramen magnum. It is relatively poorly developed compared to *Benthosuchus sushkini* (Bystrow and Efremov 1940: fig. 11). The dorsal surface of this process is concave with an oval shape.

The processus paroticus is projects dorsolaterally from the columna verticalis and forms half of the ventral part of the paroccipital process with the descending flange of the tabular. Posterolaterally, a wide foramen is present as in *Benthosuchus sushkini* (Bystrow and Efremov 1940: fig. 11) or *Lyrocephaliscus euri* (Mazin and Janvier 1983: fig. 4.)

The processus subtympanicus is an anterolaterally-projected sheet of bone. It is rectangular, but the left one has a rounded anterior extremity that makes it longer than the right one.

The occipital condyles lie in posteromedial position in relation to the columna verticalis. They are as high as wide and well separated from each other. The posterior and medial surfaces are rough.

On the anterior side of the exoccipital, between the base of the processus lamellosus and paroticus, there is a weak rounded depression. Due to this emplacement and by comparison with *Stanocephalosaurus amenasensis* (Arbez et al. 2017), it could represent the opisthotic bone.

4.7 Preservation of internal structures

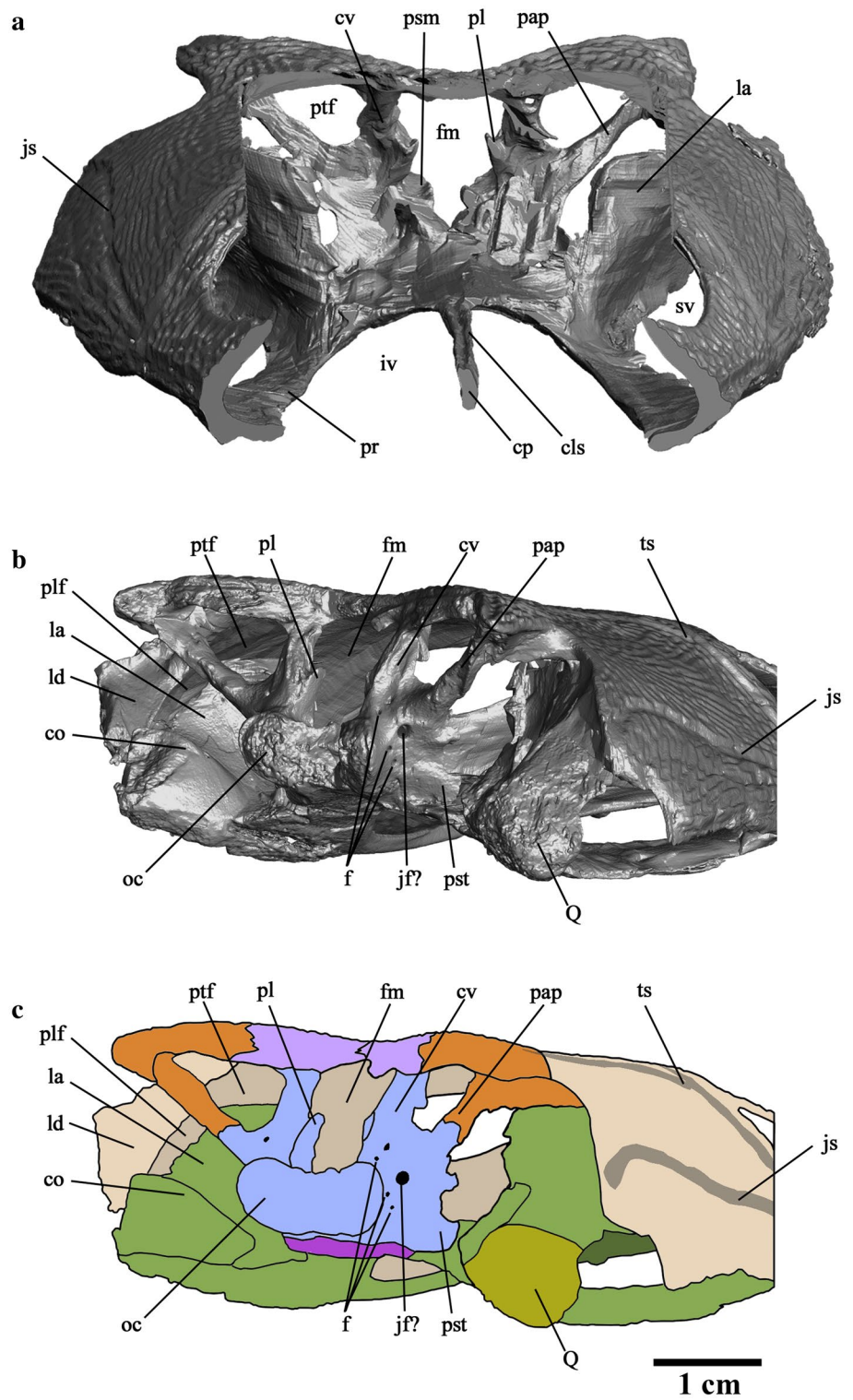
Unfortunately, the stapes, otic capsule, epipterygoid, sphenethmoid and basisphenoid are not preserved in SMNS 81782. However, this lack of preservation is not surprising, since the inner structures of stereospondyls are generally poorly ossified, likely cartilaginous, and thus rarely preserved in the fossil record. In trematosaur, the sphenethmoid is only present in *Aphaneramma* sp. (Säve-Söderbergh 1936) and *Benthosuchus sushkini* (Bystrow and Efremov 1940), and only *Aphaneramma* sp. is known to have a preserved basisphenoid (Säve-Söderbergh 1936).

Moreover, in species where the stapes, the otic capsule and the epipterygoid are preserved, these bones possess few and slight contacts with surrounding bones. For example, in *S. amenasensis* the epipterygoid is only linked to the pterygoid and likely by a cartilaginous contact (Arbez et al. 2017). As explained above, because of their weak ossification these elements are easily lost during fossilisation.

4.8 Phylogenetic analyses

The phylogenetic position of the genus including all species of *Angusaurus* was assessed via cladistic analyses. We used the data matrix of Fortuny et al. (2018), which is based

Fig. 3 3D model of *Angusaurus* cf. *tsylmensis* (SMNS 81782). Neurocranium and inner parts (a), oblique view of the occiput (b) and interpretative drawing of the oblique view (c)



on Steyer (2002), but includes some coding changes due to observation of new specimens as well as additional taxa. Herein, we added two more taxa (*Syrtosuchus samarensis*, *Prothoosuchus blomi*) based on the literature (Getmanov 1989; Novikov 2016). We tested the genus *Angusaurus* in

two different analyses. In one of the analyses, we coded the three species *A. dentatus*, *A. succedaneus* and *A. tsylmensis* separately, while in the second analysis we coded all three *Angusaurus* species as a group (labeled as *Angusaurus* spp.). All the terminal taxa were considered at the specific level

(except for the *Angusaurus* cluster). *Onchiodon frossardi* and *Arhegosaurus decheni* were considered as outgroups. In total, 25 terminal taxa and 69 characters were analyzed (see supplementary material for the data matrix) using the software PAUP 4.0 beta for PC (Swofford 2001). All characters were considered equally weighted and unordered. Eight characters from the data matrix were found to be uninformative, and excluded from the analyses. These excluded characters are (same as in the previous published analysis, Fortuny et al. 2018): 16 (frontal contact orbit), 17 (interfrontal), 28 (vomer contact maxilla), 46 (paraquadrate foramen), 56 (stapes morphology), 62 (parasymphysial tooth

row), 63 (denticles coronoids) and 67 (crista articularis). We performed a heuristic search of 10,000 replicates with simple addition sequence and only one tree retained at each step during stepwise addition. Tree-Bisection-Reconnection (TBR) branch swapping was performed and zero-length branches were collapsed to yield polytomies.

The analysis treating the three *Angusaurus* species (*A. dentatus*, *A. succedaneus*, *A. tsylmensis*) as separate species, recovered five most parsimonious trees (MPT) of 202 steps (CI 0.396, RI 0.4831, RC 0.1913) (Fig. 4). The consistency index is low and reveals a high amount of homoplasy in trematosaur evolution. A bootstrap analysis with 1000

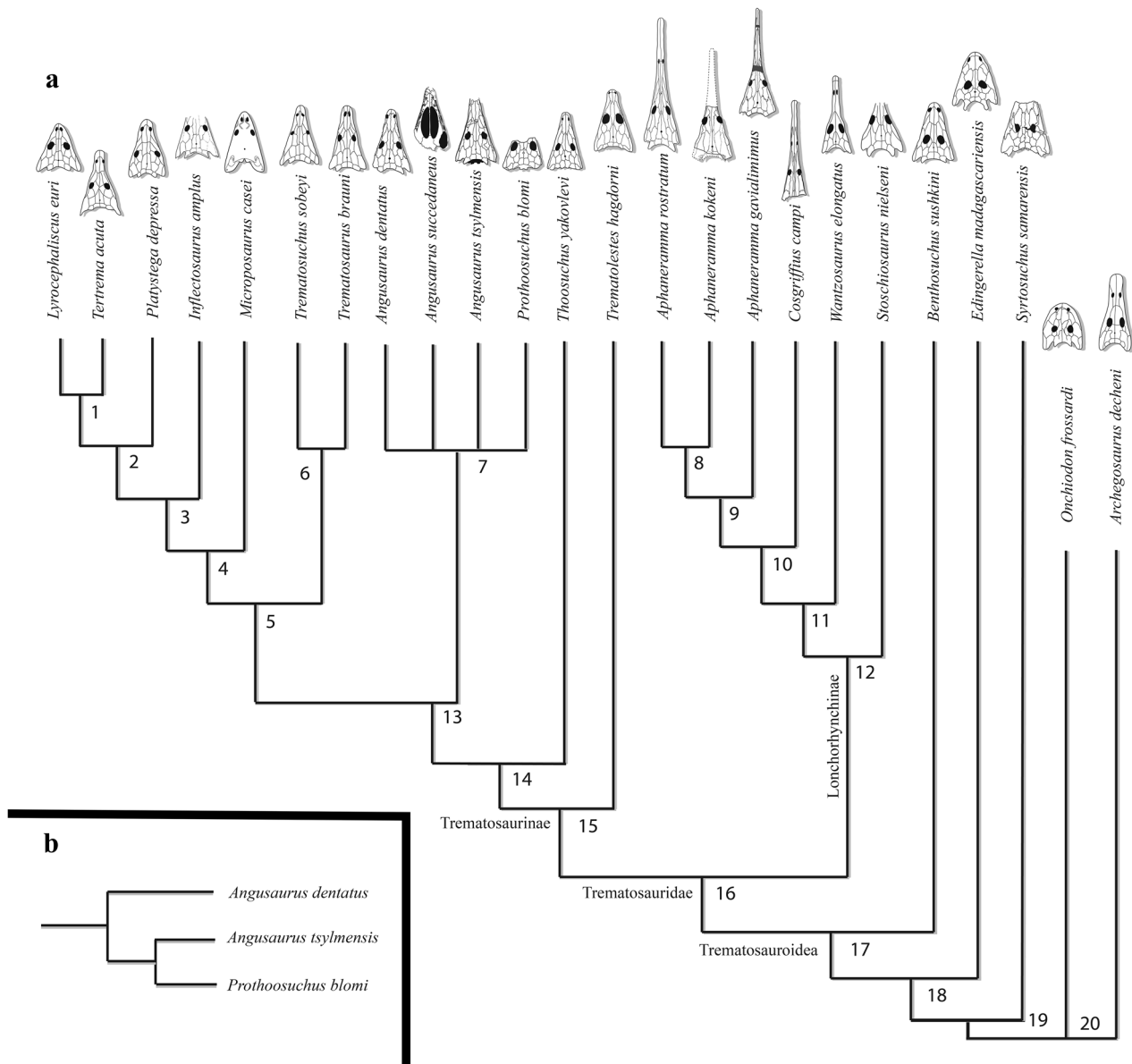


Fig. 4 Phylogenetic position of *Angusaurus* within the Trematosauroidae clade. Strict consensus tree from the five Most Parsimonious Trees (MPT) recovered under a heuristic analysis (see text) (**a**).

Results for *Angusaurus* taxa if *A. succedaneus* is removed from the analysis, rest of tree typology identical (**b**). Node numbers refer to clades (see text and supplementary material)

replicates also reveals relatively low support, with only two clades with support over 50% (Stereospondyli: 61% and *Aphaneramma rostratum*+*Aphaneramma kokeni*: 51%). The monophyletic Lonchorhynchinae (node 12) is supported by four synapomorphies: orbits facing laterally [5(1)], knife-edged cultriform process [38(2)], absence of the crista muscularis of the parasphenoid [42(0)], and carotid canal visible on the ventral surface of the parasphenoid [43(1)].

In this analysis, Lonchorhynchinae is the sister-taxon of Trematosaurinae (node 15), as found by Steyer (2002) and Fortuny et al. (2018). The Trematosaurinae clade is composed by the same taxa as in previously recovered trees (Fortuny et al. 2018), but with some taxa in different positions (see below). Trematosaurinae is supported by twelve synapomorphies: orbits facing laterally [5(1)], nostril in lateral position [9(1)], pineal large relative to the size of the skull roof [11(1)], frontal extended behind orbit [15(1)], premaxilla/nasal suture straight [24(1)], ventral opening of the orbit in the anterior half of the interpterygoid vacuity [27(2)], presence of paired anteropalatal vacuities [31(2)], anteropalatal vacuities between the premaxilla/vomerine suture [32(1)], knife-edged cultriform process of the parasphenoid [38(2)], absence of the crista muscularis of the parasphenoid [42(0)], absence of the crista falciformis of the squamosal, in occipital view [53(0)], and Meckelian foramen elongate [65(1)]. The clade joining Trematosaurinae and Lonchorhynchinae is the Trematosauridae (clade 16). This clade is supported by five synapomorphies: nostril ovoid [8(1)], otic notch deep and open [20(1)], anteropalatal vacuities between the premaxilla/vomerine suture. [32(1)], occipital condyles widely separated from each other [50(0)], and well-developed crista medialis [66(1)]. As previously recovered by Fortuny et al. (2018), Trematosauridae and *Benthosuchus sushkini* form the clade Trematosauroida (node 17), which is supported by ten synapomorphies: small orbit [4(1)], quadrate condyle anterior to the occipital condyle [22(1)], interpterygoid vacuity not widened at all [33(2)], anterior branch of the pterygoid entirely wide [35(1)], ventrally directed tabular in occipital view [48(1)], foramen magnum ventrally constricted [51(1)], presence of the crista falciformis of the squamosal in occipital view [53(1)], deep posttemporal fenestra [54(1)], posterior end of the mandible situated behind the quadrate condyle [57(1)], and extended dentary symphysis [60(1)].

The taxon *Syrtosuchus samarensis* (Novikov 2016) is here found as the most basal taxon of the analyzed taxa. Novikov (2016) erected this taxon and considered it “as the most primitive benthosuchid taxon assigned to a separate benthosuchid subfamily, Syrtosuchinae(…)” (Novikov 2016: p. 307). In the present analysis this taxon represents a stem trematosauroid, and potentially a stem capitosauroid. Future analyses focusing on basal trematosauroids and related taxa

should clarify the position of this basal taxon and assess its phylogenetic position.

Lonchorhynchinae is composed by the same taxa (and equal positions) found by Fortuny et al. (2018), while Trematosaurinae is composed by the same taxa as in Fortuny et al. (2018), with the addition of *Prothoosuchus blomi* (not included in previous analyses). *Trematolestes hagdorni* is found as the most basal trematosaurine, representing the sister taxon of *Thoosuchus yakovlevi*. In turn, this latter taxon is the sister taxon of the clade formed by all three *Angusaurus* species analyzed as well as *Prothoosuchus blomi*, resulting in a polytomy between these four taxa (node 7, Fig. 4). It should be mentioned that a heuristic analysis (under the same variables) but not including *Angusaurus succedaneus* resulted in only one Most Parsimonious Tree (MPT) with the same topology as above but with *Angusaurus dentatus* recovered as the sister taxon of the clade composed by *Angusaurus tsylmensis* and *Prothoosuchus blomi* (Fig. 4). In a similar way, the analysis grouping all three *Angusaurus* species (labeled as *Angusaurus* spp.) resulted in only one MPT with the same general topology but recovering a clade formed by *Angusaurus* spp. and *Prothoosuchus blomi*.

Regarding the rest of trematosaurines, this latter analysis recovered a clade (node 6) composed by *Trematosaurus brauni* and *Trematosuchus sobeyi*, not recovered in previous analyses by Fortuny et al. (2018) but present in analyses made by Steyer (2002). Finally, *Inflexosaurus* and *Microposaurus* appear as derived trematosaurines (Damiani 2004; Novikov 2007; Warren 2012; Fortuny et al. 2018) and closely related to *Platystega*, *Lyrocephaliscus* and *Tertrema*.

5 Discussion

5.1 Taxonomic and ontogenetic assessment of SMNS 81782

Trematosauroids are characterized by the formation of the anterior portion of a long and narrow cultriform process from the posterior projections of the vomer (Schoch and Milner, 2000), a feature also found in thoosuchids and in the studied specimen. In the specimen described neither the frontal nor the lacrimal bones have any contact with the orbits, a derived feature also present in *Thoosuchus* and other trematosauroids (Damiani and Yates 2003). In capitosauroids the lacrimal does not contact the orbits but the frontal maintains its orbit contact, causing a wide separation between the prefrontal and the postorbital, like in *Edingerella* (Schoch and Milner 2000; Maganuco et al. 2009).

In the trematosaurines *Thoosuchus* and *Angusaurus*, as well as SMNS 81782, the supraorbital sulcus runs anteriorly through the nasal, between the nares, and then enters the

lacrimal, whereas in most capitosauroids this sulcus runs along almost the entire nasal and enters the lacrimal posteriorly (Fig. 1; Damiani and Yates 2003; Maganuco et al. 2009; Schoch and Milner 2000). The presence of the occipital canal is typical of many trematosauroids like *Thoosuchus* and *Angusaurus*, while its absence is characteristic of capitosauroids, like *Edingerella* (Maganuco et al. 2009).

Small trematosaurids exclusively known from the Lower Triassic of the Eastern European Platform are usually referred to Thoosuchidae. This group is characterized by a narrow skull with concave lateral margins and a longer snout than in other trematosauroids, and by the lateral margin of the skull being straight or slightly convex like in *Benthosuchus* (Schoch and Milner 2000; Novikov 2012a). All these features are also true for the specimen studied herein.

The flexura lacrimalis is represented by a Z-shaped flexura on the lacrimal of the infraorbital sulcus. This flexura is present in thoosuchids, such as *Thoosuchus* and *Angusaurus*, as well as in the specimen described, being softer and less sharp in the latter than in other trematosauroids like *Benthosuchus* (Getmanov 1989; Schoch and Milner 2000; Damiani and Yates 2003; Novikov 2012a).

The jugals of thoosuchids are thinner in comparison with other trematosauroids. These features can be also shared with primitive trematosauroids. In adult benthosuchids, the jugal can have the width of its orbits (Schoch and Milner 2000). The palatine of thoosuchids, as well as the specimen analyzed here, has no posterior process or contact with its pterygoid, whereas the palatines of *Benthosuchus* connects with the pterygoid (Novikov 2012a; Schoch and Milner 2000).

Taking into account all these features and the fact that the specimen SMNS 81782 cannot be referred to Benthosuchidae—because members of this family have a bilobed anterior palatal vacuity—, nor to Plastystegididae—because its members have a very broad postorbital skull—this specimen is assigned with confidence to Thoosuchidae (sensu Schoch and Milner 2000). Regarding *Thoosuchus*, the specimen cannot be referred to this genus based on orbit size (bigger in *Thoosuchus*) as well as the elongation of the postorbital and postfrontal in the studied specimen relative to *Thoosuchus*. The genus *Angusaurus* (Figs. 5, 6, 7), as previously mentioned, is characterized by posteriorly elongated postorbitals and postfrontals (Schoch and Milner 2000), as in the studied specimen. The main part of the body of the jugal of the studied specimen is situated in a posterior position, behind the orbits, (Getmanov 1989). The body of the parasphenoid of *Angusaurus* is slightly longer (Getmanov 1989) than in the studied specimen (Figs. 1, 2, 3, 6). As in SMNS 81782, the pineal foramen of the genus *Angusaurus* is situated at half the distance between the orbits and the occipital, in a

middle position on the parietals (Getmanov 1989). All these characters allow us to assign the studied specimen to the genus *Angusaurus*.

To date, four *Angusaurus* species have been erected: *A. dentatus*, *A. succedaneus*, *A. weidenbaumi* and *A. tsylmensis* (Figs. 5, 6, 7). Unfortunately the diagnostic characters for each one of these species are tenuous. In addition, some of the specimens used in published descriptions are differently preserved in the homologous regions, which precludes in some cases anatomical comparison. *A. dentatus* (type species of the genus) and *A. tsylmensis* are erected based on almost complete skulls (but missing some parts of the pterygoids and parasphenoid in *A. dentatus*, as well as any mandible remain known for this taxon) (Figs. 5, 6, 7) whereas *A. weidenbaumi* is only known by a snout (not presenting the typical *flexura lacrimalis* sulcus found in the rest of *Angusaurus* taxa) and *A. succedaneus* by a partial skull with well-preserved palate and mandible but missing most of the skull roof and occiput, but associated with a well preserved mandible (Figs. 5, 6, 7).

Hence, the key anatomical characters in the analyzed specimen are: (1) (character 36; see supplementary material) posterior branch of the pterygoid, narrow and elongate (*A. dentatus*) or short and wide (*A. tsylmensis* and SMNS 81782); (2) presence of flexura lacrimalis (*A. dentatus*, *A. succedaneus*, *A. tsylmensis* and SMNS 81782) or absence (*A. weidenbaumi*); (3) the anterior and lateral border of the basal plate of the parasphenoid are almost straight (*A. dentatus*) or concave (*A. succedaneus*, *A. tsylmensis* and SMNS 81782); (4) ventral surface of the parasphenoid plate smoothed (*A. dentatus* and SMNS 81782) or dot-shaped (*A. succedaneus*, *A. tsylmensis*); (5) postemporal fenestra high (the fenestra itself, not its position) (*A. dentatus* and SMNS 81782) or low (*A. tsylmensis*).

Nonetheless, the last two characters should be taken with caution: the ornamentation is known to vary with ontogeny (Steyer 2000), and the height difference of the fenestra is not really pronounced and cannot be excluded for preservation reasons (compression).

Overall, all these characters suggest affinities of the studied specimen with *A. tsylmensis* or *A. succedaneus*, and prevent the assignment of the former to *A. dentatus* and *A. weidenbaumi*. However, an attribution to *A. succedaneus* is not possible to conclude due to the poor state of preservation of some diagnostic characters and anatomical areas (e.g. quadrate ramus) in the holotype (and unique published specimen) of *A. succedaneus*, and it cannot be discarded that some shared characters with this taxon may be due to ontogeny. Thus, based on the shared characters discussed, SMNS 81782 is referred to as *A. cf. tsylmensis*. The lack of information on the stratigraphy of the studied

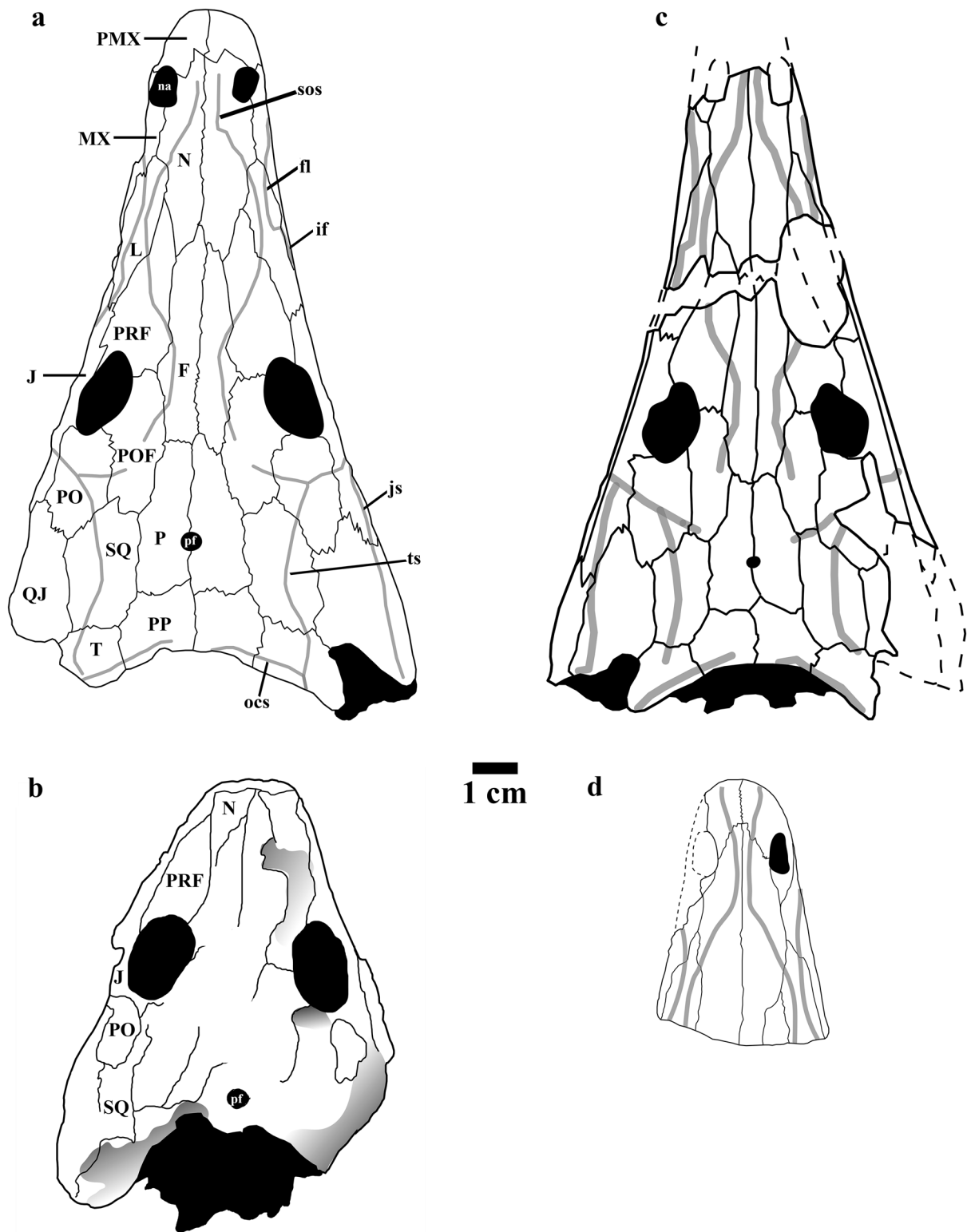
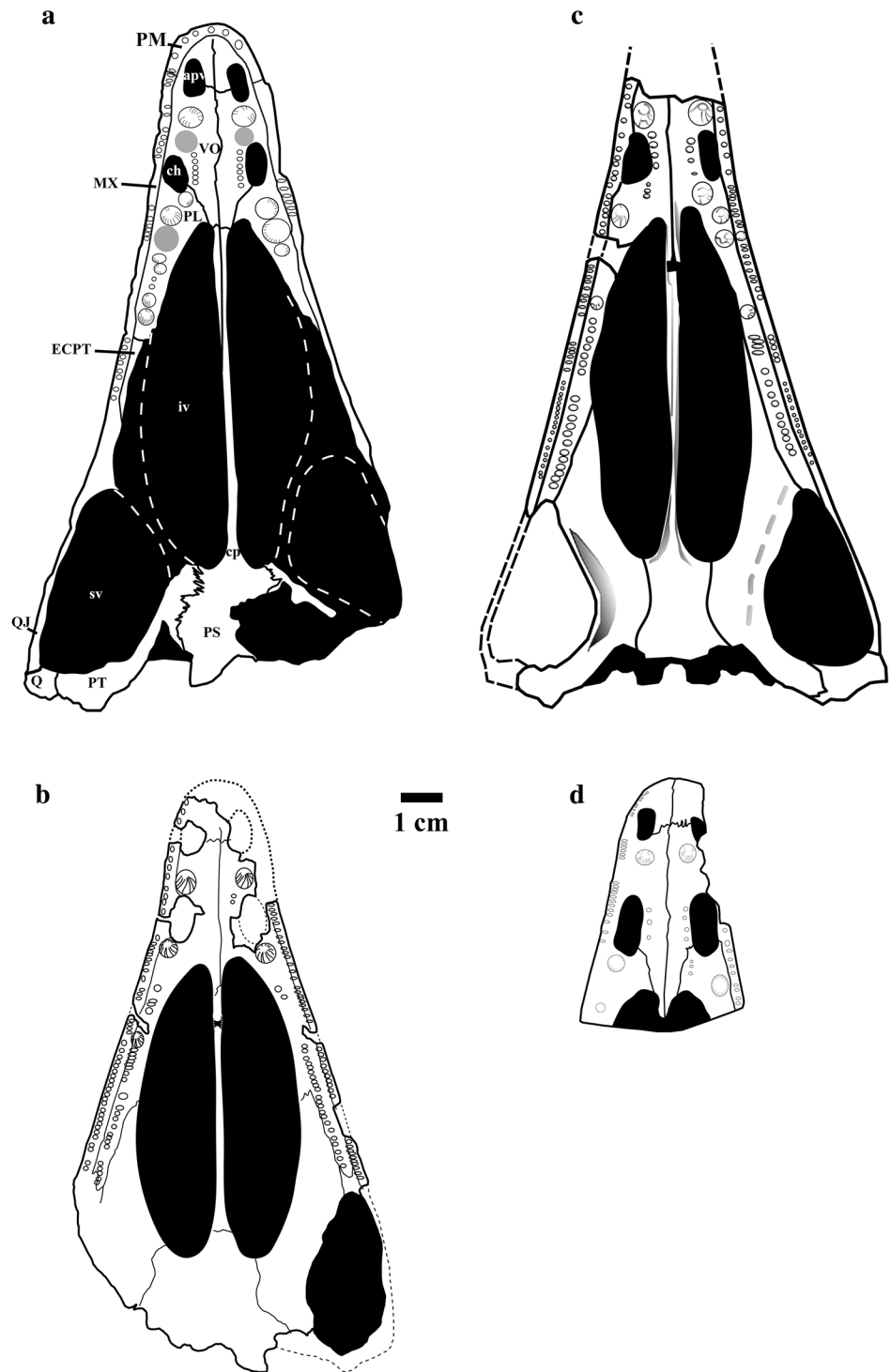


Fig. 5 The genus *Angusaurus* from the Early Olenekian (Lower Triassic) of Russia. Dorsal view of: *A. dentatus* (a), *A. succedaneus* (b), *A. tsylmensis* (c), *A. weidenbaumi* (d). Redrawn after Getmanov

1989, Novikov 1990 and personal observations (J.F.). Note that lines for cranial sutures in *A. succedaneus* are tentative

Fig. 6 The genus *Angusaurus* from the Early Olenekian (Lower Triassic) of Russia. Ventral view of: *A. dentatus* (a), *A. succedaneus* (b), *A. tsylmensis* (c) and *A. weidenbaumi* (d). Redrawn after Getmanov 1989, Novikov 1990 and personal observations (J.F.)

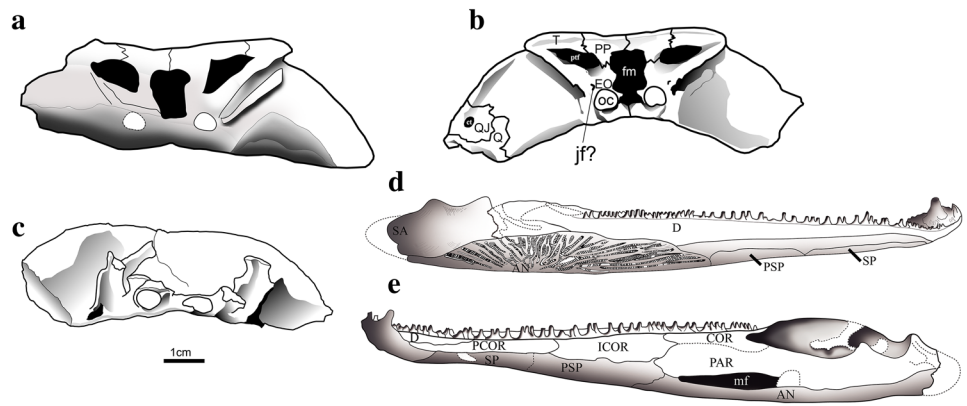


specimen, as well as potential interspecific, intraspecific, ontogenetic or preservation variability preclude a confident assignment to *A. tsylmensis*.

As previously mentioned, the approximate length of the adult skull for *Angusaurus* is about 200 mm (Getmanov 1989), while the studied skull has a length of 112.3 mm, indicating a possible juvenile condition. The

radial sculpture of the dermal bones is typically found in growing bones and it can thus be a larval or juvenile character, although this ornamentation can also be found in some adult stereospondyls. Polygonal sculpture comes from the radial sculpture pattern (Witzmann et al. 2010), and SMNS 81782 has mainly polygonal ornamentation in the ossification centre of the skull roof. Although we

Fig. 7 The genus *Angusaurus* from the Early Olenekian (Lower Triassic) of Russia. Occipital view of: *A. dentatus* (a), *A. succedaneus* (b), *A. tsylmensis* (c) and mandible in labial (d) and lingual (e) side of *A. succedaneus*. Redrawn after Getmanov 1989, Novikov 1990 and personal observations (J.F.)



interpret SMNS 81782 as a sub-adult, we cannot reject that this specimen could be an adult considering that the radial pattern can also be present in adults. Sometimes the degree of ossification and the sutural fusion is indicative of a developmental age but in temnospondyls it might not be a significant indicator (Steyer 2000).

5.2 *Angusaurus*: taxonomic and phylogenetic considerations

As previously mentioned, four *Angusaurus* species have been erected to date (*A. dentatus*, *A. succedaneus*, *A. weidenbaumi* and *A. tsylmensis*) (Figs. 5, 6, 7). This genus is apparently uniform, with few diagnostic characters allowing differentiation between species. This fact raises the question of the validity of these species. When three species (*A. dentatus*, *A. succedaneus*, and *A. tsylmensis*) are included separately in the phylogenetic analysis, it results in a polytomy (Fig. 4). In particular, two of these species (*A. dentatus* and *A. succedaneus*) are virtually equal under the character matrix (depending on the missing data), and it should be considered that both species were recovered from the same subhorizon.

Previously, Schoch and Milner (2000) also questioned the validity of *A. succedaneus*, suggesting it as potential junior synonym of *A. dentatus*. Nonetheless, these authors maintained the validity of these species.

The main reason why we do the same here is because of the difficulty to directly compare *A. dentatus*, *A. succedaneus*, and *A. weidenbaumi* (see above). Although these three taxa are recovered from the same subhorizon (see below), there is a lack of homologous regions preserved in these three species. However, we cannot discard the possibility that future findings may potentially reveal that only one or two of these species are valid, thus raising the possibility of evaluating the intraspecific variability of the genus *Angusaurus*.

On the other hand, *A. tsylmensis* is considered more confidently as a valid species because it was recovered from a

different (younger) subhorizon (Ust-Mylian) and presents the morphological characters previously discussed (e.g. short and wide posterior branch of the pterygoid; the more anterior position of the quadrate condyle, etc.) (Figs. 5, 6, 7).

5.3 Biostratigraphy and paleobiogeography of basal trematosauroids, trematosaurids and trematosaurines

Most trematosaurines have been recovered from the Russian European platform, with the exception of some taxa (e.g. *Trematolestes hagdorni*, *Trematosaurus brauni*, *Trematosuchus sobeyi* etc.) from Central Europe, Spitsbergen, South Africa, Madagascar, North America and Australia. Regarding the communities of Eastern Europe, three different main regions (or realms) can be recognized: (a) the central and northern regions of the East European Platform (Moscow–Mezen, and presumably Pechora, Synclines), and the Timan–North Ural Region; (b) the southern CisUrals, including the Obshchii Syrt Plateau; and (c) the southern regions of the East European Platform (the slope of the Voronezh Anticline) (Shishkin et al. 2006).

From a chronological point of view, these main regions span much of the Lower and Middle Triassic (from the Induan to the Ladinian), with a succession up to 3.5 km and subdivided in different superhorizons (or gorizonts, here used as synonyms but see Tverdokhlebov et al. 2002 for discussion): Vetlugian superhorizon (Induan—Olenekian), Yarenskian horizon (Olenekian), Donguz horizon (Anisian) and Bukobay horizon (Ladinian). Up to seven faunal assemblages (or svitas, here used as synonyms but see Tverdokhlebov et al. 2002 for discussion) have been proposed for these superhorizons and horizons: Kopanskaya (Induan), Staritskaya, Kzylsaiskaya, Gostevskaya, and Petropavlovskaya (all Olenekian), Donguz (Anisian), and Bukobay (Ladinian) (see Ochev and Shishkin 2000; Tverdokhlebov et al. 2002: fig. 2; Shishkin et al. 2006 for comprehensive reviews, correlation between horizon and assemblages, as well as additional references).

Herein, we focus on basal trematosauroids, trematosaurids and trematosaurines recovered from the Vetlugian superhorizon and associated with the *Benthosuchus*—*Wetlugasaurus* tetrapod complexes. This superhorizon spans the following subhorizons (or gorizonts), from base to top: Vokhmian, Rybinskian, Sludkinskian and Ust-Mylian (Ochev and Shishkin 2000: fig. 1 for a discussion). Vetlugian superhorizon is dated as Olenekian with the exception of the Vokhmian subhorizon (Induan). This latter subhorizon is of particular interest for trematosauroids, since different taxa (as part of the Kopanskaya faunal assemblage) have been recovered from this subhorizon and particularly from the main region of the southern CisUrals and the Obshchii Syrt Plateau. Remarkably, *Syrthosuchus* and primitive *Benthosuchus* taxa (e.g. *B. gusevae*) have also been recovered, as well as other temnospondyls such as *Tupilakosaurus* and *Wetlugasaurus samariensis* (Tverdokhlebov et al. 2002; Novikov 2012a, 2016).

Nonetheless, a recent proposed new horizon, Zaplavneniskian Horizon, has been proposed. Its faunal assemblage is still controversial but suggests a late developmental stage of the *Tupilakosaurus* fauna (see Shishkin and Novikov 2017 and references therein).

The Rybinskian Gorizont (Staristskaya faunal assemblage) is mainly composed of fluvial sandstones, with the usual features of arid zones (Tverdokhlebov et al. 2002). *Benthosuchus sushkini* and the genus *Thoosuchus* are the most characteristic taxa for this gorizont, forming a *Benthosuchus-Thoosuchus* group, typical of this subhorizon but also including other temnospondyl members as *Wetlugasaurus*, *Trematotegmen otschevi* and *Qantas samarensis* (Shishkin et al. 2000; Tverdokhlebov et al. 2002; Novikov 2012b). The Sludkinskian subhorizon (Kzylsaiskaya faunal assemblage) is very similar to the previous one, but in the uppermost part of the section the fluvial deposits are sometimes replaced by deltaic sediments. The fauna is similar to the Rybinskian subhorizon, particularly on the lowermost part of the section. In this faunal assemblage *Wetlugasaurus angustifrons* is the most common tetrapod, but it also contains other temnospondyls like *Wetlugasaurus kzilsajensis*, *Benthosuchus bashkiricus*, *Thoosuchus tuberculatus*, *Angusaurus dentatus*, *Angusaurus succedaneus*, *Angusaurus weidenbaumi*, *Prothoosuchus blomi*, and *Prothoosuchus samariensis*, and the first appearance of the capitosaur genus *Parotosuchus* (Shishkin et al. 2000; Tverdokhlebov et al. 2002).

Finally, the Ust-Mylian subhorizon (Gostevskaya faunal assemblage and equivalent to the assemblage III of Novikov 1994) constitutes a complex of lacustrine-deltaic accumulations where the most characteristic temnospondyl fauna is composed of *Wetlugasaurus malachovi* as well as *Angusaurus tsylmensis* and *Vyborososaurus mirus*. It also represents the first appearance of the genus *Trematosaurus* as well as the capitosaur *Vladlenosaurus alexeyevi* and indeterminate

plagisaurids (Ochev and Shishkin 2000; Tverdokhlebov et al. 2002; Shishkin et al. 2006). It should be remarked that this faunal assemblage is the most poorly known due to difficulties recognizing this fauna outside of the northeastern areas of the Russian platform.

Later, the temnospondyl fauna was replaced by the *Parotosuchus* group, which dominated the Yarenskian subhorizon (Tverdokhlebov et al. 2002).

Taking into account these previous considerations, the Early Olenekian time is of particular interest to understand the origin and diversification of basal trematosaurines. This time is characterized by a eustatic maximum (Lozovsky 1992), which resulted in the expansion of lakes in Eastern Europe (Shishkin et al. 2006) and represented by two subhorizons corresponding to the Early Olenekian: the Sludkinskian (early Early Olenekian) and the Ust-Mylian (late Early Olenekian) (Ochev and Shishkin 2000; Tverdokhlebov et al. 2002). The chronological origin of trematosauroids, trematosaurids and trematosaurines is debated, since the earliest fossil evidence for these groups ranges from (?Late) Induan to Early Olenekian, known from different continents. On the one hand, basal trematosauroids such as *Syrthosuchus* sp. and *Benthosuchus gusevae* are recorded from the (?Late) Induan (Vokhmian subhorizon) from the Southern Cis-Ural area—Obshchii Syrt Plateau (Novikov 2012a, 2016). Nonetheless, the poorly known genus *Tirratourhinus* from the Arcadia Formation (central Queensland, Australia) has been dated as (Early) Induan and proposed as a trematosaurine (Nield et al. 2006; Warren et al. 2006; but see discussion about alternative dating and taxonomic position in Shishkin et al. (2006) and Novikov (2012b).

On the other hand, recent phylogenetic analyses place the genus *Trematolestes* as the most basal trematosaurine (Fortuny et al. 2018, present study), but also resembling in some characters lonchorhynchines (Schoch 2006). This well-known taxon is recorded from the Middle Triassic (Lower Keuper facies, Upper Ladinian) of Germany and is closely related to the Early Triassic Malagasy genus *Tertremoides* (Schoch 2006). If this is correct, *Tertremoides*, recovered from the NW of Madagascar (Ankitokazo basin, Sakamena Group, Ambilobe region), has to be considered a basal trematosaurine. This genus is found in similar levels among other trematosaurids such as *Wantzosaurus* and *Aphaneramma* (Steyer 2002; Fortuny et al. 2018). The Ankitokazo basin was originally dated as Induan (Scythian A1), but later re-dated as Olenekian based on conchostracans (Yanbin et al. 2002).

Overall, the first trematosauroid datum implies an open paleobiogeographical scenario where basal trematosauroids may have originated during the Induan at the Cis-Uralian-Obshchii Syrt Plateau region of the Eastern Europe platform (Novikov 2016), but without excluding the possibility of an Australian (Nield et al. 2006) or East Greenland (see

below) trematosauroid origin and/or even suggesting paleobiogeographical link between East Greenland, Cis-Uralian-Obshchii Syrt Plateau (Russia) and Queensland (Australia). Otherwise, the trematosauroid split between short-snouted forms (e.g. trematosaurines) and long snouted ones (lonchorhynchines), is particularly unclear based on the presence of the most basal members of these groups during the Late Induan- (?Early) Olenekian in different continents: East Greenland (*Stoschiosaurus*, Late Induan, Nielsen 1935; Zaton et al. 2016); Eastern Europe (*Thoosuchus*—Early Olenekian, Ochev and Shishkin 2000) and Madagascar (*Tertremoides*—Olenekian unprecise age, Maganuco and Pasini 2009; *Wantzosaurus*—Olenekian unprecise age, Steyer 2002). Taking into account these data, we come up with, at least, two hypotheses: (A) the first appearance of basal trematosauroids and trematosaurids is older than current fossil record evidence suggests; (B) supposed basal lonchorhynchines and trematosaurines (as *Stoschiosaurus* or *Tertremoides*) may be basal trematosauroids instead of lonchorhynchines or trematosaurines, and consequently a taxonomical re-assignment and definition of these groups is required (provided that new material is found).

Lastly, the trematosaurine radiation that started in Eastern Europe with the appearance of *Thoosuchus* and subsequent related genera such as *Angusaurus*, *Prothoosuchus* and *Vyborosaurus* during the Early Olenekian potentially expanded to Central European basins (with paleobiogeographical connection between the southern regions of the East European Platform, Shishkin et al. 2006) represented by the presence of the genus *Trematosaurus* in both areas (Schoch and Milner 2000; Novikov 2010). The *Trematosaurus* radiation during the Late Olenekian also derived in the presence of advanced trematosaurines such as *Inflexosaurus*, *Platystega*, *Tertrema* and *Lyrocephaliscus* in the Eastern European platform and Spitsbergen (Svalbard), and also in southern latitudes such as South Africa (Karoo basin) during the Late Olenekian-Early Anisian, represented by *Trematosuchus* and *Microposaurus* (Schoch and Milner 2000).

6 Conclusions

The genus *Angusaurus*, present from the early Early Olenekian to late Early Olenekian in Russia, is a trematosaurine characterized by skulls about 200 mm long (adult size), small orbits (compared to the close genus *Thoosuchus*), elongated postorbitals and postfrontals, with a very reduced interchoanal tooth row and a ventral opening of the orbits in the middle part of the skull, presenting a mandible with an elongate postglenoid process (PGA) and well developed suprangular. The new specimen described herein sheds some light on the ontogeny of this genus, with implications for

other trematosaurines. The genus *Angusaurus* includes four species with few key anatomical characters. While the type species, *Angusaurus dentatus*, and *Angusaurus tsylmensis* are well supported, the validity of *Angusaurus succedaneus* and *Angusaurus weidenbaumi* is controversial and new findings are required to fully evaluate their validity and better understand the intraspecific variability of this genus. Deciphering the evolution of trematosaurines during the Early Olenekian is key to understanding the origin and diversification of basal trematosaurines with implications to clarify the different paleobiogeographical scenarios that remain still open.

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
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Affiliations

Meritxell Fernández-Coll¹ · Thomas Arbez² · Federico Bernardini^{3,4} · Josep Fortuny^{2,5} 

✉ Josep Fortuny
josep.fortuny@icp.cat

¹ Faculty of Life and Environmental Sciences, University of Iceland, Askja, Sturlugata 7, 101 Reykjavík, Iceland

² Centre de Recherches sur la Paléobiodiversité et les Paléoenvironnements, UMR 7207 CNRS-MNHN-UPMC, Muséum national d'Histoire naturelle, CP38, 8 rue Buffon, 75005 Paris, France

³ Centro Fermi, Museo Storico della Fisica e Centro di Studi e Ricerche, 'Enrico Fermi', Piazza del Viminale 1, 00184 Rome, Italy

⁴ Multidisciplinary Laboratory, the 'Abdus Salam' International Centre for Theoretical Physics, Via Beirut 31, 34151 Trieste, Italy

⁵ Institut Català de Paleontologia M. Crusafont, C\de les Columnes, s/n, Universitat Autònoma de Barcelona, 08193 Cerdanyola del Vallès, Spain

Table 1. Measurements of *Angusaurus cf. tsylmensis* (SMNS 81782). Measurements in mm.

1	Skull length	112.3	Distance from the premaxilla to the posterior end of the postparietals
2	Skull width	74.1	Maximum width of the skull
3	Skull height	28.1	Maximum height of the skull
4	Right orbit length	19.1	Maximum length of the right orbit
5	Right orbit width	13.0	Maximum width of the right orbit
6	Left orbit length	17.9	Maximum length of the left orbit
7	Left orbit width	12.3	Maximum width of the left orbit
8	Posterior interorbital distance	39.0	Distance between the posterior edge of both orbits
9	Interorbital distance	23.3	Distance between the wider part of both orbits
10	Distance from right orbit to the premaxilla	77.2	Distance from the posterior edge of the right orbit to the premaxilla
11	Distance from right orbit to the right otic notch	31.4	Distance from the posterior edge of the right orbit to the right otic notch
12	Distance from left orbit to the premaxilla	76.3	Distance from the posterior edge of the left orbit to the premaxilla
13	Distance from left orbit to the left otic notch	31.8	Distance from the posterior edge of the left orbit to the left otic notch
14	Pineal foramen diameter	4.0	Pineal foramen diameter

15	Distance from Pineal foramen to the end of postparietal	19.5	Distance from the posterior edge of the Pineal foramen to the posterior end of postparietal
16	Distance from Pineal foramen to the orbits	8.7	Distance from Pineal foramen to the posterior edge of the orbits
17	Distance between both Tabular horns	46.3	Distance between the end of both tabular horns
18	Minimum distance between exoccipitals condyles	8.1	Distance between the distal point innermost of the exoccipitals condyles
19	Maximum distance between exoccipitals condyles	16.2	Distance between the distal point outermost of the exoccipitals condyles
20	Parasphenoid length	18.1	Maximum length of the parasphenoid bone
21	Parasphenoid width	15.3	Maximum width of the parasphenoid bone
22	Maximum right pterygoid width	8.8	Distance between the parasphenoid bone and the right supratemporal vacuity
23	Maximum left pterygoid width	10.0	Distance between the parasphenoid bone and the left supratemporal vacuity
24	Right pterygoid width	8.4	Right pterygoid width between right supratemporal vacuity and right interpterygoid vacuity
25	Left pterygoid width	8.0	Left pterygoid width between left supratemporal vacuity and left

			interpterygoid vacuity
26	Cultriform process posterior width	13.9	Width of the cultriform process in a posterior position
27	Cultriform process medial width	0.8	Width of the cultriform process in a middle position
28	Cultriform process anterior width	6.3	Width of the cultriform process in an anterior position
29	Right supratemporal vacuity length	37.6	Maximum length of the right supratemporal vacuity
30	Right supratemporal vacuity width	19.3	Maximum width of the right supratemporal vacuity
31	Left supratemporal vacuity length	34.6	Maximum length of the left supratemporal vacuity
32	Left supratemporal vacuity width	18.6	Maximum width of the left supratemporal vacuity
33	Right interpterygoid vacuity length	63	Maximum length of the right interpterygoid vacuity
34	Right interpterygoid vacuity width	16.2	Maximum width of the right interpterygoid vacuity
35	Left interpterygoid vacuity length	64.1	Maximum length of the left interpterygoid vacuity
36	Left interpterygoid vacuity width	15.3	Maximum width of the left interpterygoid vacuity
37	Right choana length	11.9	Maximum length of the right choana
38	Right choana width	4.2	Maximum width of the right choana

39	Left choana length	11.9	Maximum length of the left choana
40	Left choana width	4.3	Maximum width of the left choana
41	Right exoccipital condyle diameter	6.1	Right exoccipital condyle diameter
42	Left exoccipital condyle diameter	5.7	Left exoccipital condyle diameter
43	Total foramen magnum length	14.0	Maximum length of the foramen magnum
44	Maximum foramen magnum width	9.4	Maximum width of the foramen magnum
45	Right posttemporal fenestra length	4.2	Total length of the right posttemporal fenestra
46	Right posttemporal fenestra width	10.1	Total width of the right posttemporal fenestra
47	Left posttemporal fenestra length	4.9	Total length of the left posttemporal fenestra
48	Left posttemporal fenestra width	8.7	Total width of the left posttemporal fenestra

Appendix 1

Character list

Characters of the skull roof

1. Skull roof elongate (midline length > maximum width; 1) or not (0).
2. Position of the centre of the orbit along the dorsal midline of the skull: orbit posteriorly (0) or anteriorly (1) situated.
3. Short (0), constricted (1), or elongated (2) snout (preorbital part of the skull twice longer than the postorbital one; 2).
4. Large (0) or small orbit (maximum size < 14 per cent of the dorsal midline length; 1).
5. Orbit facing laterally (1) or dorsally (0).
6. Rounded (0) or oval (1) orbit.
7. Slightly concave (0) or semicircular (1) posterior margin of the skull roof.
8. Nostril rounded (0), ovoid (1), or elongate (2).
9. Nostril in lateral position (1) or not (0).
10. Skull roof bulged (1) or not (0) at the level of the orbits.
11. Pineal foramen small (0) or large (1) relative to the size of the skull roof.
12. Pineal foramen rounded (0) or not (1).
13. Presence (1) or absence (0) of sensory-line grooves in front of the orbits.
14. Presence (0) or absence (1) of sensory-line grooves behind orbits.
15. Frontal extended behind orbit (1) or not (0).
16. Frontal in contact with orbit (1) or not (0).
17. Presence (1) or absence (0) of the interfrontal.
18. Presence (1) or absence (0) of the centroparietal.
19. Septomaxilla visible (0) or not (1) on the dorsal side of the skull.
20. Otic notch deep and narrow (0), or deep and open (1), or shallow and open (2).
21. Tabular rounded (0), pointed (1), or hook-shaped (2).
22. Quadrate condyle posterior (0) or anterior (1) to the occipital condyle.
23. Presence (1) or absence (0) of an anterodorsal dentary foramen.
24. Premaxilla/nasal suture posteriorly directed (the premaxilla is partly extended posterior to the nostril; 0), straight (1), or anterior directed (2).
25. Presence (1) or absence (0) of a prenasal growth zone.
26. Presence (1) or absence (0) of a 'temporal fossa' (sensu Damiani 1998; a slight depressed region of the skull roof anteriorly to the otic notches).
27. Ventral opening of the orbit at the level of the posterior half (0), in the mid part (1), or in the anterior half (2) of the interpterygoid vacuity.

Characters of the palate

28. Vomer in contact (1) or not (0) with the maxilla.
29. Choana rounded (0) or elongate (1).
30. Choana overlapping the nostril (1) or not (0).
31. Absence (0) or presence of a single (1) or of a paired (2) anteropalatal vacuity.
32. Anteropalatal vacuity(ies) posterior (0), between (1), or anterior (2) to the premaxilla/vomerine suture.
33. Interpterygoid vacuity posteriorly (0) or anteriorly (1) widened, or not widened at all (2).
34. Presence (1) or absence (0) of a crista obliqua on the ventral surface of the pterygoid.
35. Anterior branch of the pterygoid laterally extended (0), entirely wide (1), or narrow (2).

36. Posterior branch ('quadrate ramus') of the pterygoid narrow and elongate (0), or short and wide (1).
37. Presence (0) or absence (1) of an area asparta of the pterygoid.
38. Wide (0), narrow (1) or knife-edged (2) cultriform process of the parasphenoid.
39. Elongate (0) or very short (or absent) (1) suture between the palatine and the pterygoid.
40. Very short (0) or elongate (1) suture between the pterygoid and parasphenoid.
41. Wide (width>length; 0) or narrow (length > width; 1) parasphenoid plate.
42. Presence (1) or absence (0) of the crista muscularis of the parasphenoid.
43. Carotid canal visible (1) or not (0) on the ventral surface of the parasphenoid.
44. Presence (0) or absence (1) of ectopterygoidal tusks.
45. Suture between the exoccipital and the pterygoid visible (1) or not (0) in ventral view.

Characters of the occiput

46. Presence (1) or absence (0) of the paraquadrate foramen (foramen for the chorda tympani).
47. Flattened (0) or deep (1) occiput, in occipital view.
48. Dorsally (0) or ventrally (1) directed tabular, in occipital view.
49. Small (0) or large (1) occipital condyle.
50. Occipital condyles widely separated (0) or not (1) from each other.
51. Foramen magnum ventrally constricted (1) or not (0).
52. Curved (0) or straight (1) dorsal margin of the foramen magnum.
53. Presence (1) or absence (0) of the crista falciformis of the squamosal, in occipital view.
54. Shallow (0) or deep (1) posttemporal fenestra.
55. Elongate (0), triangular (1) or rounded (2) posttemporal fenestra.
56. Wide (0) or narrow (1) proximal part of the stapes.

Characters of the mandible

57. Posterior end of the mandible situated at the same level (0) or behind (1) the quadrate condyle.
58. Mandible deep (0) or shallow (1) in lateral view.
59. Presence (1) or absence (0) of an anterior constriction of the ventral outline of the mandible, in lateral view.
60. Reduced (0) or extended (1) dentary symphysis.
61. Presence (1) or absence (0) of a parasymphysial tusk.
62. Presence (0) or absence (1) of a parasymphysial tooth row.
63. Presence (1) or absence (0) of denticles on coronoids 2 and 3.
64. Presence (1) or absence (0) of sensory-line grooves on the mandible.
65. Meckelian foramen short (0) or elongate (1).
66. Poorly developed or absent (0), or well-developed (1) crista medialis (sensu Damiani 1998); a vertical blade on the midline of the postglenoid area (PGA sensu Jupp and Warren 1986).
67. Presence (1) or absence (0) of the crista articularis (sensu Damiani 1998); a vertical blade on the postero-lingual border of the PGA.

Characters of the postcranial skeleton

68. Interclavicle narrow (1) or not (0).

69. Interclavicle with (0) or without (1) dorsal crest(s) (trabeculae sensu Bystrow and Efremov 1940; i.e. relatively wide and flat walls posteriorly and antero-laterally directed on the dorsal surface of the interclavicle).

Institutional Abbreviations

NHMUK: The Natural History Museum, London, UK; **MGUH:** Museum Geologicum Universitatis Hafniensis, Kobenhavn, Denmark; **MNB:** Museum für Naturkunde, Berlin, Germany; **MNHN:** Muséum national d'Histoire naturelle, Paris, France; **MSNM:** Museo di Storia Naturale di Milano, Italy; **PIN:** Paleontological Institute, Moscow, Russian Federation; **SMNS:** Staatliches Museum für Naturkunde in Stuttgart, Germany; **UA:** Université d'Antananarivo, Madagascar.

Terminal taxa	Specimens	Literature
<i>Angusaurus dentatus</i>	PIN 4196/1	Getmanov 1989; Damiani and Yates, 2003
<i>Angusaurus succedaneus</i>		Getmanov, 1989
<i>Angusaurus tsylmensis</i>	PIN 4333/6	Novikov, 1990
<i>Aphaneramma gavialimimus</i>	UA-Amb007, MNHN-6703	Fortuny et al. in press
<i>Aphaneramma rostratum</i>	MNHN 20-IBSEN 1969, MNHN Rotunda 1964, MNHN St 62, MNHN Tr A 64	Säve-Söderbergh 1936, 1937; Welles 1993; Smith-Woodward 1904; Wiman 1916
<i>Aphaneramma kokeni</i>		Huene 1920; Welles, 1993
<i>Archegosaurus decheni</i>	MNB Am 114-119, 121-131, 137 MNHN 1868-1-4 1870-479-480, MNHN 1884-10, 21-25;	Gubin 1997; Jaeckel 1896; Witzmann 2006
<i>Benthosuchus sushkini</i>	PIN 2424/4, PIN 2424/10, PIN 2-19-2252, PIN 2252/41	Bystrow and Efremov, 1940
<i>Cosgriffius campi</i>		Welles 1993
<i>Edingerella madagascariensis</i>	MNHN MAE3000-3009; MSNM V2992	Damiani 1998; Lehman 1961; Warren 1980; Maganuco et al. 2009; Steyer 2003
<i>Inflectosaurus amplius</i>		Jaeckel 1922; Shishkin 1960; Novikov 2007
<i>Lyrocephaliscus euri</i>	MNHN F SVT 520	Säve-Söderbergh 1935, 1936, 1937; Mazin and Janvier 1983
<i>Microposaurus casei</i>		Damiani 2004
<i>Onchiodon frossardi</i>	MNHN 1908-20-26	Werneburg and Steyer 1999

<i>Platystega depressa</i>		Säve-Söderbergh 1936; Wiman 1916
<i>Prothoosuchus blomi</i>		Getmanov, 1989
<i>Stoschiosaurus nielseni</i>	MGUH At. 6, 7, 9, 12, 23, 43, 45, 46	Säve-Söderbergh 1935
<i>Syrthosuchus samarensis</i>		Novikov, 2016
<i>Thoosuchus yakovlevi</i>	SMNS 58880	Damiani and Yates, 2003; Novikov, 2007
<i>Tertrema acuta</i>		Säve-Söderbergh 1936; Wiman 1916
<i>Trematolestes hagdorni</i>	SMNS 81790-91, SMNS 90022,	Schoch 2006
<i>Trematosaurus brauni</i>	BMNH 30270, 36354-75, 40042 MNB Am943 1/3 MNHN AC9573	Burmeister 1849; Drevermann 1920; Lehman 1966, 1979;
<i>Trematosuchus sobeyi</i>		Haughton 1915; Shishkin and Welman 1994
<i>Wantzosaurus elongatus</i>	MNHN MAE 3030, 3034;	Lehman 1961; Steyer 2002

Apomorphy lists:

Branch	Character	Steps	CI	Change

node_20 --> node_19	4	1	0.200	0 --> 1
node_19 --> node_18	13	1	0.500	0 --> 1
	39	1	1.000	0 --> 1
	40	1	1.000	0 --> 1
	50	1	0.333	0 --> 1
node_18 --> node_17	4	1	0.200	1 --> 0
	22	1	0.200	0 --> 1
	33	1	0.286	0 --> 2
	35	1	0.400	0 --> 1
	48	1	0.333	0 --> 1
	51	1	1.000	0 --> 1
	53	1	0.333	0 --> 1
	54	1	1.000	0 --> 1
	57	1	0.333	0 --> 1
	60	1	1.000	0 --> 1
node_17 --> node_16	8	1	0.250	2 --> 1
	20	1	0.333	0 --> 1
	32	1	0.400	0 --> 1
	50	1	0.333	1 --> 0
	66	1	1.000	0 --> 1
node_16 --> node_12	5	1	0.500	0 --> 1
	38	1	1.000	0 --> 2
	42	1	0.200	1 --> 0
	43	1	0.250	0 --> 1
node_12 --> node_11	9	1	0.200	0 --> 1
	25	1	0.500	0 --> 1

	31	1 0.667 1 --> 2
node_11 --> node_10	8	1 0.250 1 --> 2
	23	1 1.000 0 --> 1
	27	1 0.250 0 --> 2
node_10 --> node_9	4	1 0.200 0 --> 1
	15	1 0.250 0 --> 1
	35	1 0.400 1 --> 2
	47	1 0.500 0 --> 1
	55	1 0.500 0 --> 1
node_9 --> <i>A. gavialimimus</i>	8	1 0.250 2 --> 1
	21	1 0.400 0 --> 2
	41	1 0.200 0 --> 1
	44	1 0.250 0 --> 1
node_9 --> node_8	8	1 0.250 2 --> 1
	9	1 0.200 1 --> 0
	18	1 1.000 0 ==> 1
	21	1 0.400 0 --> 2
	24	1 0.286 2 --> 1
	36	1 0.250 0 --> 1
	41	1 0.200 0 --> 1
	42	1 0.200 0 --> 1
	44	1 0.250 0 --> 1
	49	1 0.333 0 --> 1
	50	1 0.333 0 --> 1
	64	1 0.500 0 --> 1
	65	1 0.500 0 --> 1
node_8 --> <i>A. kokeni</i>	33	1 0.286 2 ==> 0
node_8 --> <i>A. rostratum</i>	27	1 0.250 2 ==> 1
node_10 --> <i>C. campi</i>	4	1 0.200 0 --> 1
	15	1 0.250 0 --> 1

	31	1 0.667 2 --> 0
node_11 --> <i>W. elongatus</i>	7	1 0.333 0 ==> 1
	8	1 0.250 1 --> 2
	11	1 0.250 0 ==> 1
	20	1 0.333 1 ==> 2
	22	1 0.200 1 ==> 0
	23	1 1.000 0 --> 1
	27	1 0.250 0 ==> 1
	30	1 0.500 0 ==> 1
	33	1 0.286 2 ==> 0
	35	1 0.400 1 --> 2
	47	1 0.500 0 --> 1
	55	1 0.500 0 --> 1
node_12 --> <i>S. nielsenii</i>	25	1 0.500 0 --> 1
	37	1 0.500 0 ==> 1
node_16 --> node_15	5	1 0.500 0 --> 1
	9	1 0.200 0 --> 1
	11	1 0.250 0 ==> 1
	15	1 0.250 0 ==> 1
	24	1 0.286 2 --> 1
	27	1 0.250 0 --> 2
	31	1 0.667 1 --> 2
	38	1 1.000 0 --> 2
	42	1 0.200 1 --> 0
	53	1 0.333 1 ==> 0
	65	1 0.500 0 --> 1
node_15 --> node_14	19	1 0.250 1 ==> 0
	22	1 0.200 1 ==> 0
	55	1 0.500 0 --> 1
	58	1 0.333 1 ==> 0

	68	1 0.250 1 ==> 0
	69	1 0.333 1 ==> 0
node_14 --> node_14	3	1 0.333 2 ==> 1
	4	1 0.200 0 ==> 1
	21	1 0.400 0 --> 1
	24	1 0.286 1 --> 2
	61	1 0.500 0 --> 1
node_13 --> node_5	8	1 0.250 1 --> 0
	9	1 0.200 1 ==> 0
	41	1 0.200 0 ==> 1
	48	1 0.333 1 --> 0
	53	1 0.333 0 ==> 1
node_5 --> node_4	12	1 0.500 0 --> 1
	29	1 0.333 1 ==> 0
	49	1 0.333 0 ==> 1
	52	1 1.000 0 --> 1
	57	1 0.333 1 --> 0
	59	1 0.333 0 --> 1
	64	1 0.500 0 --> 1
node_4 --> node_3	27	1 0.250 2 ==> 1
	32	1 0.400 1 --> 2
node_3 --> <i>I. amplus</i>	5	1 0.500 1 ==> 0
	22	1 0.200 0 ==> 1
	55	1 0.500 1 ==> 0
	56	1 1.000 1 ==> 0
node_3 --> node_2	33	1 0.286 2 ==> 1
	35	1 0.400 1 ==> 0
	36	1 0.250 0 ==> 1
	68	1 0.250 0 --> 1
node_2 --> node_1	3	1 0.333 1 ==> 0

	11	1 0.250 1 --> 0
	20	1 0.333 1 --> 0
node_1 --> <i>L. euri</i>	1	1 0.500 1 ==> 0
	4	1 0.200 1 ==> 0
	20	1 0.333 0 --> 2
	24	1 0.286 2 ==> 0
	27	1 0.250 1 ==> 2
	41	1 0.200 1 ==> 0
	42	1 0.200 0 ==> 1
	43	1 0.250 0 ==> 1
	45	1 0.500 0 ==> 1
	48	1 0.333 0 --> 1
	55	1 0.500 1 ==> 2
node_1 --> <i>T. acuta</i>	8	1 0.250 0 ==> 2
	9	1 0.200 0 ==> 1
	19	1 0.250 0 ==> 1
	29	1 0.333 0 ==> 1
	37	1 0.500 0 ==> 1
	47	1 0.500 0 ==> 1
node_2 --> <i>P. depressa</i>	10	1 0.500 0 ==> 1
	15	1 0.250 1 ==> 0
	32	1 0.400 2 --> 1
	44	1 0.250 0 ==> 1
node_4 --> <i>M. casei</i>	2	1 0.500 0 ==> 1
	8	1 0.250 0 --> 2
	10	1 0.500 0 ==> 1
	25	1 0.500 0 ==> 1
	33	1 0.286 2 ==> 0
node_5 --> node_6	3	1 0.333 1 ==> 0
	20	1 0.333 1 ==> 2

	21	1 0.400 1 --> 0
	32	1 0.400 1 --> 0
	45	1 0.500 0 ==> 1
node_6 --> <i>T. brauni</i>	8	1 0.250 0 --> 1
	19	1 0.250 0 ==> 1
	22	1 0.200 0 ==> 1
	24	1 0.286 2 ==> 1
node_6 --> <i>T. sobeyi</i>	6	1 0.500 1 ==> 0
	13	1 0.500 1 ==> 0
	14	1 0.500 0 ==> 1
	17	1 1.000 0 ==> 1
	34	1 0.333 1 ==> 0
	35	1 0.400 1 ==> 0
node_13 --> node_7	27	1 0.250 2 ==> 1
node_ <i>A.dentatus</i> + <i>[A. tsylmensis+P.bloomi</i> --> node_ <i>A.tsyl+P.bloomi</i>	36	1 0.250 0 ==> 1
node_ <i>A.tsyl+P.bloomi</i> --> <i>P. bloomi</i>	15	1 0.250 1 ==> 0
node_14 --> <i>T. yakovlevi</i>	26	1 0.500 0 ==> 1
	27	1 0.250 2 ==> 0
	32	1 0.400 1 ==> 2
	42	1 0.200 0 ==> 1
	57	1 0.333 1 ==> 0
node_15 --> <i>T. hagdorni</i>	2	1 0.500 0 ==> 1
	3	1 0.333 2 ==> 0
	12	1 0.500 0 ==> 1
	33	1 0.286 2 ==> 1
	34	1 0.333 1 ==> 0
	36	1 0.250 0 ==> 1
	41	1 0.200 0 ==> 1
	43	1 0.250 0 --> 1

	59	1 0.333 0 ==> 1
node_17 --> <i>B. sushkini</i>	8	1 0.250 2 --> 1
	20	1 0.333 0 --> 1
	30	1 0.500 0 ==> 1
	32	1 0.400 0 --> 1
	43	1 0.250 0 --> 1
	44	1 0.250 0 ==> 1
	50	1 0.333 1 --> 0
	58	1 0.333 1 ==> 0
	61	1 0.500 0 ==> 1
	66	1 1.000 0 --> 1
	68	1 0.250 1 ==> 0
	69	1 0.333 1 ==> 0
node_18 --> <i>E. madagascariensis</i> 3	1	0.333 2 ==> 1
	4	1 0.200 1 --> 0
	7	1 0.333 0 ==> 1
	9	1 0.200 0 --> 1
	16	1 1.000 0 ==> 1
	21	1 0.400 0 ==> 1
	22	1 0.200 0 --> 1
	24	1 0.286 2 ==> 1
	26	1 0.500 0 ==> 1
	33	1 0.286 0 ==> 1
	35	1 0.400 0 --> 1
	41	1 0.200 0 ==> 1
	48	1 0.333 0 --> 1
	51	1 1.000 0 --> 1
	53	1 0.333 0 --> 1
	54	1 1.000 0 --> 1
	57	1 0.333 0 --> 1

	59	1 0.333 0 ==> 1
	60	1 1.000 0 --> 1
	63	1 1.000 0 ==> 1
	67	1 1.000 0 ==> 1
node_19 --> <i>S. samarensis</i>	13	1 0.500 0 --> 1
	20	1 0.333 0 --> 1
	33	1 0.286 0 --> 2
	39	1 1.000 0 --> 1
	40	1 1.000 0 --> 1
node_20 --> <i>A. decheni</i>	4	1 0.200 0 --> 1
	7	1 0.333 0 ==> 1
	11	1 0.250 0 ==> 1
	14	1 0.500 0 ==> 1
	21	1 0.400 0 ==> 1
	27	1 0.250 0 ==> 1
	35	1 0.400 0 ==> 2
	43	1 0.250 0 --> 1
	44	1 0.250 0 ==> 1
	49	1 0.333 0 ==> 1
	50	1 0.333 0 --> 1
node_20 --> <i>O. frossardi</i>	1	1 0.500 1 ==> 0
	3	1 0.333 2 ==> 0
	6	1 0.500 1 ==> 0
	8	1 0.250 2 ==> 0
	19	1 0.250 1 ==> 0
	24	1 0.286 2 ==> 0
	28	1 1.000 1 ==> 0
	29	1 0.333 1 ==> 0
	31	1 0.667 1 ==> 0
	34	1 0.333 1 ==> 0

42 1 0.200 1 ==> 0

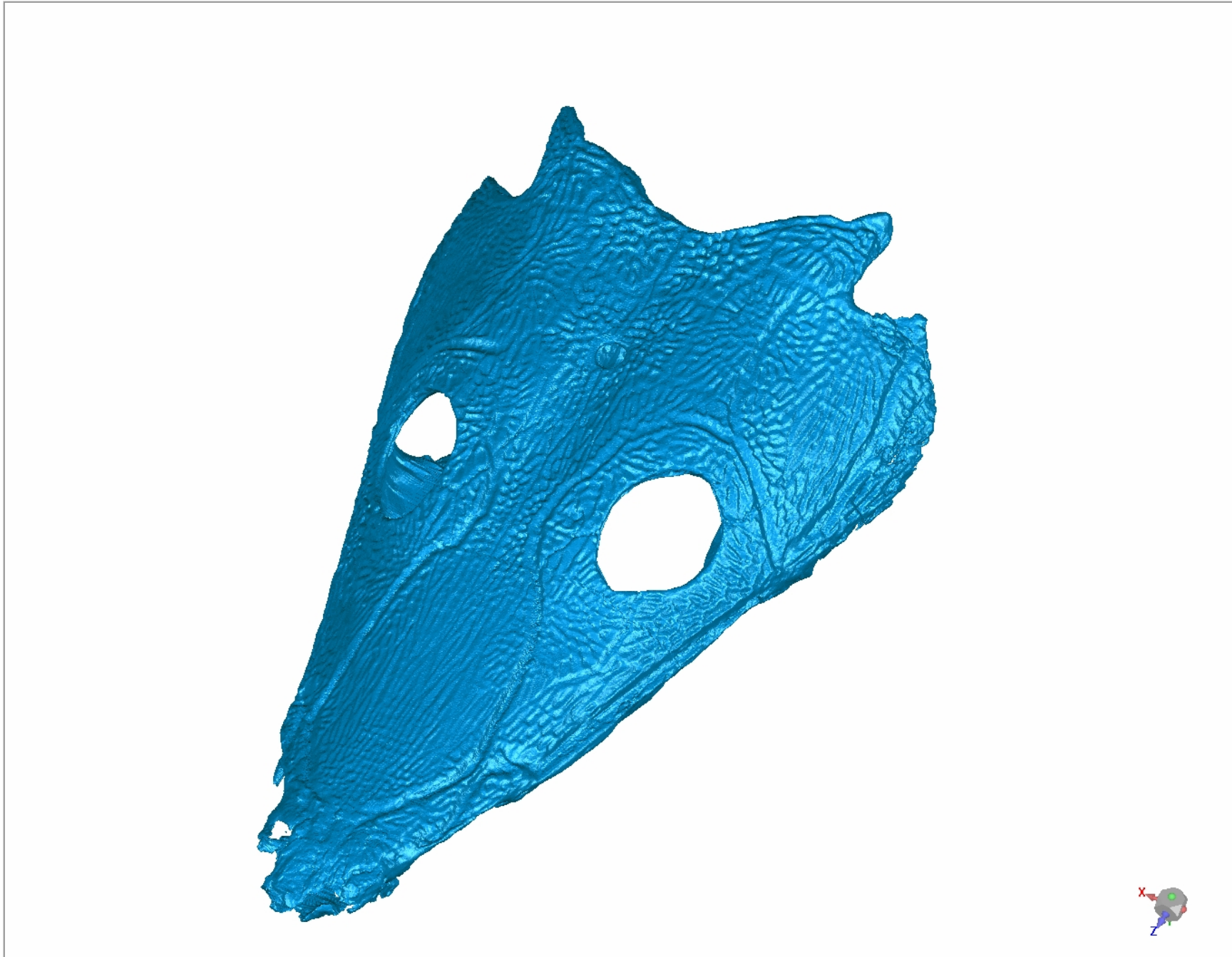
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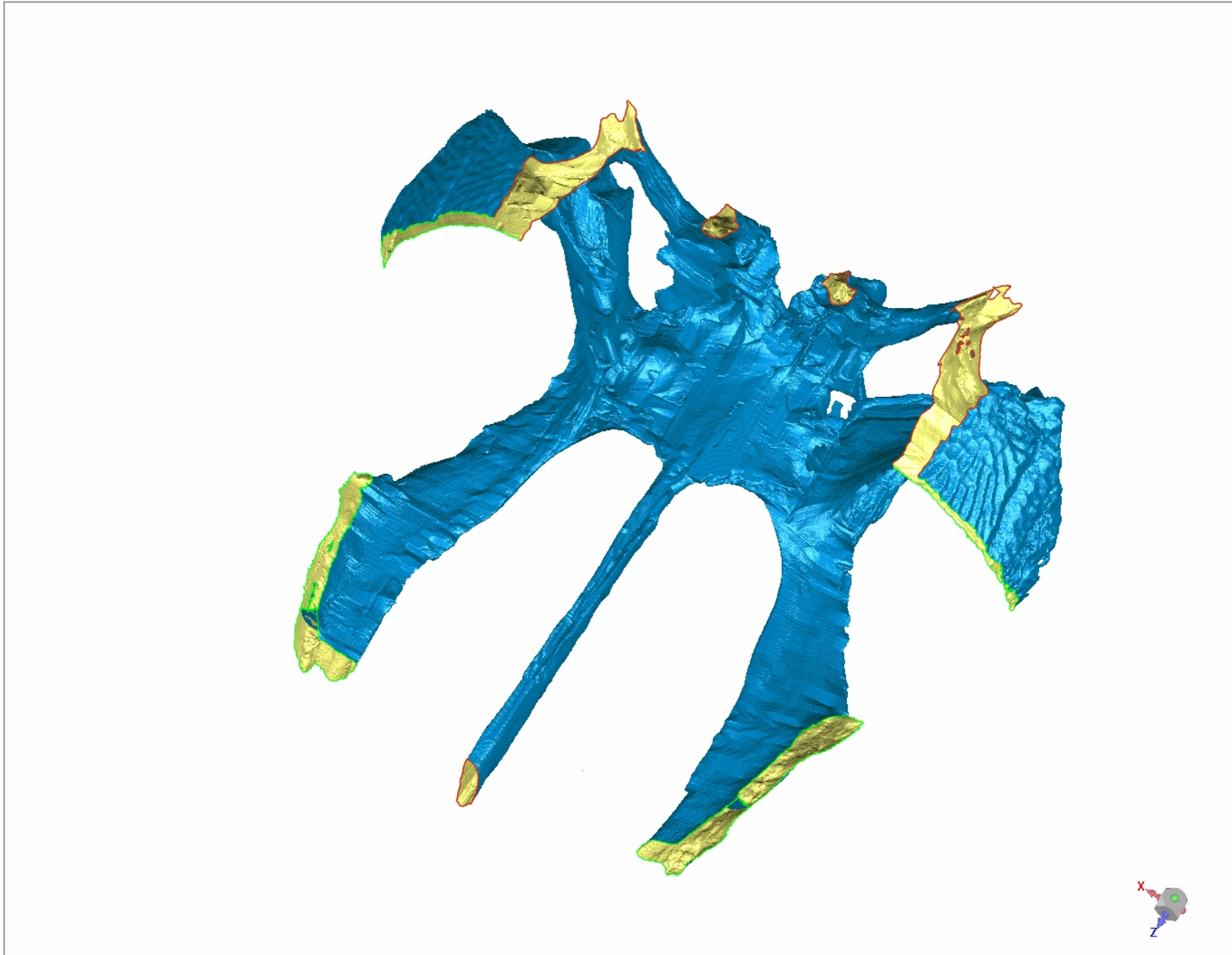
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68 1 0.250 1 ==> 0

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