

A new elasmosaurid plesiosaurian from the Early Cretaceous of Russia marks an early attempt at neck elongation

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Plesiosaurian marine reptiles evolved a wide range of body shapes during the Jurassic and Cretaceous, including long-necked forms. Many Late Cretaceous members of the clade Elasmosauridae epitomized this part of the plesiosaurian morphological spectrum by evolving extremely long necks through somitogenesis (resulting in an increase in the number of cervical centra) and differential growth (resulting in the elongation of cervical centra). However, the early evolution of elasmosaurids remains poorly understood because of a generally poor Lower Cretaceous fossil record. We describe a new elasmosaurid, *Jucha squalia* **gen. et sp. nov.**, from the upper Hauterivian (Lower Cretaceous) of Ulyanovsk (European Russia), in addition to other elasmosaurid remains from the same area. *Jucha squalia* is one of the oldest and basalmost elasmosaurids known and lacks a series of features that otherwise characterize the group, such as the heart-shaped intercoracoid fenestra and the median pectoral bar. However, *Jucha squalia* marks an early attempt at cervical elongation through differential growth. The data we gathered on the shape of cervical centra among elasmosaurids suggest multiple episodes of elongation and shortening. However, the precise patterns are obscured by an unstable phylogenetic signal.

ADDITIONAL KEYWORDS: cervical elongation – differential growth – Hauterivian – marine reptiles – Xenopsaria.

INTRODUCTION

Elasmosaurids are a clade of plesiosaurian marine reptiles notably characterized by an extreme

elongation of the neck, which contains several dozens of centra, sometimes > 70 (O’Keefe, 2002; Kubo *et al.*, 2012; Sachs *et al.*, 2013; Soul & Benson, 2017). Elasmosaurids are part of a wider clade, Xenopsaria, that diversified markedly during the Cretaceous (Benson & Druckenmiller, 2014), evolving a range of distinct morphologies with varying relative neck lengths (O’Keefe, 2002; Benson & Druckenmiller, 2014; Otero, 2016; Serratos *et al.*, 2017; Fischer *et al.*, 2018).

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However, the early evolution of Xenopsaria and the initial diversification of its main clades, Elasmosauridae and Leptocleididae, are poorly known and in a state of flux, because only a few taxa are known from the Early Cretaceous and these frequently switch phylogenetic positions (e.g. compare [Kear et al., 2006](#); [Benson et al., 2013](#); [Hampe, 2013](#); [Benson & Druckenmiller, 2014](#); [Sachs et al., 2016](#); [Serratos et al., 2017](#); [Fischer et al., 2018](#); [Páramo-Fonseca et al., 2019b](#); [O’Gorman, 2020](#)).

Only a few elasmosaurid taxa are known from the entire Early Cretaceous: *Lagenanectes richterae* Sachs, Hornung & Kear, 2017 from the Hauterivian–Barremian of Germany, the yet unnamed Speeton plesiosaurs (NHMUK PV R8623 and SCARB 200751) from the Hauterivian of the UK ([Benson & Druckenmiller, 2014](#)), *Callawayasaurus colombiensis* (Welles, 1962) from the Aptian of Colombia, *Leivanectes bernardoi* Páramo-Fonseca et al., 2019b from the Aptian of Colombia, *Wapuskianectes betsynichollsae* Druckenmiller & Russell, 2006 from the Albion of Canada and *Eromangasaurus australis* Kear, 2005 from the Albion of Australia ([Sachs, 2005a](#); [Kear, 2007](#)). All the other putative elasmosaurid specimens from the Berriasian–Hauterivian interval are undetermined postcranial remains, from Western Europe ([Fournier et al., 1982](#)), Russia ([Dubeikovskiy & Ochev, 1967](#); [Berezin & Aleksandrov, 2016](#); [Zverkov & Kiselev, 2018](#)), Argentina ([Lazo & Cichowolski, 2003](#); [O’Gorman et al., 2015a](#)) and Colombia ([Páramo-Fonseca, 2015](#)). Moreover, *La. richterae* was recovered as a basal elasmosaurid in two analyses ([Sachs et al., 2017](#); [Madzia & Cau, 2020](#)) and as a basal leptocleidid or basal xenopsarian in two others ([Sachs et al., 2018](#); [Páramo-Fonseca et al., 2019b](#), respectively), further demonstrating the need for additional data to document the early history of elasmosaurids better.

We describe a new taxon and two other specimens from the upper Hauterivian of European Russia. This new taxon is recovered as one of the earliest elasmosaurids and lacks several features that otherwise characterize the group. We also analyse the patterns of cervical elongation through the evolutionary history of elasmosaurids. The new taxon marks an early case of cervical elongation, which re-evolves only by the Late Cretaceous.

MATERIAL AND METHODS

INSTITUTIONAL ABBREVIATIONS

AMNH, American Museum of Natural History, New York, NY, USA; ANSP, Academy of Natural Sciences of Drexel University, Philadelphia, PA, USA; BGR, Bundesanstalt für Geowissenschaften und Rohstoffe, Hannover, Germany; GPMM, Geomuseum der

Universität Münster, Westfalen, Germany; KUV, Natural History Museum, University of Kansas, Lawrence, KS, USA; LACM, Natural History Museum of Los Angeles County, Los Angeles, CA, USA; MCS, Museo de Cinco Saltos, Río Negro Province, Argentina; MGUAN, Museu de Geologia da Universidade Agostinho Neto, Luanda, Angola; MLP, Museo de la Plata, Buenos Aires Province, Argentina; MOR, Museum of the Rockies, Bozeman, MT, USA; MOZ, Museo Juan Olsacher, Zapala, Neuquen Province, Argentina; NHMUK, Natural History Museum, London, UK; NZGS, New Zealand Geological Survey, Lower Hutt, New Zealand; RSM, Royal Saskatchewan Museum, Regina, Saskatchewan, Canada; SCARB, Rotunda Museum, Scarborough, UK; SGO, Museo Nacional de Historia Natural, Santiago, Chile; SMNK, Staatliches Museum für Naturkunde Karlsruhe, Germany; SMU SMP, Shuler Museum of Paleontology, Southern Methodist University, Dallas, TX, USA; SSU, Geological Museum, Saratov State University, Saratov, Russia; SU, Kagoshima Prefectural Museum, Shishijima, Japan; TMP, Royal Tyrrell Museum of Paleontology, Drumheller, Alberta, Canada; UCMP, University of California Museum of Paleontology, Berkeley, CA, USA; UPM, Undory Palaeontological museum, Undory, Ulyanovsk Region, Russia; YKM, Ulyanovsk Regional Museum of Local Lore named after I. A. Goncharov, Ulyanovsk, Ulyanovsk Region, Russia; YSPU, Ushinsky State Pedagogical University, Yaroslavl, Russia.

GEOGRAPHY AND STRATIGRAPHY

The right bank of the Volga River exposes Hauterivian to Barremian successions in the northern part of the Ulyanovsk district (European Russia). From the Slantsevy Rudnik village (formerly Zakharyevsky Rudnik) to the Polivno settlement (north of Ulyanovsk; [Fig. 1](#)), these successions are restricted to the upper Hauterivian–lower Barremian interval of the Klimovka Formation ([Baraboshkin & Guzhikov, 2015](#)). The Hauterivian deposits in the area have a thickness of ~50 m and consist of dark grey, poorly lithified shales, with occasional beds and lenses of siltite ([Blagovetshenskiy & Shumilkin, 2006a](#)). Several horizons contain large carbonate concretions ranging from 0.2 to 1.5 m in length; the shape of these concretions, colour of their matrix and the sequence of their diagenetic fracture filling can be used as stratigraphic markers, delineating several horizons across the succession. Despite a long history of bio- and lithostratigraphic studies initiated by [Pavlov \(1892\)](#), there is little consensus on the details of the biostratigraphic succession of the Lower Cretaceous successions along the Volga River, because a series of local schemes co-exists ([Baraboshkin, 2004](#);

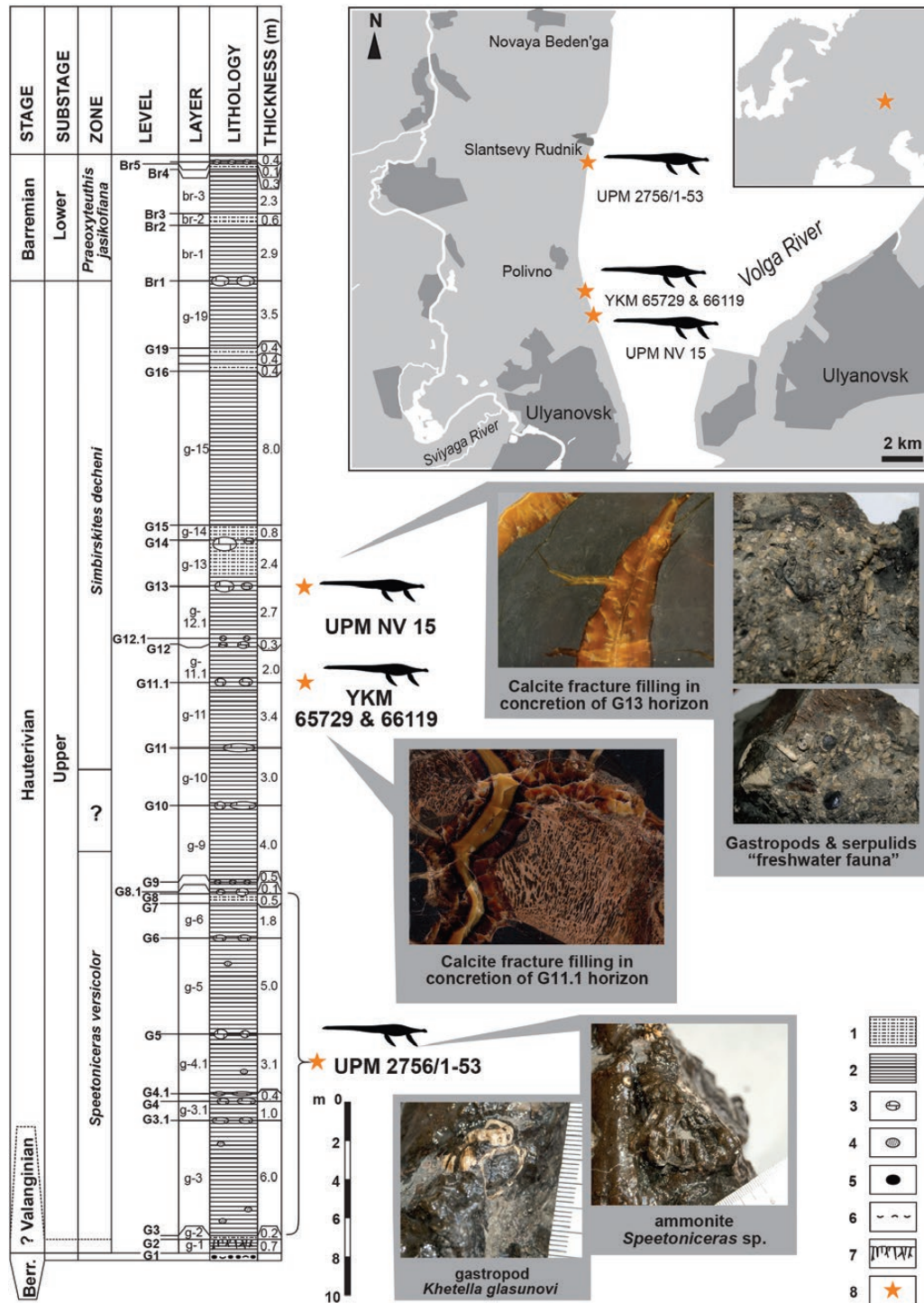


Figure 1. Spatiotemporal distribution of the specimens studied here and stratigraphic log of the relevant section of the Klimovka Formation, upper Hauterivian, Lower Cretaceous. Lithologies: 1, shale; 2, clay; 3, carbonate concretions; 4, pyrite concretions; 5, phosphorite nodules; 6, coquina bed; 7, bioturbated bed; 8, elasmosaurid remains.

Blagovetshenskiy & Shumilkin, 2006a, b; Baraboshkin & Blagoveschensky, 2010; Baraboshkin & Guzhikov, 2015, 2018). Based on our field observations (I.V.B. and I.M.S.) and layer-by-layer sampling of

invertebrates (I.V.B.), we follow the divisions proposed by Pavlow (1892, 1901), who recognized two ammonite biozones in the upper Hauterivian of the region: the *Speetonicerias versicolor* Zone in the lower part of the

Upper Hauterivian and the *Simbirskites decheni* Zone in the uppermost Hauterivian.

The specimens UPM NV 15 and YKM 65729 + 66119 were collected near the Polivno settlement (Fig. 1), and the specimen UPM 2756/1-53 was collected near the Slantsevy Rudnik village. All these specimens originate from the Klimovka Formation, but their precise position within the formation is unclear because they were collected from landslides, at water level. However, the diagenesis of the bones, the structure of the enclosing concretions, and the associated invertebrate fauna can be used to refine their stratigraphic provenance.

The concretions enclosing some elements of the specimen UPM NV 15 contain the typical gastropod assemblage of the so-called ‘freshwater fauna’ of Kabanov (1959), even though the palaeoenvironment was clearly marine (Baraboshkin et al., 2003). It includes gastropods *Crispotochus humilis* (Trautschold, 1865), *Hudlestoniella pusilla* (Tullberg, 1881), *Tornatellaea kabanovi* Blagovetshenskiy, 2017 and multiple serpulids of the genus *Ditrupa*. This combination of taxa and, in places, the abundance of invertebrate fossils are unique to the local stratigraphic intervals G13–G15, corresponding to the lower half of the *Simbirskites decheni* Zone (Blagovetshenskiy & Shumilkin, 2006a). Within this interval, the sequence of light orange then white calcite fracture filling is typical of G13 horizon, enclosed within the *Simbirskites decheni* Zone (Fig. 1).

The specimen YKM 65729 + 66119 is preserved in two large clayey-carbonate concretions of irregular shape. Although no invertebrate remains are found with this specimen, the diagenetic filling sequence of the fractures in the concretion by dark then yellowish calcite is characteristic of G11.1 horizon, corresponding to the basal part of the *Simbirskites decheni* Zone.

The specimen UPM 2756/1-53 is covered by an external layer of pyrite that fills inner pores and that fused with the bone surface, which is typical of the vertebrate remains from the *Speetonicerias versicolor* Zone in the region (Fischer et al., 2015, 2017; I. M. Stenshin, pers. obs.). A small ammonite, *Speetonicerias* sp., and a large specimen of the gastropod *Khetella Glazunov* Guzhov, 2004 are associated with the specimen (Fig. 1). *Khetella glazunovi* is abundant in the upper Hauterivian and present in both the *Speetonicerias versicolor* and the *Simbirskites decheni* zones. However, its specimens are commonly larger in size in the *Speetonicerias versicolor* Zone; the presence of an ammonite, *Speetonicerias* sp., associated with the specimen further supports this conclusion. This indicates that UPM 2756/1-53 belongs to the *Speetonicerias versicolor* Zone, like the holotype of the pliosaurid *Makhaira rossica* Fischer et al., 2015, which was found ~200 m north.

PHYLOGENETIC ANALYSIS

We used the dataset of O’Gorman (2020), which is the most complete dataset for elasmosaurids to date. We updated the matrix by adding two putative elasmosaurid operational taxonomic units (OTUs): *La. richterae*, complementing the scores of Sachs et al. (2017) (which we obtained from B. Kear, pers. comm. April 2020) and the new species, *Jucha squalea*. Our scores for *J. squalea* are based solely on the holotype specimen. We also changed two scores of *Er. australis* based on first-hand examination of the holotype by one of us (V. Fischer): 155:0&1 and 165:1. We also added the polycotylid and pliosaurid data stemming from the work of Benson & Druckenmiller (2014) and updated by Fischer et al. (2015, 2017, 2018) and Serratos et al. (2017), in addition to data from the literature on non-elasmosaurids (Hampe, 1992; Gasparini & O’Gorman, 2014; Páramo-Fonseca et al., 2016; Wintrich et al., 2017; Madzia et al., 2018; O’Gorman et al., 2018; Páramo-Fonseca et al., 2018, 2019a). O’Gorman (2020) added several elasmosaurid-focused characters (13) but did not score for them most non-elasmosaurid OTUs; likewise, the data on polycotylids and pliosaurids we added is scored as ‘?’ for these new characters, pending a thorough scoring of these new characters for non-elasmosaurids, which is not within the scope of this paper. The resulting dataset is a matrix of 131 taxa × 283 characters.

We then analysed this dataset in maximum parsimony using TNT v.1.5 (Goloboff & Catalano, 2016) with equal and implied weighting (with the default concavity constant, $k = 3$). In a maximum parsimony framework, implied weighting has been shown to yield clearly superior results, comparable to those of a Bayesian inference (Smith, 2019). In each case, we used the parsimony ratchet to carry out a rapid investigation of a series of shortest-tree islands (200 ratchet iterations and otherwise default options, with drift activated at ten iterations). The most parsimonious trees recovered by the parsimony ratchet were subjected to branch swapping using the tree bisection and reconnection (TBR) to maximize the recovery of most parsimonious trees; we set the maximal number of retained trees to 100 000. We provide our matrices containing the most parsimonious trees (one nexus file for the equal weighted analysis and one for the implied weighted analysis) and our TNT scripts as Supporting Information (supplementary files). We estimated the support for each clade using symmetric resampling, which is suitable for both unweighted and weighted parsimony analyses (Goloboff et al., 2003). We used a 33% symmetric change probability and 10 000 replicates, using the clades of the first most parsimonious tree as the reference.

We computed the strict and the 50% majority rule consensus trees in R using the packages *ape* v.5.3 (Paradis *et al.*, 2004) and *paleotree* v.3.3 (Bapst, 2012). We time-scaled the consensus tree using the ‘equal’ method (Brusatte *et al.*, 2008), using the *strap* v.1.4 package (Bell & Lloyd, 2015) and a table describing the geological range (or age uncertainties) of each taxon in the phylogeny. This table (see [Supporting Information, Table S1; supplementary files](#)) was updated from a previous version (Fischer *et al.*, 2018) using the literature (Otero, 2016; Sachs *et al.*, 2017; O’Gorman, 2020).

PATTERNS OF NECK ELONGATION

We gathered a series of measurements on the axial and appendicular skeletons of the three specimens described here (Tables 1 and 2). We assembled a dataset of anterior cervical centrum dimensions among elasmosaurids and basal leptocleidians. Although this approach does not use total neck lengths as in the study by Soul & Benson (2017), the taxonomic sampling is markedly increased, allowing incorporation of 29 elasmosaurids as opposed to between five and eight when complete necks are required. We obtained these data first hand (*Er. australis* and *J. squalea*) and from the literature (see Table 3). By convention, we focused our data on cervical centrum 10 (or the best-preserved centrum close to centrum 10), in order to investigate roughly similar regions of the anterior part of the neck. We gathered similar data on indeterminate Early Cretaceous elasmosaurids and on taxa not currently included in our phylogenetic dataset, such as ‘*Cimoliasaurus*’ and *Styxosaurus browni* Welles, 1943, in order to obtain a more complete overview of the cervical elongation patterns in elasmosaurids. Given that the cervical centra with the greatest elongation are distributed inconsistently within elasmosaurid necks, we also assembled a second dataset, wherein we used the cervical centrum with the highest length-to-height ratio, regardless of its position within the neck (Table 3).

We computed the length-to-height ratio of each specimen and used a time-scaled, randomly selected most parsimonious tree (see above) to infer ancestral (node) values in a maximum likelihood framework, using the *phytools* v.0.7-10 package (Revell, 2012). We used these data to create phenograms of cervical centrum elongation throughout the evolutionary history of elasmosaurids. We also used the data from Soul & Benson (2017) to evaluate the correlation between neck length and cervical elongation in long-necked (‘plesiosauiromorph’) plesiosaurians (Supporting Information, Table S2; supplementary files).

RESULTS

SYSTEMATIC DESCRIPTION

SAUROPTERYGIA OWEN, 1860

PLESIOSAURIA DE BLAINVILLE, 1835

XENOPSARIA BENSON & DRUCKENMILLER, 2014

ELASMOSAURIDAE COPE, 1869

JUCHA SQUALEA GEN. NOV., SP. NOV.

Zoobank registration: Publication LSID: urn:lsid:zoobank.org:pub:8CF4F7E1-7DE3-46DE-ABC3-34C4597DABCC

Genus LSID: urn:lsid:zoobank.org:act:A5A9FD2C-86CB-4CA5-A41A-22812277A226

Species LSID: urn:lsid:zoobank.org:act:8A9BDEF1-47B6-4616-8A70-59B505DBD8FA

Etymology

The generic name comes from the Cyrillic Юxa, a snake-like demon associated with water in local Volga–Ural region folklore. In English phonology, the generic name is pronounced ‘you kha’. The specific name is the Latin for coated, covered, rugose, referring to the pyrite layer found on many bones of the holotype.

Holotype

UPM 2756/1-53, a disarticulated partial skeleton discovered in 2007 by a field crew led by one of us (G.N.U.), comprising 22 cervical centra, 19 dorsal centra including two pectorals, one sacral and four caudals, two partial coracoids, two partial humeri, one radius, one radiale, several metacarpals and phalanges, two complete femora, one tibia, one fibula, fragmentary ribs and gastralia. This specimen originates from the *Speetonicerias versicolor* Zone (upper Hauterivian, Lower Cretaceous) in the vicinity of the Slantsevy Rudnik village (Fig. 1), Ulyanovsk Oblast, European Russia.

Diagnosis

Jucha squalea is characterized by the following autapomorphies among Elasmosauridae: (1) strongly waisted anterior to middle dorsal centra, giving the centrum an hourglass shape in ventral view; (2) massive distally-thickening transverse processes in middle to posterior dorsal vertebrae; (3) absence of a heart-shaped intercoracoid fenestra; and (4) large radius that is anteroposteriorly longer than the humeral head/capitulum.

Jucha squalea is also characterized by a unique combination of features, the most salient of which are as follows: (1) elongated anterior cervicals, with a length-to-height ratio ≤ 1.36 , similar to *Er. australis*

Table 1. Vertebral measurements (in millimetres) of *Jucha squalea* and cf. *Jucha*

| Specimen | Region | Length | Medial height | Width |
|---|-------------------|--------|---------------|-------|
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Cervical | 38 | 30 | 34 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Cervical | 58 | 43 | 51 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Cervical | 64 | 49 | 59 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Cervical | 70.5 | 52 | 67 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Cervical | 69 | 56 | 66 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Cervical | 70 | 56 | 69 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Cervical | 73 | 56 | 69 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Cervical | 78 | 61 | 76 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Cervical | 80 | 65 | 80 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Cervical | 79 | 68 | 81 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Cervical | 79 | 65 | 80 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Cervical | 82 | 70 | 82 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Cervical | 84 | 70 | 85 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Cervical | 84 | 72 | 87 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Cervical* | 79 | 71 | 87 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Cervical* | 83 | NA | 93 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Cervical* | 83 | 78 | 99 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Cervical* | 87 | NA | 102 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Cervical | 79 | NA | 100 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Cervical | 87 | 89 | 108 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Cervical | 85 | 89 | 108 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Cervical | 80 | 79 | 95 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Pectoral* | 75 | NA | 100 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Pectoral* | 73 | NA | 100 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Dorsal | 70 | NA | 100 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Dorsal* | 74 | NA | 108 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Dorsal* | 72 | NA | 115 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Dorsal* | ~35 | NA | 120 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Posterior dorsal* | 74 | 83 | 100 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Posterior dorsal* | 73 | NA | 89 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Posterior dorsal* | 70 | NA | 90 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Posterior dorsal* | 74 | NA | 99 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Posterior dorsal* | 72 | NA | 95 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Posterior dorsal* | 72 | NA | 99 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Posterior dorsal* | 65 | NA | 95 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Posterior dorsal* | 62 | NA | 89 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Posterior dorsal* | 59 | NA | 86 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Posterior dorsal* | 58 | NA | 91 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Posterior dorsal* | 61 | 72 | 89 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Posterior dorsal* | 58 | 69 | 79 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Posterior dorsal | 75 | NA | 81 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Sacral | 56 | 65 | 78 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Caudal* | 45 | 61 | 69 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Caudal* | 45 | NA | NA |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Caudal* | 42 | NA | 79 |
| cf. <i>Jucha</i> (UPM NV 15) | Cervical | 57.4 | 42.8 | 51.3 |
| cf. <i>Jucha</i> (UPM NV 15) | Cervical | 63 | 45.9 | 56.8 |
| cf. <i>Jucha</i> (UPM NV 15) | Cervical | 65.1 | 47 | 57.7 |

Table 1. Continued

| Specimen | Region | Length | Medial height | Width |
|------------------------------|-----------|--------|---------------|-------|
| cf. <i>Jucha</i> (UPM NV 15) | Cervical | 68.6 | 49 | 62 |
| cf. <i>Jucha</i> (UPM NV 15) | Cervical | 69 | 50 | 61 |
| cf. <i>Jucha</i> (UPM NV 15) | Cervical* | 70.2 | 51 | 62.2 |
| cf. <i>Jucha</i> (UPM NV 15) | Cervical* | 69.8 | 52 | 63.4 |
| cf. <i>Jucha</i> (UPM NV 15) | Cervical* | 72.5 | NA | 64.6 |
| cf. <i>Jucha</i> (UPM NV 15) | Cervical* | 72.4 | 55 | 67.3 |
| cf. <i>Jucha</i> (UPM NV 15) | Cervical* | 75.8 | NA | 69.9 |
| cf. <i>Jucha</i> (UPM NV 15) | Cervical* | 74.5 | 58.6 | 70.3 |
| cf. <i>Jucha</i> (UPM NV 15) | Cervical* | 75.2 | 59 | 71.5 |
| cf. <i>Jucha</i> (UPM NV 15) | Cervical | 74.6 | 59 | 72.5 |
| cf. <i>Jucha</i> (UPM NV 15) | Cervical* | 76.9 | 58.6 | 73.7 |
| cf. <i>Jucha</i> (UPM NV 15) | Cervical* | 82 | NA | NA |
| cf. <i>Jucha</i> (UPM NV 15) | Cervical* | 81 | NA | NA |
| cf. <i>Jucha</i> (UPM NV 15) | Cervical‡ | 81.5 | NA | NA |
| cf. <i>Jucha</i> (UPM NV 15) | Cervical‡ | 81 | NA | NA |
| cf. <i>Jucha</i> (UPM NV 15) | Cervical‡ | 81.5 | NA | NA |
| cf. <i>Jucha</i> (UPM NV 15) | Cervical* | 81.3 | 64.5 | NA |
| cf. <i>Jucha</i> (UPM NV 15) | Cervical* | 81 | NA | NA |
| cf. <i>Jucha</i> (UPM NV 15) | Cervical | 78.5 | 67.5 | 82.3 |
| cf. <i>Jucha</i> (UPM NV 15) | Cervical | 78.7 | NA | 82.7 |
| cf. <i>Jucha</i> (UPM NV 15) | Cervical* | 77.6 | NA | 84.4 |
| cf. <i>Jucha</i> (UPM NV 15) | Cervical* | 74.8 | NA | 84.7 |
| cf. <i>Jucha</i> (UPM NV 15) | Cervical | 71.5 | 68.9 | 87 |
| cf. <i>Jucha</i> (UPM NV 15) | Cervical | 74.4 | NA | 86.3 |
| cf. <i>Jucha</i> (UPM NV 15) | Cervical | 71 | NA | NA |
| cf. <i>Jucha</i> (UPM NV 15) | Caudal | 45.6 | 58.1 | 71 |
| cf. <i>Jucha</i> (UPM NV 15) | Caudal* | 46.2 | 55 | 63.6 |
| cf. <i>Jucha</i> (UPM NV 15) | Caudal* | 44.9 | 55.9 | 60.4 |

*(and, when needed, ‡): Articulated series. Missing data is indicated by “NA”.

and *Kawanectes lafquenianum* (Gasparini & Goñi 1985) (O’Gorman, 2016); (2) absence of ventral notch in all cervical centra, as in *La. richterae* (Sachs *et al.*, 2017), *Callawayasaurus colombiensis* (Welles, 1962), *Zarafasaura oceanis* Vincent *et al.*, 2011 (Lomax & Wahl, 2013; O’Gorman, 2020) and unlike *Er. australis* (V. Fischer, pers. obs.) and derived elasmosaurids (Welles, 1943, 1952; Otero, 2016; O’Gorman, 2020); (3) triangular, anteroposteriorly short anteromedial process of the coracoid, as in *Styxosaurus* (Welles & Bump, 1949; Welles, 1952; Otero, 2016), *Thalassomedon haningtoni* Welles, 1943 and *Nakonanectes bradti* Serratos *et al.*, 2017; (4) small, slit-like epipodal notch (also called radioulnar/tibiofibular foramen, spatium interosseum) in both the forelimb and the hindlimb, as in *Hydrotherosaurus alexandrae* Welles, 1943, and *Morenosaurus stocki* Welles, 1943; and (5) large tibia that is anteroposteriorly longer than femoral

head/capitulum, as in *Callawayasaurus colombiensis* (Welles, 1962).

Stratum typicum

Speetonicerus versicolor Zone of the Klimovka Formation, upper Hauterivian, Lower Cretaceous.

Locus typicus

Slantsevy Rudnik, Ulyanovsk Oblast, European Russia.

Description

The specimen UPM 2756/1-53 belongs to an osteologically mature individual, having neural arches fused with the centrum (Brown, 1981), elongated

Table 2. Measurements (in millimetres) of non-axial elements of *Jucha squalea* and cf. *Jucha*

| Specimen | Element | Anteroposterior length (proximally) | Anteroposterior length (distally) | Proximodistal length |
|---|----------------|-------------------------------------|-----------------------------------|----------------------|
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Humerus | 90 | NA | NA |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Radius | 125 | NA | 155 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Left femur | 94 | 178 | 270 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Right femur | 95 | 174 | 269 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Tibia | 117 | NA | 101 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Fibula | 95 | NA | 105 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Pedal phalange | 42 | NA | 64 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Pedal phalange | 38 | NA | 58 |
| <i>Jucha squalea</i> , holotype (UPM 2756/1-53) | Pedal phalange | 40 | NA | 63 |
| cf. <i>Jucha</i> (UPM NV 15) | Humerus | NA | 165 | 295 |

Missing data is indicated by “NA”.

propodials (O’Keefe & Chiappe, 2011) and finished bone surfaces on girdle elements.

Cervical vertebrae: At least 22 disarticulated cervicals are preserved in the holotype (Figs 2, 3), with no occurrence of the atlas and axis. One of these cervicals is a single, isolated anterior cervical (Fig. 3A–C; Table 1), suggesting that the neck comprised at least 24 elements (including the atlas and the axis), and probably many more. This anterior cervical centrum is markedly smaller (especially dorsoventrally) than the more posterior cervicals preserved (it is only 30 mm high and 38 mm long, compared with 43–49 mm high and 58–64 mm long for the next two smallest anterior cervicals; Table 1). Such a marked increase in centrum size along the neck is a frequent feature in derived elasmosaurids (O’Keefe & Hiller, 2006; O’Gorman *et al.*, 2015b; Sachs *et al.*, 2018) and suggests that *J. squalea* possessed several tens of cervical centra, although the precise number is impossible to estimate at present.

The cervical vertebrae are elongated; their length-to-height ratio is between 0.95 (posteriormost cervicals) and 1.36 (occurring in the anterior third of the neck). The anterior cervical centra are clearly elongated, approaching the ‘can-shaped’ condition (Fig. 3A–O; Tables 1 and 3). This shape resembles that of *Er. australis* and differs from many Early

Cretaceous elasmosaurids that have less elongated cervical centra, such as *Callawayasaurus colombiensis* (Welles, 1962) and *La. richterae* (Sachs *et al.*, 2017) (Table 3). The cervical centra of *J. squalea* are also more elongated than in *N. bradti* (Serratos *et al.*, 2017) and many weddellonectian euelasmosauridans for which this feature is known (Tables 3).

The articular surfaces are oval in outline, with their dorsal margin depressed under the neural canal (Fig. 3F, H, M). The ventral edge of the articular surface is continuously rounded, unlike derived elasmosaurids, in which a ventral notch gives the articular surface a binocular shape (O’Keefe, 2001); such a depression is also present but faint in *Er. australis* (V. Fischer, pers. obs. on holotype QMF11050). The articular surfaces are essentially platycoelous; they are amphicoelous in *La. richterae*, *Albertonectes vanderveldei* (Kubo *et al.*, 2012), *Libonectes morgani* (Welles, 1949) and *Th. haningtoni* (Kubo *et al.*, 2012; Sachs & Kear, 2015; Sachs *et al.*, 2017). The central part of the articular surface is slightly concave, lacking the central boss observed in the indeterminate Hauterivian elasmosaurid SSU 104-a/17 (Dubeikovsky & Ochev, 1967). In all of the preserved cervical centra, there are no ventral protrusions (lips), in contrast to SSU 104-a/17 (Dubeikovsky & Ochev, 1967) and possibly *La. richterae* (Sachs *et al.*, 2017).

Table 3. Anterior cervical measurements (height, length and width; in millimetres) and position for cervical centra at or close to the position of the tenth centrum (columns 4–7) and for the relatively longest cervical centrum record (columns 8–11) (for an R-friendly version, see [Supporting Information, supplementary files](#))

| Taxon | Specimen | Source | Height, centrum 10 | Length, centrum 10 | Width, centrum 10 | Position centrum | Height of longest centrum | Length of longest centrum | Width of longest centrum | Position of longest centrum |
|--------------------------------------|-------------------------|---|--------------------|--------------------|-------------------|------------------|---------------------------|---------------------------|--------------------------|-----------------------------|
| <i>Edgarosaurus muddi</i> | Holotype, MOR 751 | Druckenmiller (2002) , from figure | 42.8 | 34.3 | NA | 10 | 42.8 | 34.3 | NA | 10 |
| <i>Brancasaurus brancai</i> | Holotype, GPMM A3.B4 | Sachs et al. (2016) | 25 | 23 | NA | 10 | 25 | 23 | NA | 10 |
| Indet. <i>Californio</i> | LACM2832 | Welles (1952) ; O'Gorman (2020) | 22 | 27 | 41 | 12 | 26 | 37 | 51 | 20 |
| <i>Thalassomedon haningtoni</i> | Holotype, CMNH1588 | Welles (1952) | 50 | 56 | 68 | 10 | 81 | 123 | 102 | 35 |
| <i>Hydrotherosaurus alexandrae</i> | Holotype, UCMP33912 | Welles (1952) | 35 | 50 | 56 | 12 | 39 | 65 | NA | 20 |
| <i>Callawayasaurus colombiensis</i> | Holotype, UCMP38349 | Welles (1962) | 32 | 38 | 49 | 10 | 50 | 65 | 71 | 20 |
| <i>Cimoliasaurus maccoyi</i> | Holotype, AM F9644 | Kear (2002b) | 25.9 | 25.29 | 35.55 | 10 | 17.7 | 21.21 | 24.12 | 2 |
| <i>Eromangasaurus australis</i> | Holotype, QM F12219 | This paper (three-dimensional surface scan) | 50.55 | 69.3 | 70.76 | NA | 50.55 | 69.3 | 70.76 | NA |
| <i>Woolungasaurus glendowerensis</i> | Holotype, QM F6890 | Persson (1960) | 45 | 62 | 58 | NA | 33 | 50 | 47 | NA |
| Indet. Japan | SU01 | Utsunomiya (2019) | 25 | 35 | 38 | 10 | 25 | 35 | 38 | 10 |
| Hauterivian indet. Argentina 1 | MOZ PV 6893 | O'Gorman et al. (2015a) | 42 | 45 | NA | NA | 33 | 42 | 48 | NA |
| Valanginian indet. Argentina 1 | MOZ PV 6890 | O'Gorman et al. (2015a) | 47 | 59 | 52 | NA | 46 | 60 | 51 | NA |
| Hauterivian indet. Argentina 2 | MOZ PV 6991 | O'Gorman et al. (2015a) | 60 | 69 | NA | NA | 52 | 63 | NA | NA |
| Hauterivian indet. Russia | SSU 104-a/17-19 | Dubeikovskiy & Ochev (1967) | 51 | 62 | 57 | NA | 51 | 62 | 57 | NA |
| Valanginian indet. Russia | YSPU1896-32 | Zverkov & Kiselev (2018) | 67.1 | 94.6 | 83 | NA | 67.1 | 94.6 | 83 | NA |
| <i>Libonectes morgani</i> | Holotype, SMU SMP 69120 | Welles (1949) | 40 | 52 | 65 | 13 | 55 | 95 | 93 | 30 |
| <i>Libonectes atlansense</i> | SMNK-PAL-3978 | Buchy (2005) | NA | 42 | NA | 10 | NA | 42 | NA | 10 |
| <i>Lagenanectes richterae</i> | Holotype, BGR Ma 13328 | Sachs et al. (2017) | 43 | 45.72 | 45.53 | NA | 49.83 | 59.13 | 53.23 | NA |

Table 3. Continued

| Taxon | Specimen | Source | Height, centrum 10 | Length, centrum 10 | Width, centrum 10 | Position | Height of longest centrum | Length of longest centrum | Width of longest centrum | Position of longest centrum |
|---|------------------------------|--|--------------------------|--------------------------|-------------------------|----------|---------------------------------|---------------------------------|--------------------------------|--------------------------------------|
| <i>Styxosaurus</i> sp. (‘ <i>Hydralmosaurus</i> ’) | AMNH1495 | Otero (2016) | 40.65 | 56.78 | 55.24 | 10 | 44.98 | 70.44 | 62.83 | 14 |
| <i>Styxosaurus browni</i> | Holotype, AMNH5835 | Otero (2016) | 32.23 | 58.05 | 51.58 | 10 | 32.23 | 58.05 | 51.58 | 10 |
| <i>Styxosaurus snowii</i> | Holotype, KUVF1301 | Welles (1952); Otero (2016) | 60 | 78 | NA | 6 | 60 | 78 | NA | 6 |
| <i>Cimoliasaurus magnus</i> | AMNH2554 | Otero (2016) | 84.04 | 75.55 | 111.84 | 9 | 84.04 | 75.55 | 111.84 | 9 |
| <i>Albertonectes</i> | Holotype, TMP2007.011.001 | Kubo <i>et al.</i> (2012) | 45 | 70 | 57 | 25 | 23 | 38 | 31 | 3 |
| <i>Elasmosaurus</i> | Holotype, ANSP 10081 | Sachs (2005b) | 31 | 51 | 43 | 10 | 54 | 93 | 65 | 32 |
| <i>platyrus</i> | | | | | | | | | | |
| <i>Vegasaurus molyi</i> | Holotype, MLP 93-I-5-1 | O’Gorman <i>et al.</i> (2015b) | 27 | 32 | 38 | 10 | 39 | 51 | 55 | 20 |
| <i>Nakonanectes bradti</i> | Holotype, MOR3072 | Serratos <i>et al.</i> (2017) | 30.4 | 34.7 | 48 | 10 | 31.4 | 40.2 | 51.6 | 13 |
| <i>Kawanectes</i> | MCS PV4 ‘C2’ | O’Gorman (2016) | 33 | 45 | 50 | NA | 33 | 45 | 50 | NA |
| <i>lafquenianum</i> | | | | | | | | | | |
| <i>Tuarangisaurus keyesi</i> | NZGS, CD426 | Wiffen & Moiseley (1986) | 32 | 42 | 43 | 8 | 30 | 40 | 41 | 5 |
| <i>Aristonectes parvidens</i> | Holotype, MLP 40-XI-14-6 | Gasparini <i>et al.</i> (2003) | 53 | 60 | 96 | NA | 53 | 60 | 96 | NA |
| <i>Aristonectes</i> | Holotype, SGO.PV.957 | Otero <i>et al.</i> (2014) | 57.5 | 56 | 75.3 | 10 | 57.5 | 56 | 75.3 | 10 |
| <i>quiriquinensis</i> | | | | | | | | | | |
| <i>Cardiocorax mukulu</i> | Holotype, MGUAN PA103 | Araújo <i>et al.</i> (2015a), from figure | 35.7 | 53.1 | 59.6 | NA | 35.7 | 53.1 | 59.6 | NA |
| <i>Terminonator</i> | Holotype, RSM P2414.1 | Sato (2003) | 37 | 62 | 55 | 8 | 24 | 43 | 39 | 3 |
| <i>ponteixensis</i> | | | | | | | | | | |
| <i>Jucha squalea</i> | Holotype, UPM 2756/1-53 | This paper | 52 | 70.5 | 67 | NA | 52 | 70.5 | 67 | NA |
| cf. <i>Jucha</i> | UPM NV 15 | This paper | 49 | 68.6 | 62 | NA | 49 | 68.6 | 62 | NA |

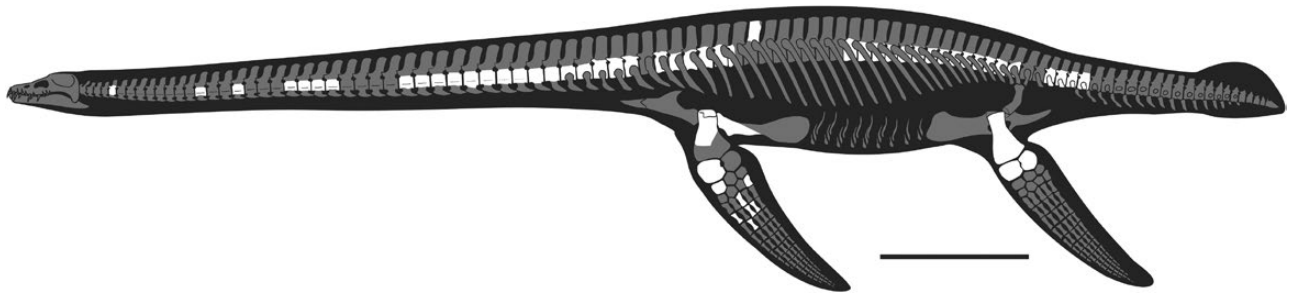


Figure 2. Skeletal reconstruction of the holotype specimen of *Jucha squalea* (UPM 2756/1-53). Preserved bones are coloured in white; the remainder of the osteology is based on *Callawayasaurus colombiensis*, *Thalassomedon haningtoni* and *Hydrotherosaurus alexandrae* (Welles, 1943, 1962).

A consistent feature of *J. squalea* is the presence of a strongly rugose peripheral band adjacent to the articular surface of cervical centra, forming irregular, anteroposteriorly oriented ridges and furrows on the lateral and, to a lesser degree, ventral surfaces of the centrum (Fig. 3G, K, L, N). A peripheral rugosity in vertebral centra is variably present in many adult plesiosauroidea (e.g. Owen, 1840; Seeley, 1874), including other Hauterivian specimens from France (Fournier *et al.*, 1982) and the UK (the Speeton plesiosaur; N. G. Zverkov, pers. obs. on NHMUK PV R8623, April 2019), but it appears much more pronounced and consistently restricted to all cervical–pectoral centra. In contrast, rugosities appear entirely absent in SSU 104-a/17 (Dubeikovskiy & Ochev, 1967) and *La. richterae* (Sachs *et al.*, 2017) and are present but faint in *Callawayasaurus colombiensis* (Welles, 1962).

Cervical centra are slightly waisted. Paired oval foramina are present on the ventral side and are separated by a rounded median keel, as in all elasmosaurids. The cervical rib facets are lozenge shaped anteriorly and become rounded posteriorly (Fig. 3G, N, P). A sharp lateral, anteroposteriorly oriented ridge [a feature frequently evolved convergently among long-necked plesiosauroidea (e.g. Noè, *et al.*, 2017; Fischer *et al.*, 2018)] is present immediately dorsal to the rib facet in anterior and middle cervical centra (Fig. 3A, G, K, N). Posterior cervicals lack this feature (Fig. 3P). The prezygapophyses are mediolaterally narrow and face dorsomedially. They are separated from one another from at least one-third of their length (Fig. 3O). No neural spine is preserved. All the preserved neural arches are fully fused to their corresponding centra; the suture is V-shaped in lateral view (Fig. 3K).

Pectoral and dorsal vertebrae: Four and a half vertebrae from the pectoral to anterior dorsal region are preserved (Fig. 3R–Z), in addition to a complete and articulated mid-dorsal to sacral series (Fig. 4).

Anterior to middle dorsal centra are markedly waisted transversely, giving the centrum an hourglass shape in ventral view (Figs 3R, X, 4C, F). Pectoral centra are wider than high, and the dorsal centra become progressively as wide as high and then slightly higher than wide throughout the dorsal series.

Another peculiar feature of *J. squalea* is the robust, distally thickening transverse processes in the middle to posterior dorsal region (Fig. 4A, C). The orientation of the long axis of the distal end of these processes varies throughout the dorsal series; it is oriented dorsoventrally in the pectorals (Fig. 3W) and almost horizontally in the midposterior dorsals (Fig. 4B), before progressively orienting posteroventrally in the posteriormost dorsals (Fig. 4H). This condition somewhat resembles that of the dorsoventrally flattened *Tatenectes laramiensis* (Knight, 1900) (O’Keefe *et al.*, 2011). The transverse processes in pectorals and anterior dorsals have a shallow ventral groove, giving the process an ‘A’-shaped cross-section (Fig. 3U); a similarly placed but much deeper ‘subdiapophyseal fossa’ has been reported in ‘*Gronausaurus wegneri*’ Hampe, 2013 [= *Brancaasaurus brancai* according to Sachs *et al.* (2016)]. In the posteriormost dorsals, the transverse processes abruptly become compressed, with a rectangular cross-section.

These posterior dorsal centra are also not waisted, unlike the previous ones, which have the autapomorphic hourglass shape in ventral view (Fig. 4F). Paired ventral foramina are present and positioned ventrolaterally throughout the dorsal series. The prezygapophyses are strongly concave, with a dorsomedial concavity; they progressively separate from one another throughout the dorsal series, becoming separated over at least one-half of their length in posterior dorsals (Fig. 4C).

An isolated, fragmentary neural spine is present; this fragment is 100 mm high and mediolaterally thickens dorsally, suggesting that it originates from the dorsal region. This fragment indicates that dorsal neural spines were substantially higher than their

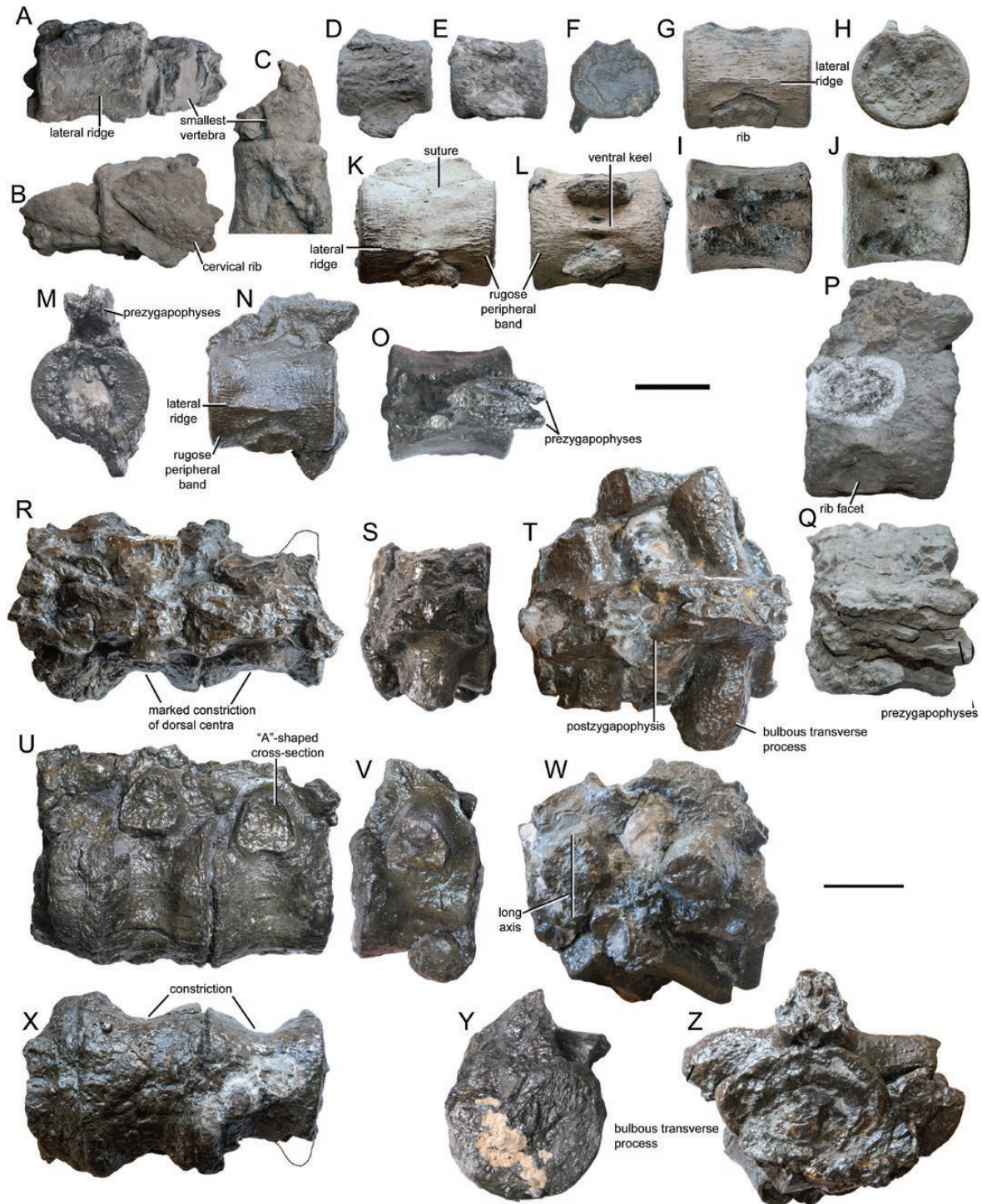


Figure 3. Cervical, pectoral and anterior dorsal vertebrae of *Jucha squalea* (UPM 2756/1-53). A–C, smallest preserved anterior cervical vertebrae in lateral view (A, B) and articular surface of the smallest preserved vertebra (C). D–O, anterior to middle cervical vertebrae in lateral (D, G, K, N), ventral (E, L, J), articular (F, H, M) and dorsal (I, O) views. P, Q, posterior cervical vertebra in right lateral (P) and dorsal (Q) views. R–Z, pectoral to anterior dorsal vertebrae in dorsal (R–T), right lateral (U–W), ventral (X) and articular (Y, Z) views. Scale bars: 50 mm.

corresponding centra, because none of the preserved centra exceeds 82 mm in dorsoventral height. A similar condition is present in one of the Speeton plesiosaurs, NHMUK PV R8623 (N. G. Zverkov, pers. obs. April 2019), and in an indeterminate Hauterivian specimen from France (Fournier *et al.*, 1982).

Sacral and caudal vertebrae: One sacral vertebra is preserved. The base of the transverse process is large again, and its long axis is vertical (Fig. 4K). The sacral centrum is similar to the posteriormost dorsals in being unwaisted ventrally. Paired foramina are present and positioned ventrolaterally.

A series of four caudal vertebrae with articulated ribs and neural arches is preserved. The caudal vertebrae are anteroposteriorly short (Fig. 4M, P). The articular surfaces are oval to subhexagonal in outline, with the width exceeding the height (Fig. 4N). The caudal ribs are thick and become dorsoventrally compressed posteriorly.

Ribs: Anterior cervical ribs have a lozenge-shaped basal cross-section (Fig. 3B, D, L); their cross-section becomes progressively oval, then circular along the cervical series. Distally, the anterior cervical rib forms a small, pointed, proximally placed anterior process and a longer posterodistal process (Fig. 3B), resembling those of *Futabasaurus suzuki* Sato, Hasegawa, & Manabe, 2006 and, possibly, *Callawayasaurus colombiensis* (Welles, 1962). This differs from elasmosaurids that have 'hatchet'-shaped cervical ribs, where the anterior and posterior processes are placed distally [e.g. *Albertonectes vanderveldei* (Kubo *et al.*, 2012), *Styxosaurus browni* (Otero *et al.*, 2016) and *Elasmosaurus platyrus* Cope, 1868 (Sachs, 2005b)].

Coracoid: Both coracoids are preserved in connection but are fragmentary (Fig. 5A–F). The coracoid symphysis is thickened ventrally (215:1) and slightly thickened dorsally, giving it an eye shape in medial view, with a median ventral protrusion (Fig. 5D). This ventral protrusion is moderately pronounced, as in *Callawayasaurus colombiensis* (Welles, 1962), and unlike the extremely protruding process of *Wa. betsynichollsae* (Druckenmiller & Russell, 2006) and *Li. morgani* (Sachs & Kear, 2017). The symphysis is undulating in ventral view and is bordered by thin anteroposterior ridges texturing the bone surface. The dorsal and ventral surfaces of the coracoid are flattened, and the dorsal surface essentially lacks a mediolateral buttress (Fig. 5D, E). A slight swelling is present on the anterior portion but does not result in an anterior depression; therefore, we scored this character as 214:2.

The anteromedial process is preserved; its anterior and medial surfaces are thickened and concave, whereas its anterolateral edge is thin and sheet-like.

The anteromedial process is anteroposteriorly short and triangular in outline; the medial surfaces of the anteromedial processes are divergent, and the angle formed by the medial and anterolateral margins of the anteromedial process is ~65° in *J. squala* (Fig. 5A, E). *Styxosaurus* spp., *Th. haningtoni* and *H. alexandrae* have an angle >50° (Welles, 1943, 1952; Welles & Bump, 1949), whereas it is only ~40° in *Callawayasaurus* and *Aphrosaurus furlongi* Welles, 1943 (Welles, 1943 (Welles, 1962, O'Gorman, 2020) and ~35° in *Wa. betsynichollsae* (Druckenmiller & Russell, 2006). The anteromedial process is located close to the ventral protrusion, indicating a weak development of the pectoral bar (Fig. 5A, D). This condition resembles that of *Styxosaurus* (Welles & Bump, 1949; Welles, 1952; Otero, 2016), *Th. haningtoni* (Welles, 1943) and, to a certain degree, *N. bradti* (Serratos *et al.*, 2017) but differs from many other elasmosaurids, whose anteromedial processes protrude far more anteriorly (Welles, 1962; Druckenmiller & Russell, 2006; Araújo *et al.*, 2015a; Sachs *et al.*, 2017; O'Gorman, 2020).

The medial surfaces of the coracoids diverge gradually posterior to the symphysis (Fig. 5A); the preserved portions of the coracoids suggest that *J. squala* lacks a posteromedial process, which forms the heart-shaped intercoracoid cavity seen in all known elasmosaurids (e.g. Welles, 1943, 1952, 1962; Druckenmiller & Russell, 2006; Otero, 2016; O'Gorman, 2020). The angle between the scapular and glenoid facets is ~150°; they are poorly demarcated and have an irregularly papillose surface (Fig. 3B, C). The coracoid of *J. squala* thus lacks several features of other known basal elasmosaurids.

Humerus: Only the proximal parts of both humeri are preserved. The humerus appears robust (Fig. 5G). Its anteroposterior width rapidly increases distally from the proximal end, resulting in a basically unwaisted shaft, resembling that of *F. suzuki* (Sato *et al.*, 2006) and unlike that of most other elasmosaurids (e.g. Welles, 1943, 1952, 1962; Otero, 2016; O'Gorman, 2020). In this aspect, the humerus exhibits similarities to some aristonectines that have paedomorphic limbs (e.g. Araújo *et al.*, 2015b), although the limbs of *J. squala* do not show any marked evidence of a delayed or slowed ossification. The humeral head (capitulum) and dorsal tuberosity are not separated by the band of periosteal bone, unlike *H. alexandrae* and *Mo. stocki* (Welles, 1952). Their surfaces are flattened and irregularly papillate, indicating the presence of an extensive cartilaginous cap *in vivo*. The dorsal tuberosity is as wide anteroposteriorly as the humeral head and is deflected postaxially (Fig. 5K).

Radius: The radius of *J. squala* is unique in being large, markedly longer anteroposteriorly than the head of the humerus. The humeral facet is oval,

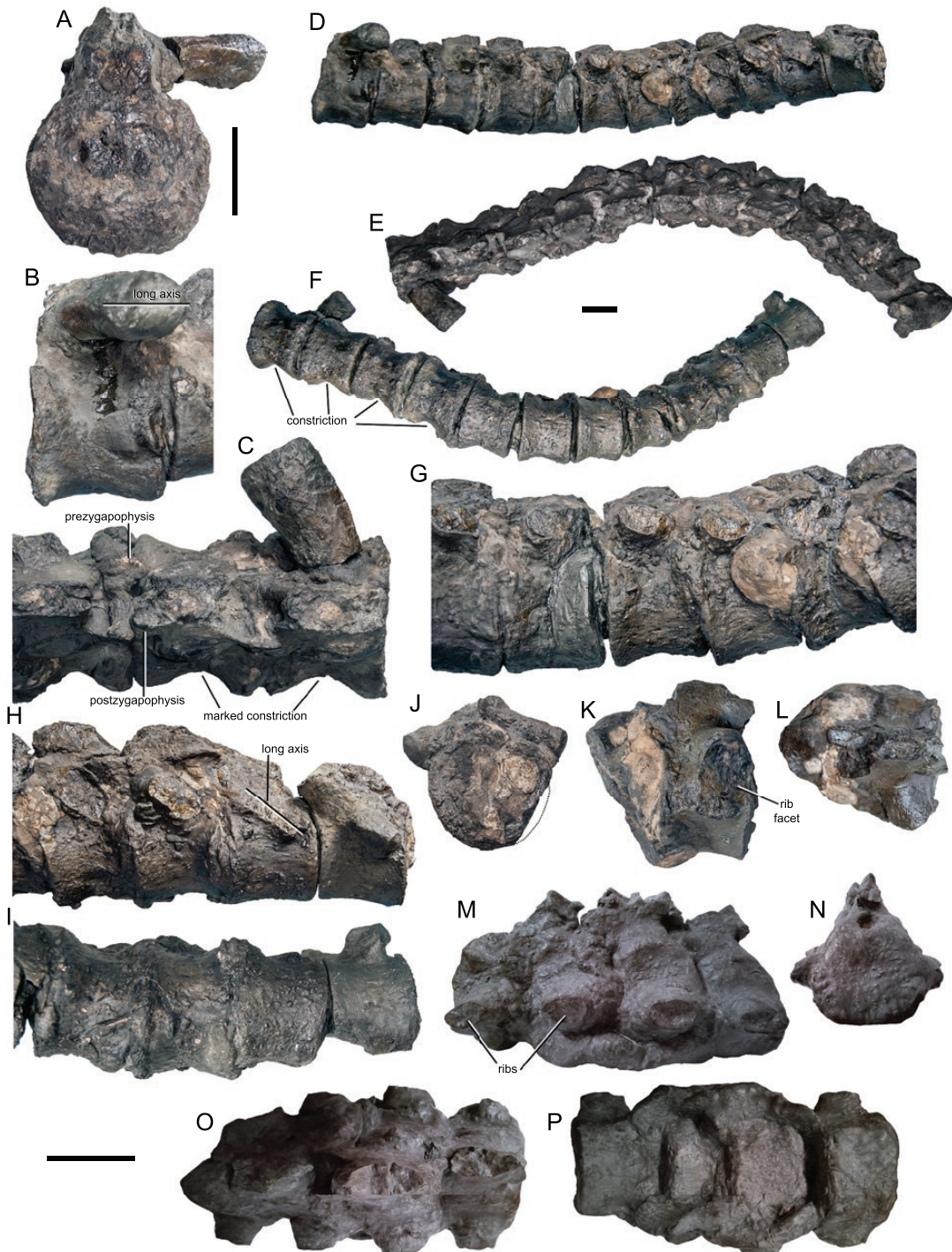


Figure 4. Dorsal, sacral and caudal vertebrae of *Jucha squalea* (UPM 2756/1-53). A–C, middle dorsal vertebra in anterior (A), left lateral (B) and dorsal (C) views. D–F, articulated middle to posterior dorsal vertebrae in left lateral (D), dorsal (E) and ventral (F) views. G, H, magnified regions of (A), dorsal vertebra in lateral view. I, J, posterior-most dorsal vertebrae in ventral (I) and posterior (J) views. K, L, sacral vertebra in left lateral (K) and dorsal (L) views. M–P, articulated caudal vertebrae in left lateral (M), posterior articular (N), dorsal (O) and ventral (P) views. Scale bars: 50 mm.

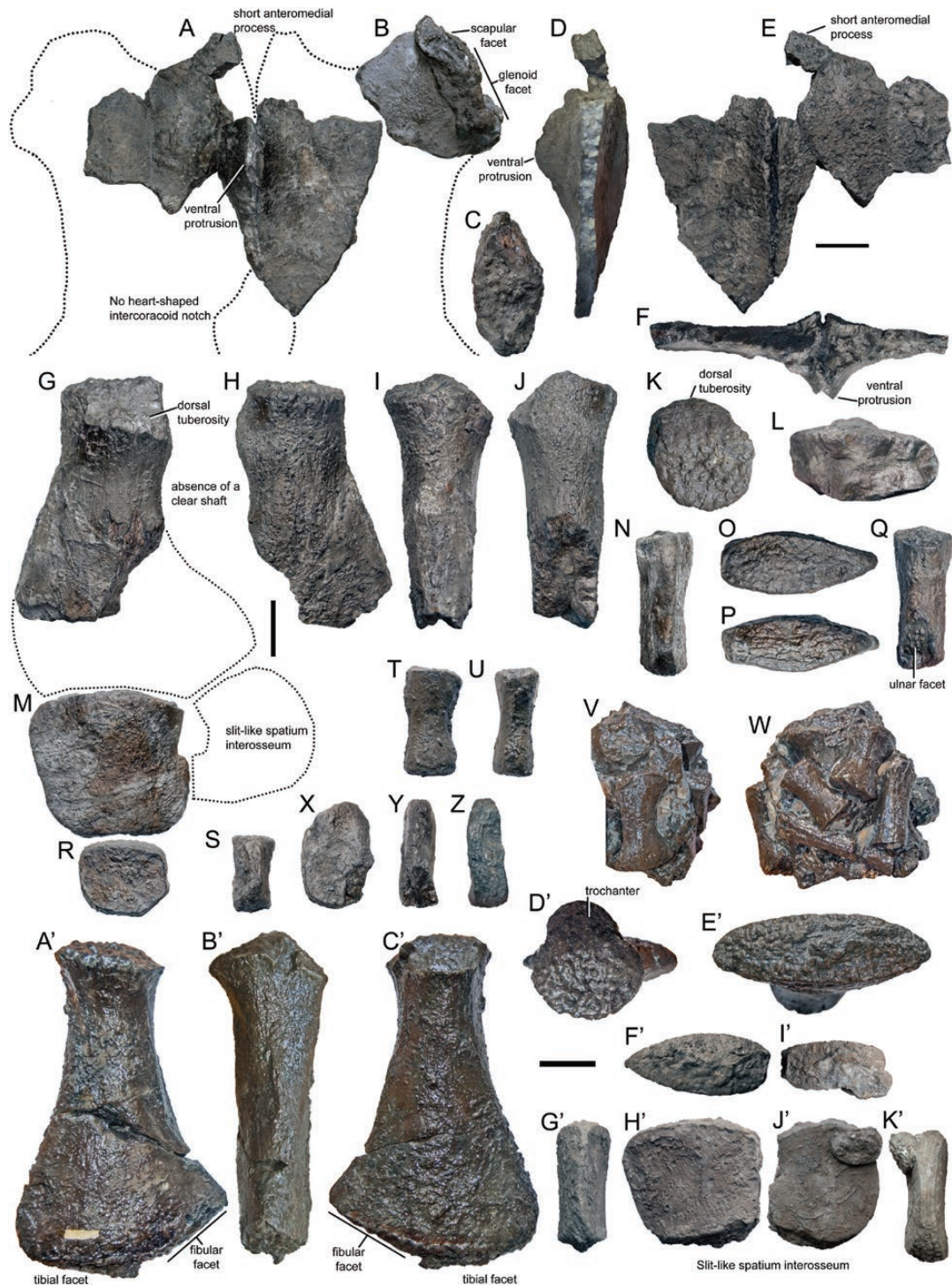


Figure 5. Appendicular skeleton of *Jucha squalea* (UPM 2756/1-53). A–F, articulated coracoids in ventral (A, B), lateral (D), dorsal (E) and anterior (F) views; C, articular view of glenoid portion. G–L, partial left humerus in dorsal (G), ventral (H), anterior (I), posterior (J), proximal (K) and distal (L) views. M–Q, radius in dorsal (M), anterior (N), proximal (O), distal (P) and posterior (Q) views. R, S, radiale in dorsal (R) and anterior (S) views. T, U, metacarpal or proximal phalanx. V, W, phalanges; X–Z, fifth metacarpal in dorsal (X), posterior (Y) and anterior (Z) views. A'–E', right femur in ventral (A'), posterior (B'), dorsal (C'), proximal (D') and distal (E') views. F'–H', left tibia in proximal (F'), anterior (G') and dorsal (H') views. I'–K', left fibula in proximal (I'), dorsal (J') and posterior (K') views. Scale bars: 50 mm.

rugose and slightly convex (Fig. 5M). The anterior edge is straight and anteriorly tapering and is made of finished bone (Fig. 5N). The posterior facet of the radius is bifid, because of the presence of a median concavity forming the anterior edge of the radioulnar foramen (Fig. 5M, Q). A radial notch is thus present, but the contribution of the radius to the radioulnar foramen is small, as in *H. alexandrae*, *Mo. stocki* and *Aphrosaurus furlongi* (Welles, 1952; O’Gorman, 2020) but unlike ‘*Gronausaurus wegneri*’ (Hampe, 2013), *Callawayasaurus colombiensis* (Welles, 1962) and several derived elasmosaurids (Welles, 1949, 1952; Sato *et al.*, 2006; Otero *et al.*, 2014; Hiller *et al.*, 2017). Proximally, the articulation with the ulna is barely noticeable and possibly absent, unlike ‘*Gronausaurus wegneri*’ [= *Brancausaurus brancai* according to Sachs *et al.* (2016)] and many elasmosaurids (Welles, 1962; Hampe, 2013; O’Gorman, 2020). The radius simply forms a flattened, oblique surface (Fig. 5Q). On the contrary, the distal ulnar facet is prominent and bordered by a sharp, raised edge. A facet for the intermedium is present, but it is markedly smaller than the radiale facet and is poorly demarcated from it; this condition (character state 263.2) appears rare, being restricted to some aristonectines and *J. squalea* (Otero *et al.*, 2014; Araújo *et al.*, 2015b).

Radiale: The radiale is thick and anteriorly tapering and possesses four articular facets for the humerus, intermedium, distal carpal 2 + 3 and distal carpal 1 (Fig. 5R). The radiale is much smaller than the radius (anteroposterior length of 77 mm, compared with 125 mm for the radius). Contrary to the radius, the anterior surface of the radiale lacks finished bone (Fig. 5S).

Metacarpal V: A proximodistally elongated, ear-shaped element is interpreted here as the metacarpal V. This tapers posteriorly and possesses a small posterior notch (Fig. 5X, Y). The distal facet is flat and thick, whereas the ulnare facet is triangular and anteromedially facing.

Femur: The femur is short and robust. Like the humerus, the femur is not waisted; it starts expanding anteroposteriorly distal to the first third of total femoral length (Fig. 5A’, C’). This condition is similar to *Callawayasaurus colombiensis* (Welles, 1962; O’Gorman, 2020) and *Mo. stocki* (Welles, 1952) and differs from the longer shafts seen in ‘*Gronausaurus wegneri*’ (Hampe, 2013), *La. richterae* (Sachs *et al.*, 2017), *Th. haningtoni*, *H. alexandrae*, *Aphrosaurus furlongi* (Welles, 1952; O’Gorman, 2020), *F. suzuki* (Sato *et al.*, 2006), ‘*Woolungasaurus glendowerensis*’ (Elasmosauridae indet.) (Persson, 1960; Sachs, 2004)

and a Hauterivian specimen from France (Fournier *et al.*, 1982); the condition appears somewhat variable in *Styxosaurus* spp. (Welles, 1952; Otero, 2016). The femur appears straight in anteroposterior and dorsoventral views and is therefore not sigmoid (Fig. 5A’–C’), although a slight diagenetic flattening cannot be ruled out. The long axis of the femur is slightly deflected posterodistally, but a marked postaxial deflexion is absent, unlike the femur in *Styxosaurus* spp., where the posterior expansion is longer (Welles, 1952, 1962). This condition appears similar to that of *Callawayasaurus colombiensis* (Welles, 1962; O’Gorman, 2020), although the anterodistal surface is less rounded than in *J. squalea*. The dorsal trochanter is thick, semicircular in cross-section, and slightly narrower (anteroposteriorly) than the femoral head (Fig. 5D’). There is no separation of the dorsal trochanter from the femoral head by periosteal bone, unlike in *Mo. stocki* (Welles, 1952) and, to smaller extent, *Callawayasaurus colombiensis* (Welles, 1962). The distal surface is rounded in dorsoventral view, making the distal facets hardly discernible, as in *Callawayasaurus colombiensis* (Welles, 1962).

Tibia: The tibia resembles the radius in being anteroposteriorly longer than its corresponding propodial proximal head; this condition is shared only with *Callawayasaurus colombiensis* (Welles, 1962). The tibia is large and bulky; a posterior notch is present, but it is so small that it is invisible in dorsoventral view (Fig. 5H’). This condition again resembles that of *Callawayasaurus colombiensis* (Welles, 1962), although the tibial contribution appears even more reduced in *J. squalea*.

Fibula: The fibula is rounded and slightly smaller than the tibia (Fig. 5I’–K’). The fibula possesses a small anterior notch, unlike the fibula in *Th. haningtoni*, *Aphrosaurus furlongi* and *Styxosaurus* sp. (Welles, 1962; Otero, 2016; O’Gorman, 2020). As a result, the tibioulnar foramen is small and slit-like, resembling but smaller than those of *Callawayasaurus colombiensis*, *H. alexandrae* and *Mo. stocki* (Welles, 1943, 1962), in addition to that of an indeterminate Hauterivian elasmosaurid from France (Fournier *et al.*, 1982).

CF. JUCHA

Referred specimens

UPM NV 15, a fragmentary disarticulated skeleton comprising 29 cervicals, at least three caudals, cervical and dorsal ribs, a partial left humerus, one epipodial element and several phalanges. This specimen originates from the *Simbirskites decheni*

Zone (upper Hauterivian, Lower Cretaceous) of the Polivno locality (Fig. 1), Ulyanovsk Oblast, European Russia.

YKM 65729 + 66119, a fragmentary disarticulated skeleton comprising one cervical, ?three pectorals, 17 dorsal and two caudal vertebrae, many ribs, one partial femur, ischium and pubis. This specimen also originates from the *Simbirskites decheni* Zone (upper Hauterivian, Lower Cretaceous) of the Polivno locality (Fig. 1), Ulyanovsk Oblast, European Russia.

Preliminary note

Although these specimens are generally similar and compatible with the holotype of *J. squalia* (in addition to being spatiotemporally close), their incompleteness and poor overlap with the holotype does not allow an unambiguous referral to the same species. We therefore describe them as cf. *Jucha*, focusing our efforts on the differences from the holotype of *J. squalia*.

Description

Cervical vertebrae: A total of 29 partly articulated cervical centra are preserved in UPM NV 15, including some from the anterior half of the neck. The anterior cervical centra of UPM NV 15 are elongated, with a length-to-height ratio reaching 1.4. Peripheral ridges and furrows appear more rugose than in the holotype of *J. squalia*, forming a pitted texture (Fig. 6A, E). The lateral ridge is either absent or present but is fainter and anteroposteriorly shorter in UPM NV 15 than in the holotype of *J. squalia*. A partial cervical rib is preserved in UPM NV 15 (Fig. 6I–K). It bears several mediolateral ridges on its proximal part and has a longer shaft than in the ribs preserved in the holotype of *J. squalia*.

Pectoral and dorsal vertebrae and ribs: Three pectorals are preserved in YKM 65729 + 66119. These centra also possess the rugose peripheral band, although it appears less conspicuous than in the cervical centra of the other specimens (Fig. 7G). The dorsal neural spine is longer than the dorsoventral height of their corresponding centrum (125 vs. 79 mm; Fig. 7A, I). The dorsal surface of the dorsal neural spines is not expanded and bears a flat surface that is convex in lateral view (Fig. 7B, I, K). Dorsal transverse processes of YKM 65729 + 66119 appear more slender than in the holotype of *J. squalia* and are markedly inclined dorsolaterally (Fig. 7B, I, J); their rib facets have an oval outline, with their long axis set almost vertically. The dorsal ribs are thick, with an oval cross-section. They have a weakly sigmoidal profile in anteroposterior view: medially, the dorsal surface curves dorsolaterally and then curves ventrolaterally, as in *Albertonectes*

vanderveldei and unlike '*Gronausaurus wegneri*' (Kubo *et al.*, 2012; Hampe, 2013). The ribs of UPM NV 15 bear several deep longitudinal ridges proximally but lack a posterior sulcus, unlike the ribs of '*Gronausaurus wegneri*' (Hampe, 2013).

Sacral and caudal vertebrae and ribs: Two vertebrae of YKM 65729 + 66119 are interpreted as sacrals (but might be anteriormost caudal, lacking chevron facets). These vertebrae are anteroposteriorly short and are preserved with their articulated ribs (Fig. 7B, N). The ventral surface is flat to slightly concave and is rugose. The rib facets are large and oval in outline, deeply concave and occupying most of the dorsoventral height of the centrum (Fig. 7M). The associated ribs are robust and expand distally; they bear a protruding, posteroventrally projecting median ventral processes (Fig. 7N).

Three caudal vertebrae are preserved in UPM NV 15. Their centra are anteroposteriorly short. The chevron facets are semi-oval and contact the posterior edge of the centrum, whereas there are no traces of chevrons on the anterior edge (Fig. 6P, U). No ventral or lateral keel is present.

Forefin: A nearly complete left humerus is preserved UPM NV 15, in addition to a fragmentary epipodial element and numerous phalanges (Fig. 8A–K). The humerus differs slightly from that of the holotype of *J. squalia* in having a longer shaft and a more pronounced waist. The anterior surface of the humerus is essentially straight, whereas the posterior margin is strongly convex (Fig. 8A, C); the humerus appears slightly sigmoidal, bearing similarities to those of *Callawayasaurus colombiensis* and *Mo. stocki* (Welles, 1962). The radial facet faces distally, whereas the ulnar facet is deflected, facing posterodistally. This condition also resembles those of *Callawayasaurus colombiensis* and *Mo. stocki* (Welles, 1962). The dorsal tuberosity is shifted postaxially, as in the holotype of *J. squalia*. The fragmentary epipodial element lacks its posterior and distal surfaces; therefore, its absolute and relative sizes are unknown (Fig. 8F). The contribution to the radioulnar foramen is preserved, but it is difficult to assess its shape unambiguously. The proximal surface is convex and oval in outline (Fig. 8G). Phalanges are elongated and oval in cross-section.

Pubis: A complete pubis is preserved in YKM 65729 + 66119 but partly obscured by matrix and other elements. The pubis has a squared outline, being as mediolaterally wide as it is anteroposteriorly long, as in other elasmosaurids (e.g. Welles, 1952). The acetabular portion is covered by matrix (Fig. 7C). The ventral surface is slightly convex, and the medial

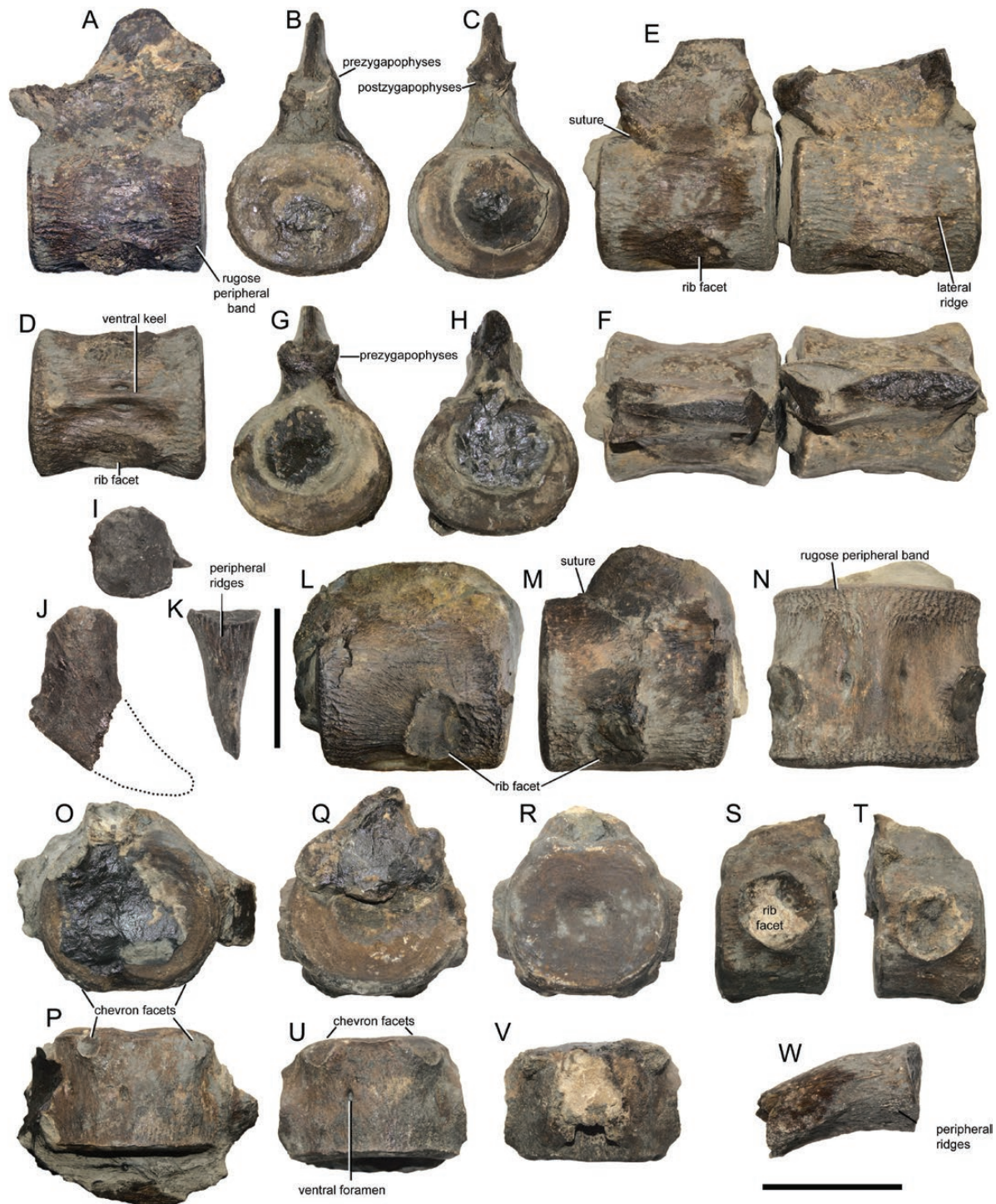


Figure 6. Axial skeleton of *cf. Jucha* UPM NV 15. A–D, anterior to middle cervical vertebra in left lateral (A), anterior (B), posterior (C) and ventral (D) views. E–H, two articulated anterior to middle cervical vertebrae in lateral (E), dorsal (F), anterior (G) and posterior (H) views. I–K, posterior cervical rib in proximal (I), dorsal (J) and posterior (K) views. L–N, posterior cervical centra in lateral (L, M) and ventral (N) views. O, P, caudal centrum in posterior (O) and ventral (P) views; Q–T, caudal centrum in anterior (Q), posterior (R), lateral (S, T), ventral (U) and dorsal (V) views. W, proximal portion of dorsal rib. Scale bars: 50 mm.

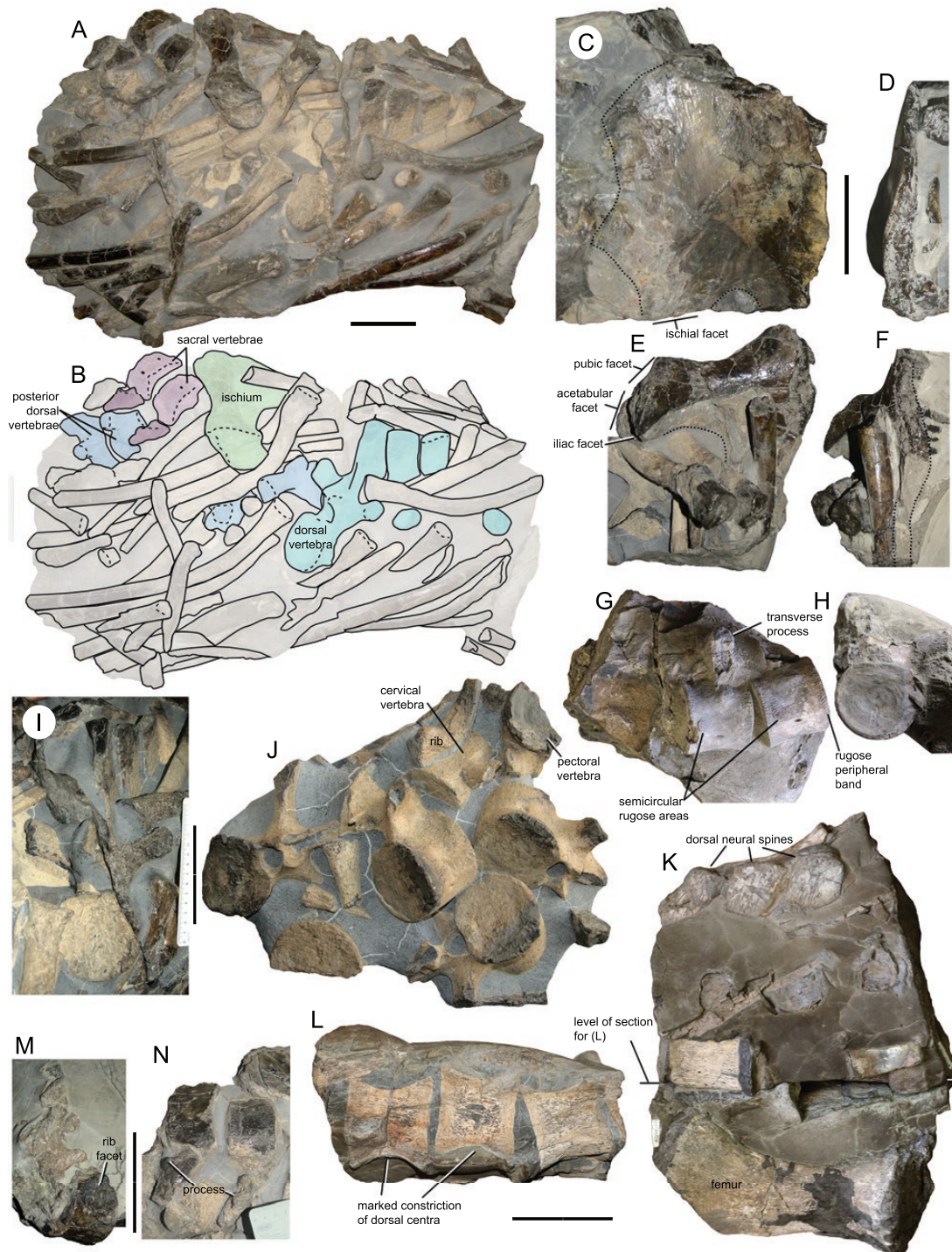


Figure 7. Axial and appendicular skeleton of cf. *Jucha* YKM 65729 + 66119. A, B, main block with vertebrae, ribs and pelvic girdle elements. C, D, left pubis in dorsal (C) and medial (D) views. E, F, left ischium in dorsal (E) and medial (F) views. G, H, pectoral vertebrae in left lateral (G) and posterior (H) views. I, mid-dorsal vertebra in posterior view. J, association of dorsal, cervical and pectoral vertebrae and ribs. K, concretion enclosing dorsal vertebrae and femur; L, cross-section of the same concretion. M, sacral vertebra in lateral view, N, sacral vertebrae and articulated ribs in ventral view. Scale bars: 100 mm.

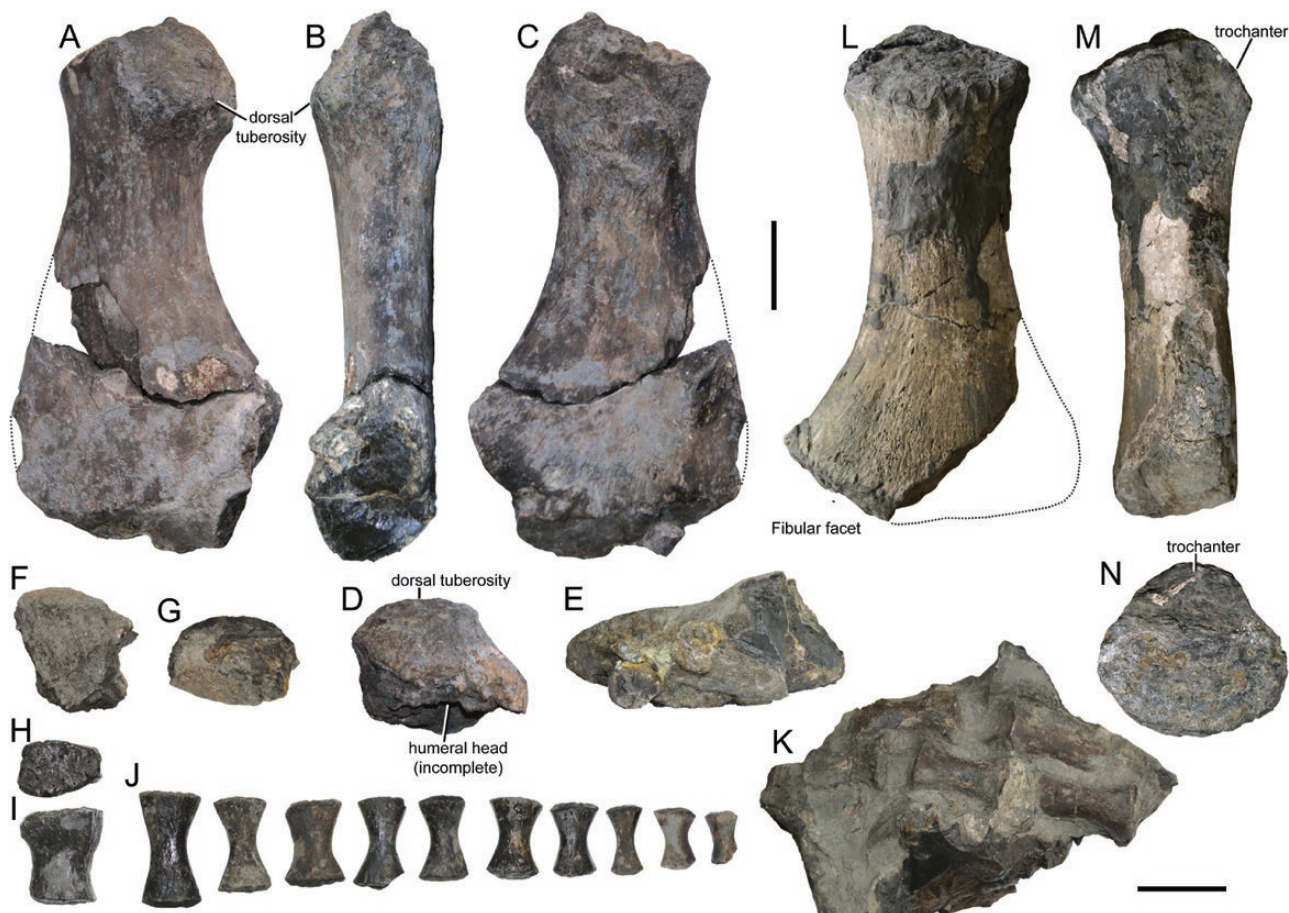


Figure 8. Appendicular skeleton of cf. *Jucha*. A–K, UPM NV 15, partial forelimb. A–E, left humerus in dorsal (A), posterior (B), ventral (C), proximal (D) and distal (E) views. F, G, partial epipodial element in dorsal (F) and proximal (G) views. H, I, possible first metacarpal in proximal (H) and dorsal (I) views. J, isolated phalanges. K, articulated phalanges in concretion. L–N, left femur of YKM 65729 + 66119 in ventral (L), anterior (M) and proximal (N) views. Scale bars: 50 mm.

symphysis is dorsoventrally compressed and sigmoidal (Fig. 7D).

Ischium: The left ischium is almost completely preserved in YKM 65729 + 66119, but most of the ischial blade is obscured by ribs. The ischium appears nearly as wide as long and is hatchet shaped (Fig. 7B, E). The anterior margin is shallowly concave, and the anteromedial process is short and ventrally curving, indicating the absence of a pelvic bar, unlike *El. platyurus*, *Li. morgani* and *Ka. lafquenianum* (Welles, 1943, 1952; O’Gorman, 2016; Sachs & Kear, 2017). The acetabular head is gracile, being weakly expanded anteroposteriorly (the ratio of the ischial neck-to-acetabular process length is ~0.8). The iliac and acetabular facets are poorly demarcated from the large, anteriorly facing facet for the articulation with the pubis; the acetabular contribution of the ischium appears small compared with the pubic facet (Fig. 7E). The ischium forms an anterior buttress connecting the acetabular head and the dorsomedial process. This

buttress gives the ischial symphysis a sigmoid shape (Fig. 7F).

Femur: A partial left femur is preserved in YKM 65729 + 66119. The femoral head of YKM 65729 + 66119 is massive and subcircular in outline, whereas it is dorsomedially flattened in the holotype of *J. squalea*. The dorsal trochanter is weakly expressed; its proximal surface makes an angle of ~110° with the proximal surface of the femoral head, as in UPM 2756/1-53. The trochanter is clearly narrower anteroposteriorly than the femoral head and is slightly inclined anteriorly (Fig. 8N). Although the femoral size is close to that of the holotype of *J. squalea*, it differs in the presence of a long shaft; the maximal constriction is set at midlength, whereas the maximal constriction is much more proximal in the holotype of *J. squalea* (cf. Figs 5A’, 8L). As a result, the distal expansion is restricted to the distal third of the femur, unlike in the holotype of *J. squalea* and the holotype of

Callawayasaurus colombiensis, where it starts more proximally [this is less clear in the referred specimen of *Callawayasaurus colombiensis* (Welles, 1962); this feature might thus be variable intraspecifically]. The femur is straight in anteroposterior view, lacking any evidence for a dorsal deflection.

PHYLOGENETIC PLACEMENT

Our implied weighting maximum parsimony analyses recovered 100.000+ most parsimonious trees, with a length of 146.62811 steps. The equal weight analysis recovered 100.000+ most parsimonious trees, with a length of 1842 steps. The general structure of the strict consensus trees (Fig. 9; Supporting Information, Figs S1, S2) and the composition of the major plesiosaurian clades do not differ from those obtained in previous iterations of the dataset (Benson & Druckenmiller,

2014; Fischer *et al.*, 2018; O’Gorman, 2020) and will not be discussed here.

We recover *J. squala* as one of the most basal elasmosaurids (Fig. 9), regardless of the optimality criterion used. However, basal polytomies are recovered in both the equal and implied weighting analyses, involving taxa frequently regarded as early elasmosaurids. For the implied weighting, this basal polytomy contains *Er. australis*, *Wa. betsynichollsae* and postcranial skeletons not yet formally described from the Hauterivian of the Speeton Clay Formation of England (NHMUK PV R8623 and SCARB 200751), known as the ‘Speeton Clay Plesiosaurian’ (Benson & Druckenmiller, 2014; Otero, 2016; Sachs *et al.*, 2017; Serratos *et al.*, 2017; O’Gorman, 2020). Unexpectedly, two elasmosaurine OTUs join this basal polytomy: *El. platyrus* and *N. bradti*. The basal polytomy is much larger in the equal weight analysis, mainly

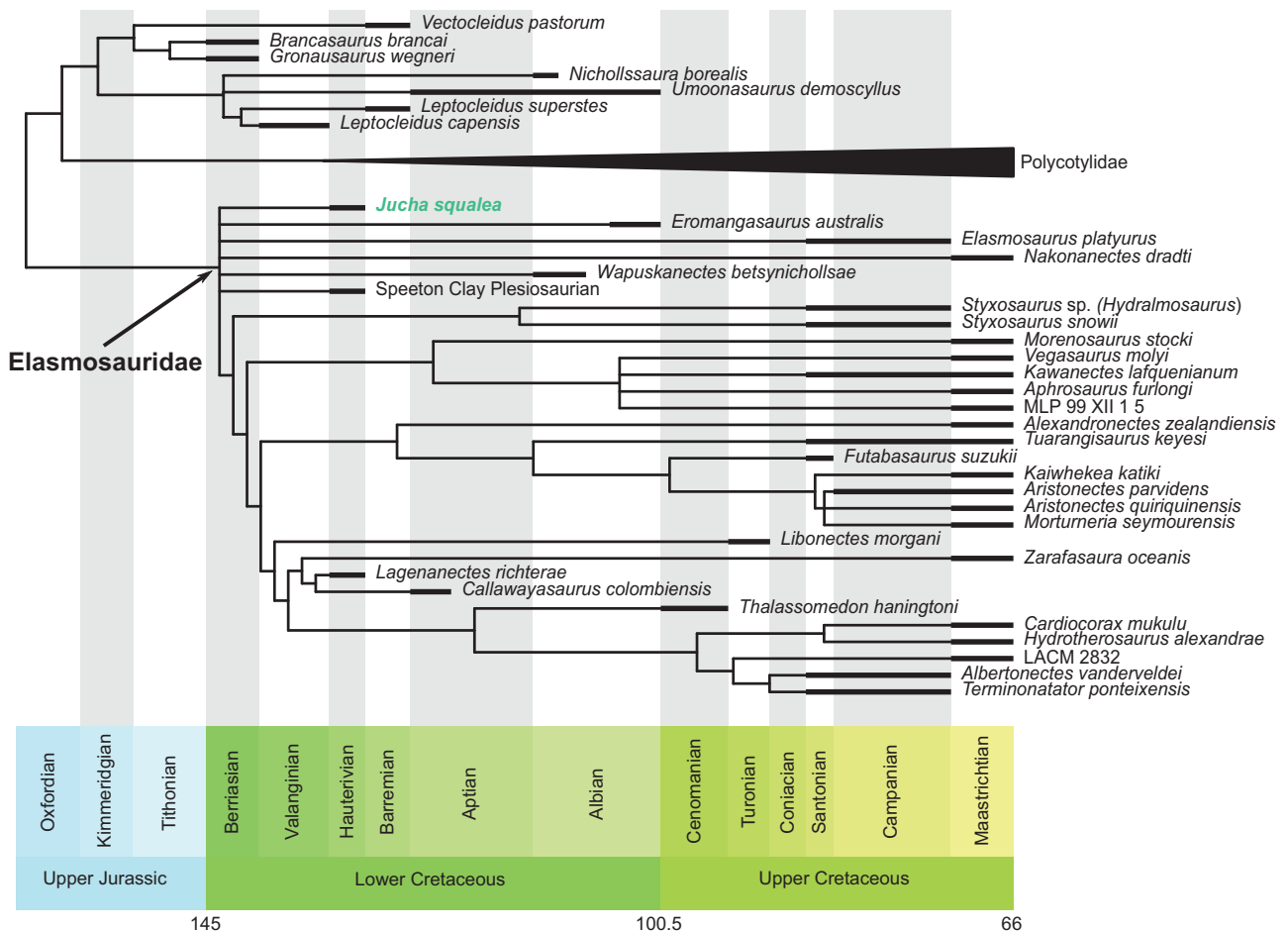


Figure 9. Phylogeny of xenopsarian plesiosaurians: time-scaled strict consensus of the implied weighting maximum parsimony analysis. The analysis was conducted on Plesiosauria as a whole, but only the relevant subset is presented here (see also Supporting Information, Figs S1, S2). Despite a polytomy at the base of Elasmosauridae, *Jucha squala* is recovered as a basal elasmosaurid.

because the Maastrichtian taxon *Alexandronectes parvidens* Cabrera, 1941 (whose scores were not altered from O’Gorman, 2020) is often recovered as a basal xenopsarian. In equal weights, *J. squala* forms a clade of early elasmosaurids with *Callawayasaurus colombiensis* and *Er. australis* (Supporting Information, Fig. S2). The clade Elasmosaurinae is recovered as in the study by O’Gorman (2020), in addition to non-aristonectine weddellonectians (Supporting Information, Fig. S2). The support for the trees arising from each method is low, as in the original analysis [although O’Gorman (2020) estimated support via a Bremer decay index, whereas we used symmetric resampling]. No method yields a better supported topology than the other, although the support for Elasmosauridae is slightly better in implied weighting than in equally weights (21 vs. 19; see Supporting Information, Figs S3, S4).

The addition of *J. squala* and *La. richterae* and the implied weighting framework result in most parsimonious trees that differ substantially from those of O’Gorman (2020). Generally, the topology we recovered in implied weighting appears less congruent with stratigraphy: *La. richterae* and *Callawayasaurus colombiensis* are recovered as fairly derived, closely related to the Late Cretaceous taxa *Albertonectes vanderveldei*, *Cardiacorax mukulu* Araújo et al. 2015, *H. alexandrae*, *Li. morgani*, *Th. haningtoni* and *Z. oceanis*. This has the effect of dragging several deeply nested nodes into the Early Cretaceous. In this topology, the clade Euelasmosaurida is restricted to a single OTU, *Li. morgani*. Weddellonectians are recovered as closely related, but the clade as presently defined is paraphyletic in our consensus tree. The clade with the most profound modifications is Elasmosaurinae, whose members are recovered as either basal (*El. platyurus* and *N. bradti*) or derived forms (*Albertonectes vanderveldei*, *H. alexandrae* and *Terminonatator ponteixensis* Sato, 2003).

PATTERNS OF CERVICAL ELONGATION

Phylogenetic uncertainties set aside, *J. squala* unambiguously represents an early attempt at cervical elongation (Fig. 10; Supporting Information, Fig. S5), recording high length-to-height ratios compared with other Early Cretaceous taxa. Indeed, the holotype of *J. squala* (1.36) is surpassed only by *Er. australis* (1.37), cf. *Jucha* (1.4), an indeterminate elasmosaurid from the Valanginian of Russia (1.41) and some cervical centra of the holotype of ‘*Woolungasaurus glendowerensis*’ (1.51; regarded as an indeterminate Aptian elasmosaurid by Kear, 2007), which differs from the much lower values of *Callawayasaurus*

colombiensis (1.19), *La. richterae* (1.06) and non-elasmosaurid xenopsarians (0.80–0.92). The lowest value among Early Cretaceous elasmosaurids is that of the anterior cervicals of ‘*Cimoliasaurus maccoyi*’ Etheridge, 1904 (another indeterminate Aptian elasmosaurid Kear, 2002a), with a value of 0.97 (more posterior cervicals record a higher value of 1.19). The full range of cervical elongation values recorded by Early Cretaceous elasmosaurids (0.97–1.51) is thus substantial (Fig. 10; Supporting Information, Fig. S5), but still a far cry from what elasmosaurids evolved during the Late Cretaceous, especially during the Campanian–Maastrichtian interval (0.9–1.8).

Several taxa exceed the value of *J. squala* during the Late Cretaceous: *Albertonectes vanderveldei*, *Cardiacorax mukulu*, *El. platyurus*, *H. alexandrae*, *Styxosaurus* spp. and *Te. ponteixensis*. Although there is evidence that elasmosaurid taxa as a whole record longer cervical centra during the Late Cretaceous (Fig. 10; Supporting Information, Fig. S5), the uncertainties of the phylogenetic relationships of elasmosaurids (see above) and the indeterminate status of ‘cimoliasaurids’ make it hard to identify unambiguous episodes of cervical reduction, besides *N. bradti* and aristonectines (Serratos et al., 2017).

DISCUSSION

Neck length in sauropterygians is more strongly driven by changes in the number of vertebra than by a modification of their individual shape (Soul & Benson, 2017). As a result, the incomplete preservation of the neck in the type specimen precludes a precise evaluation of the length of the neck of *J. squala*. However, centrum shape also remains an important parameter, and the data of Soul & Benson (2017) indicate that the number of cervical centra in long-necked plesiosaurians is well correlated (Pearson’s $r = 0.8, P < 0.001$; see Supporting Information, Table S2) with their average length. Cervical centrum elongation is thus a useful parameter to investigate neck elongation when complete skeletons are missing, as is the case for the early evolution of elasmosaurids. The anterior cervical centra of *J. squala* are clearly more elongated than those of other early xenopsarians, yielding one of the highest values of the Early Cretaceous; it now appears clear that elasmosaurids attained a substantial range of cervical elongation values early in their history (Fig. 10; Supporting Information, Fig. S5), thanks to the co-occurrence of *J. squala*, *La. richterae* and indeterminate elasmosaurids from the Valanginian–Hauterivian interval of Russia and Argentina (O’Gorman et al., 2015a; Sachs et al., 2017; Zverkov

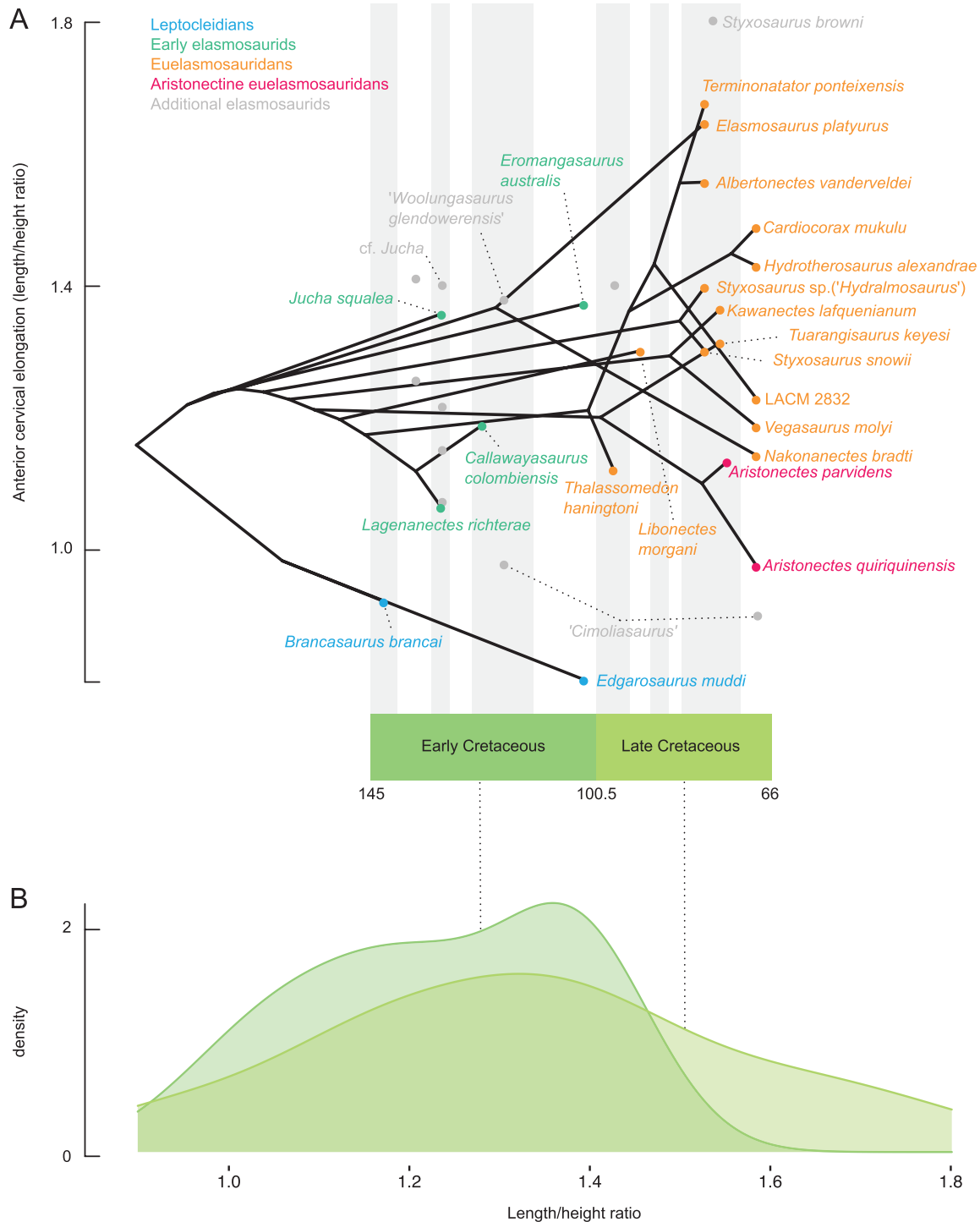


Figure 10. Patterns of vertebral elongation in anterior cervical centra in elasmosaurids. A, phenogram using the anterior cervical centra (at or close to the tenth cervical centrum); additional elasmosaurids not included in the phylogeny have also been mapped (grey dots). B, density distribution of cervical elongation values for the Early and Late Cretaceous.

& Kiselev, 2018). However, this range is dwarfed by that of Campanian–Maastrichtian elasmosaurids (Fig. 10; Supporting Information, Fig. S5), and our

results corroborate the hypothesis of a complex rather than trended evolution of relative neck lengths in elasmosaurids (see Serratos *et al.*, 2017).

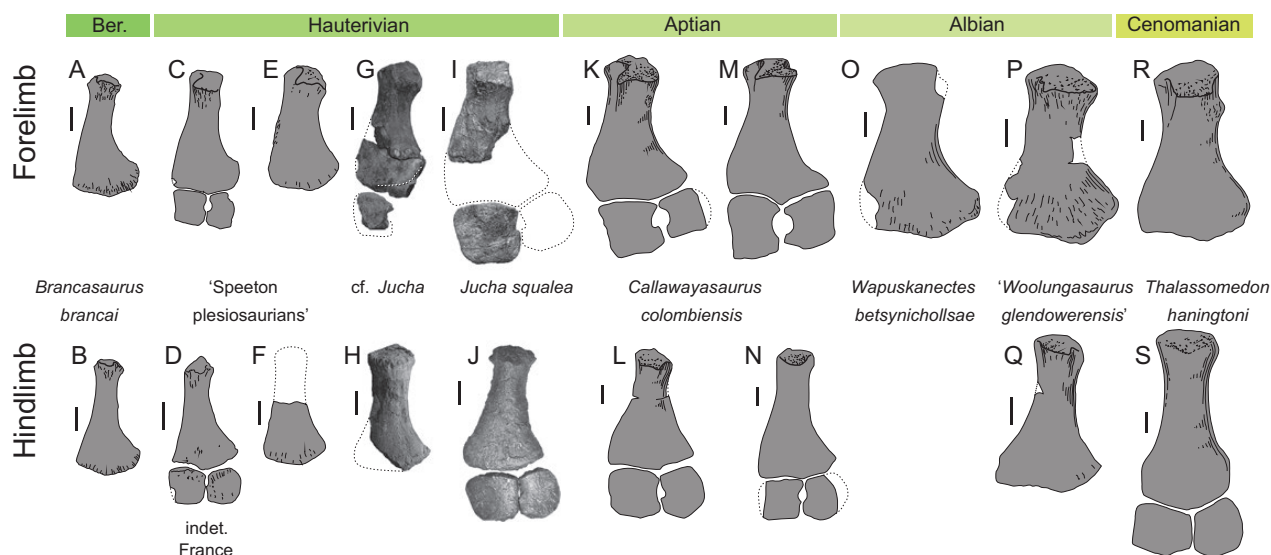


Figure 11. Comparative anatomy of Berriasian–Cenomanian long-necked leptocleidian (A, B) and elasmosaurid (C–S) limbs (propodial + epipodium). A, B, *Brancasaurus brancai* [holotype, GPMM A3.B4; modified from Sachs *et al.* (2016), mirrored for comparative purposes]. C, E, F, the so-called Speeton Clay plesiosaurs (C, SCARB 200751; E, F, NHMUK PV R8623; based on personal observations by N. G. Zverkov). D, indeterminate elasmosaurid from France (modified from Fournier *et al.*, 1982). G, H, cf. *Jucha* (G, UPM NV 15; H, YKM 65729 + 66119, mirrored for comparative purposes). I, J, *Jucha squalea* (UPM 2756/1–53). K–N, *Callawayasaurus colombiensis* (K, L, holotype UCMP 38349; M, N, referred specimen SGC MGJRG.2018.V.1; both modified from Welles, 1962). O, *Wapuskaneetes betsynichollsae* (TMP 98.49.02, in ventral view, modified from Druckenmiller & Russell, 2006). P, Q, indeterminate elasmosaurid [holotype of ‘*Woolungasaurus glendowerensis*’ QM D 6890; QMF3567 in Sachs (2004), modified from Persson (1960) and mirrored for comparative purposes]. R, S, *Thalassomedon haningtoni* (holotype, CMNH 1588, modified from Welles, 1962). All specimens (except for O) are shown in dorsal view. Scale bars: 50 mm.

Jucha squalea departs from many other elasmosaurids, including Early Cretaceous ones, by possessing bulky propodials and by lacking a median pectoral bar and a heart-shaped intercoracoid fenestra. Despite a generally poor fossil record, Early Cretaceous elasmosaurids appear dissimilar, displaying a range of coracoid and propodial shapes, with varying degrees of elongation and postaxial deflection (Fig. 11). This, in turn, suggests that these taxa did not differ only in relative neck lengths but also in the shapes of their flippers. Although it is tempting to use this as evidence for high early disparity in elasmosaurids [which appears common in xenopsarians, being documented in leptocleidians (Benson *et al.*, 2013) and polycotylids (Fischer *et al.*, 2018)], only an increased sampling among early elasmosaurids could determine how peculiar the morphologies of *Jucha*, *Callawayasaurus*, *Lagenanectes* and *Wapuskaneetes* are.

An unstable phylogenetic signal is another factor currently preventing a thorough understanding of the elasmosaurid diversification. Our analyses of the dataset from O’Gorman (2020) in implied and equal weighting frameworks yielded topologies that are, in places, clearly at odds with those obtained by the

original author, and which were used to define or redefine suprageneric clades. This suggests that at least some of the features that support Elasmosaurinae and Euelasmosaurida might be homoplastic. However, this mismatch is not a new problem; the instability of elasmosaurid relationships has been discussed by Serratos *et al.* (2017), who found low congruence between the results of the analyses of the last semi-decade. At any rate, the wealth of new elasmosaurid data published in the recent years coupled with the ever-increasing knowledge on the behaviour of phylogenetic methods and parameters (Bapst *et al.*, 2016; O’Reilly *et al.*, 2016; Rosa *et al.*, 2019; Smith, 2019) might solve this long-standing conundrum, along with increased and optimized taxonomic and character sampling.

CONCLUSIONS

We describe a new basal elasmosaurid from the upper Hauterivian of European Russia, *J. squalea*. This taxon represents one of the geologically oldest occurrences of elasmosaurids. It lacks a series of features that otherwise characterize the group, such as the pectoral

bar and the heart-shaped intercoracoid fenestra, and thus documents some of the earliest stages of the elasmosaurid radiation. *Jucha squalea* marks an early attempt at cervical elongation in elasmosaurids via differential growth, possessing anterior cervical centra that are much more elongated than those of other early xenosaurs. The cervical shape values we gathered suggest that elasmosaurids underwent multiple episodes of cervical shortening, notably during the Early Cretaceous. However, the precise patterns of cervical elongation and character acquisition in elasmosaurids are obscured by an unstable phylogenetic signal. Indeed, our implied weighting maximum parsimony analysis does not recover the clades Euelasmosaurida, Aristonectinae and Elasmosaurinae as currently defined, suggesting that homoplasy plays a pervasive role in elasmosaurid phylogenetics.

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SUPPORTING INFORMATION

Additional Supporting Information may be found in the online version of this article at the publisher's web-site:

Table S1. Temporal data used to provide a time scale for the phylogenetic tree. See also the Supporting Information (supplementary file 10 ‘SUPP_ranges.txt’).

Table S2. Body plan from [Soul & Benson \(2017\)](#), focusing on long-necked plesiosaurians (i.e. with Triassic sauropterygians, thalassophoneans, rhomaleosaurids and polycotylids removed). The contribution column (‘contrib’) indicates the average length of cervical centra, obtained by dividing the length of the neck by the number of cervical centra. See [Soul & Benson \(2017\)](#) and references therein for the data source. See also the Supporting Information (supplementary file ‘SUPP_S&B2017_long_necked.csv’).

Figure S1. Strict consensus of the most parsimonious trees arising from our implied weight cladistic analysis.

Figure S2. Strict consensus of the most parsimonious trees arising from our equal weight cladistic analysis.

Figure S3. Clade support by symmetric resampling, implied weighting analysis.

Figure S4. Clade support by symmetric resampling, equal weighting analysis.

Figure S5. Patterns of vertebral elongation in cervical centra in elasmosaurids. A, phenogram using the relatively longest centra of the neck, regardless of its position; additional elasmosaurids not included in the phylogeny have also been mapped (grey dots). B, density distribution of cervical elongation values for the Early and Late Cretaceous.