

New Late Cretaceous zhelestid mammal from the Bayanshiree Formation, Mongolia

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Numerous exquisitely preserved mammal fossils unearthed from Upper Cretaceous strata in the Gobi Desert of Mongolia have played a key role in understanding Mesozoic mammalian evolution. These splendid mammal fossils have been recovered mainly from the Baruungoyot and Djadokhta formations, but only two fragmentary remains of mammals have been collected from the underlying Bayanshiree Formation. Here we report a new species of Zhelestidae, *Ravjaa ishihi* gen. et sp. nov., based on a new mammal specimen recently discovered from the Bayanshiree Formation at the Bayan Shiree locality. The new specimen is represented by a well-preserved partial right dentary with the distal portion of an ultimate premolar and the first to the third molars. The gross observation and the phylogenetic analyses demonstrate zhelestid affinities: the subequally tall protoconid and metaconid, closely approximated hypoconulid and entoconid. Zhelestidae was widely distributed from Eurasia to North America in the Late Cretaceous but has not previously been recovered in the abundant Late Cretaceous mammalian fauna of Mongolia. This species represents the first zhelestid from the Mongolian Upper Cretaceous strata and contributes to further understanding of their paleobiogeographic and ecological insights.

Key words: Mammalia, Eutheria, Cretaceous, Bayanshiree Formation, Mongolia.

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Introduction

The Upper Cretaceous strata of Mongolia provided over 20 mammalian species, mostly known from the aeolian-domi-

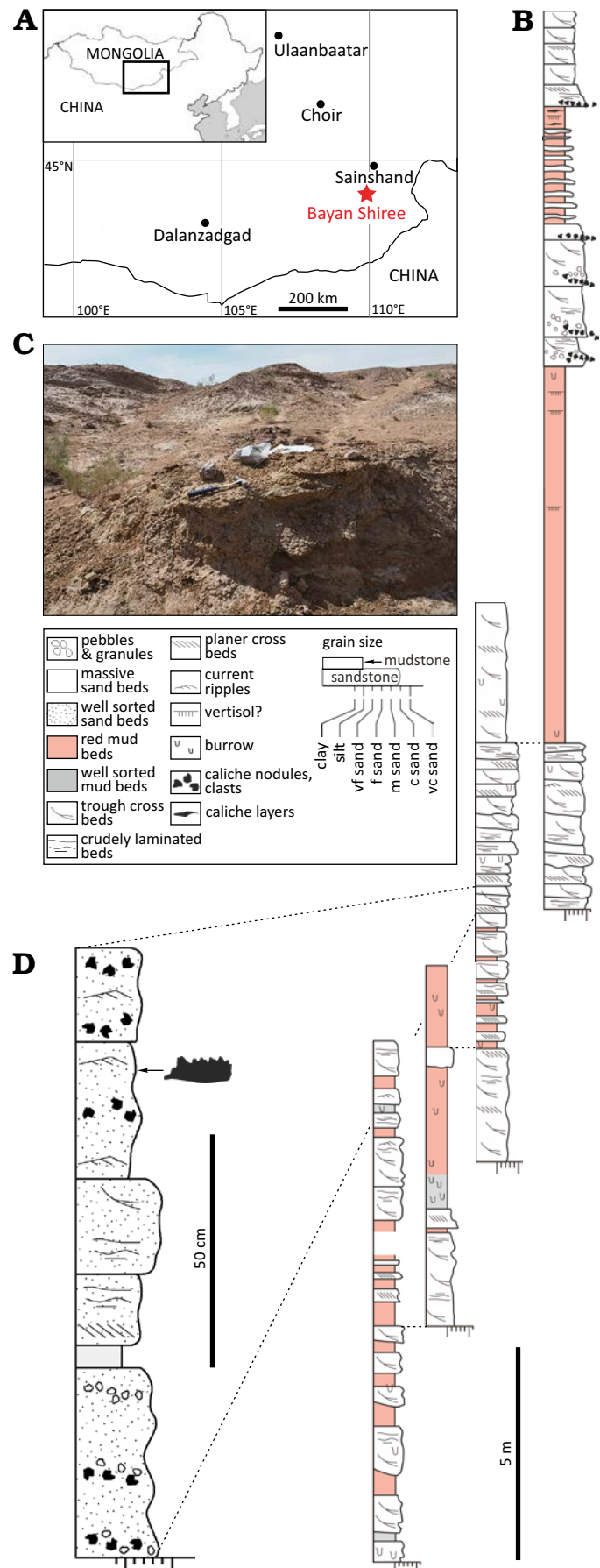
nated Djadokhta and Baruungoyot formations (16 species from the former and seven species from the latter, Lopatin 2020). The mammalian fossils from these aeolian-dominated formations include nearly complete skulls and skel-

etons (e.g., Kielan-Jaworowska 1970; Novacek et al. 1997; Wible et al. 2007; Hurum and Kielan-Jaworowska 2008). The mammalian faunas in these strata are dominated by multituberculates (Averianov and Archibald 2005; Kielan-Jaworowska et al. 2004) with some metatherians (e.g., Gregory and Simpson 1926; Trofimov and Szalay 1994) and eutherian Asioryctitheria (e.g., Kielan-Jaworowska 1969, 1975; Novacek et al. 1997), Zalambdalestidae (e.g., Gregory and Simpson 1926; Kielan-Jaworowska 1975; Fostowicz-Freluk 2016), and Cimolestidae (e.g., Wible et al. 2007). Mammalian remains from the overlying Nemegt Formation, which is fluvial-dominated and partially contemporaneous with the Baruungoyot Formation, are scarce but represented by two taxa, a fragmentary cranial material of a multituberculate and a relatively well-preserved cranium of a metatherian (e.g., Kielan-Jaworowska and Sochava 1969; Szalay and Trofimov 1996). However, the understanding of mammalian fauna in the Bayanshree and the Javkhlant formations, the lowermost Upper Cretaceous strata in Mongolia, are much more limited; conference abstracts of unnamed eutherians from the Javkhlant Formation (Giallombardo and Novacek 2006; Giallombardo 2007) and the recent description of a metatherian *Tsagandelta dashzevegi* Rougier et al., 2015, and an asioryctitherian *Bayshinoryctes shuvalovi* Lopatin & Averianov, 2023, from the Bayanshree Formation. Of particular interest in Mongolia is the absence of zhelestids, which are one of the Late Cretaceous eutherian clades (Wible et al. 2007, 2009; Archibald and Averianov 2012), well-known from contemporaneous Cenomanian to Maastrichtian strata across multiple continents: Asia, Europe, North America, and potentially Africa (Wible et al. 2009; Archibald and Averianov 2012). Among these continents, Asia represents the highest taxonomic diversity and the numeric abundance of zhelestids, mostly from central Asian countries such as Uzbekistan and Kazakhstan. Here we describe the first zhelestid specimen from Mongolia, which is also the third mammalian material from the Bayanshree Formation. This specimen provides new insights into the paleobiogeographical hypothesis of this clade that previous studies provoked (Nessov et al. 1998).

Institutional abbreviations.—MPC, Mongolian Paleontological Center, Mongolian Academy of Sciences, Ulaanbaatar, Mongolia (formerly Institute of Paleontology, Mongolian Academy of Science).

Nomenclatural acts.—This published work and the nomenclatural acts it contains have been registered in ZooBank: urn:lsid:zoobank.org:pub:CC5A9DA0-2082-410D-9DA0-B54744E8197F.

Fig 1. Map of the locality of *Ravjaa ishiii* gen. et sp. nov. in Mongolia. **A.** Southern Mongolia with the position of the Bayan Shiree locality. **B.** Combined stratigraphic columns at Bayan Shiree locality. Photograph (C) and stratigraphic column (D) of the layer from which the specimen was derived. The locality is indicated by a specimen silhouette.



Material and methods

The new eutherian specimen described here, MPC-M 100s/001, was recovered from the Bayanshree Formation exposed at the Bayan Shree locality of the Gobi Desert, Mongolia (Fig. 1A). This formation is widely distributed throughout the eastern and southern Gobi (Martinson 1982) and approximately 35 m of this formation is exposed at Bayan Shree (Fig. 1B). The depositional age of the Bayanshree Formation had been recognized as Cenomanian to Santonian based on biostratigraphical data and K-Ar dating on basalt (e.g., Jerzykiewicz and Russel 1991; Shuvalov 2000; Averianov and Sues 2012). This age is supported by the recent U-Pb dating recovered 95.9 ± 6.0 and 89.6 ± 4.0 Ma (Cenomanian–Santonian) on caliches recovered from the Khongil Tsav locality, which is 19 km east of the Bayan Shree locality (Kurumada et al. 2020). The section at Bayan Shree mainly consists of mudstone and sandstone. Laterally continuous red mudstone layers, each about 5 to 10 m thick, are observed throughout the section. Most of these mudstone layers are directly overlain by sandstones exhibiting fining-upward sequences with well-developed sedimentary structures such as trough cross-stratification, planar cross-stratification, and current ripple lamination. The sandstone layers of the middle to upper part tend to be thicker, laterally continuous and have well-developed sedimentary structures, but those of the lower part are thinner and less laterally continuous with poorly developed sedimentary structures. These lithostratigraphic features suggest that the section was composed of fluvial deposits. The lower part of the sections is likely to be braided river deposits, whereas the middle and upper parts are deposited with the influence of meandering rivers.

In 2019, a member of the joint expedition of the Institute of Paleontology and Geology (currently the Institute of Paleontology) of the Mongolian Academy of Sciences and Okayama University of Science, Japan found a layer with abundant microvertebrate remains (Ishigaki et al. 2019). The fossil remains from the layer include isolated bones of fishes, anurans, lizards, turtles, crocodylians, non-avian dinosaurs, and the mammal described herein. The mammal specimen was surface-collected from near the top of the mound, indicating the specimen is not a float from the higher stratigraphic position (Fig. 1C). The specimen-bearing layer is in an approximately 1-meter thick point bar sandstone of a laterally accretional meandering channel deposit. Although the sedimentological feature of the host facies indicates that the specimen was transported prior to the final burial event, the pristine preservation of the specimen without abrasion suggests that the distance of the transportation was not significant.

We describe MPC-M 100s/001 following Nessov et al. (1998) for dental terminologies unless noted. Molar measurements were taken based on the scheme of Archibald (1982). The comparable morphological data of other zhelestid specimens were compiled from previous literature. We con-

ducted micro X-ray computed tomography (CT) scanning on MPC-M 100s/001 in addition to the gross observation. The scan images were obtained by Nikon XT H 225 ST at the Industrial X-ray/CT Systems Rental Lab, Nikon Solutions CO., LTD., in Yokohama, Kanagawa Prefecture, Japan. The obtained CT images were rendered using VGSTUDIO MAX 3.4 (Volume Graphics, Germany) and segmented using 3DSlicer version 5.3 (Fedorov et al. 2012).

The phylogenetic analyses were performed using TNT 1.5, May 2022 edition “No Taxon Limit” (Goloboff and Catalano 2016), based on the data matrix of Averianov and Archibald (2016), which is composed of a broad taxonomic sampling of Cretaceous eutherian mammals, including zhelestids. We added *Valentinella vitrollense* Tabuce et al., 2004, *Mistralestes arcensis* Tabuce et al., 2013, and *Azilestes ragei* Gheerbrant and Teodori 2021, to this dataset (see SOM 1, Supplementary Online Material available at http://app.pan.pl/SOM/app70-Okoshi_etal_SOM.pdf). The data matrix consists of 95 taxa and 416 characters (see Appendix 1 for scorings of the newly described specimen in this study), including *Nanolestes drescheriae* Martin, 2002, as the outgroup. All of the characters are equally weighed and not ordered. The maximum number of trees was set to 99 999, and then a traditional search with 100 random seeds and 1000 replicates of Wagner trees, followed by the TBR branch swapping that held 100 trees per replicate, was performed. The branch supports were evaluated by bootstrap resampling using standard absolute frequencies (10 000 replicates). The resulting phylogenetic relationships were visualized using R version 4.3.0, employing two packages: APE version 5.8 (Paradis et al. 2004) and strap version 1.6-1 (Bell and Lloyd 2015). The locality map was produced using GMT6 (Wessel et al. 2019).

Systematic palaeontology

Mammalia Linnaeus, 1758

Theria Parker and Haswell, 1897

Eutheria Gill, 1872

Zhelestidae Nessov, 1985a

Genus *Ravjaa* nov.

Zoobank LSID: urn:lsid:zoobank.org:act:500CED48-4C63-406B-8D17-50FCD17513D3.

Etymology: Derived from the 5th Noyon Khutugt lama of the Gobi, Dulduityn Danzanravjaa, the famous Buddhist scholar who was deeply respected by people in the Dornogobi region, where this specimen was unearthed. Given that the genus name “*Ravjaa*” originates from the personal name of a male figure, it should be considered masculine.

Type species: *Ravjaa ishiii* sp. nov.; see below.

Diagnosis.—As for the type species by monotypy.

Stratigraphic and geographic range.—Bayanshree Formation, Cenomanian–Santonian, Bayan Shree, Gobi Desert, Mongolia.

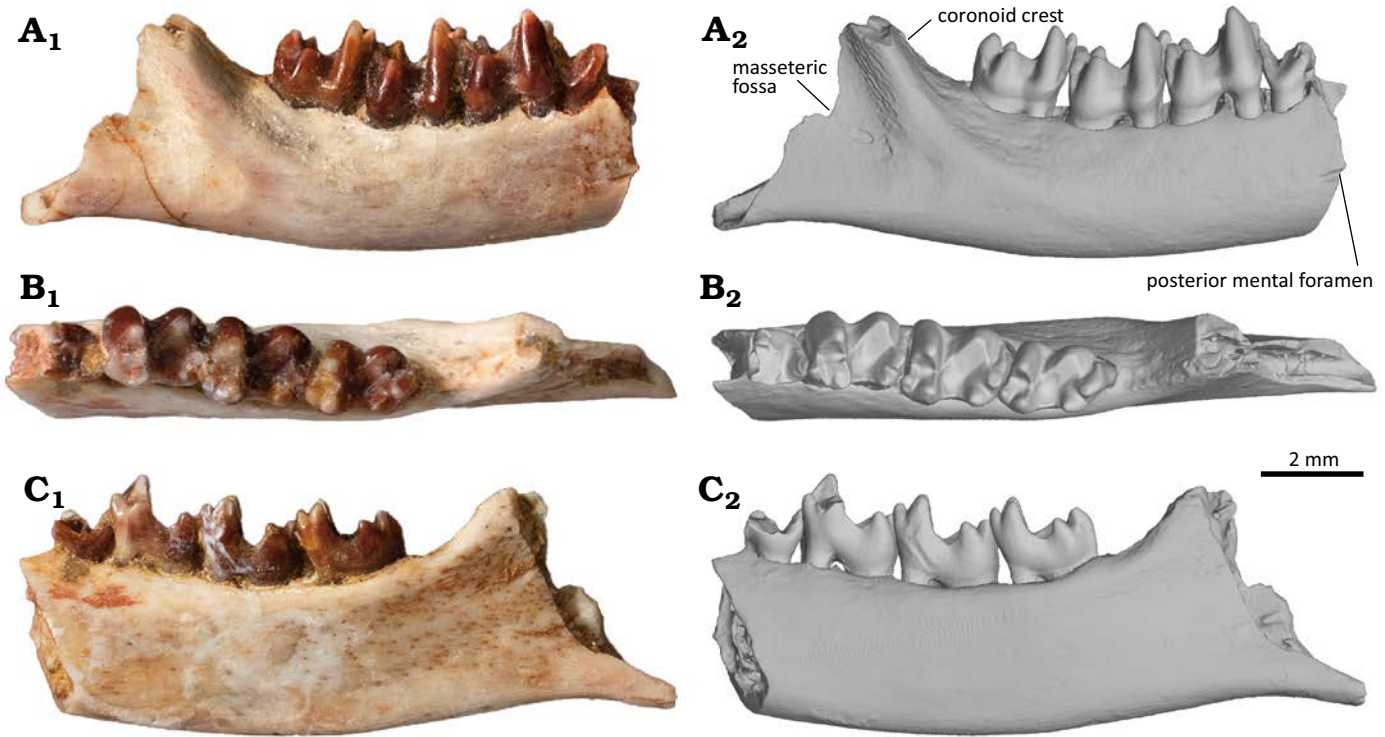


Fig. 2. Holotype of the zhelestid eutherian mammal *Ravjaa ishiii* gen. et sp. nov. (MPC-M 100s/001) from the Upper Cretaceous Bayanshiree Formation, Dornogovi, Mongolia. Right mandible in labial (A), occlusal (B) and lingual (C) views. Photographs (A₁–C₁) and 3D model (A₂–C₂) produced from a CT scan.

Ravjaa ishiii sp. nov.

Figs. 2–4.

Zoobank LSID: urn:lsid:zoobank.org:act:416ECE52-5728-493F-8DAB-2632FF8228D4.

Etymology: In honor of Ken-ichi Ishii, the director of Hayashibara Museum of Natural Sciences, Okayama, Japan, for his leading role in the Japanese team of the Mongolia-Japan joint expedition, which tremendously contributes to Mongolian paleontology.

Holotype: MPC-M 100s/001, partial right dentary with a distal portion of the ultimate premolar, damaged m1, and well-preserved m2–3.

Type locality: Bayan Shiree, located approximately 25 km southwest of Dzüünbayan, Dornogovi, Mongolia.

Type horizon: Cenomanian–Santonian, Upper Cretaceous, Bayanshiree Formation.

Diagnosis.—A zhelestid eutherian that differs from all other zhelestids in: the faint coronoid crest of the dentary with the shallow masseteric fossa, the concave dorsal alveolar margin of the dentary, the complete absence of the labial mandibular foramen, and the significantly high m2 crown that reaches approximately 70% of the height of the horizontal ramus directly below it on labial view.

Ravjaa ishiii sp. nov. differs from *Borisodon kara* Archibald & Averianov, 2012, in the rounded mesiolingual vertical crest of the paraconid, the coronoid process tilting 110–115°, and the lack of Meckelian groove; differs from *Gallolestes* spp. (*Gallolestes pachymandibularis* Lillegraven, 1976, and *Gallolestes agujaensis* Cifelli, 1994) in the less bulbous, more anteroposteriorly compressed protoconid; differs from *Azilestes ragei* in the lack of hypolophid; differs from

Parazhelestes spp. (*Parazhelestes robustus* Nesson, 1993, and *Parazhelestes mynbulakensis* Nesson, 1985b) in the absence of a marked tuberosity or concavity in the medial side of the area of the coronoid facet between the m3 and base of the coronoid process; differs from *Borisodon kara* and *Eoungulatum kudukensis* Nesson et al., 1998, by the absence of coronoid facet; differs from *Eozhelestes* in the complete labial attachment of the cristid oblique to the notch of the protocristid; differs from *Zhalmouzia bazhanovi* Averianov et al., 2014, in the lack of cingulid of the lingual and labial side of molars; differs from *Borisodon kara* and *Eozhelestes mangit* Nesson, 1997, by the trigonid height is less than twice the talonid height; differs from *Avitotherium utahensis* Cifelli, 1990, and *Lainodon orueetxebarriai* Gheerbrant & Astibia, 1994, by the slight labial shift of the paraconid; differs from *Sheikhdzheilia rezvyii* Averianov & Archibald, 2005, and *Lainodon orueetxebarriai* by presence of the labial postcingulid; differs from *Mistralestes arcensis*, the entocoid subequal to the hypoconulid; differs from *Mistralestes arcensis* in the precingulid with a distinct posteroventral cingular shelf; differs from *Eozhelestes mangit*, *Sheikhdzheilia rezvyii*, *Lainodon orueetxebarriai*, *Mistralestes arcensis* in the transversed protocristid; differs from *Zhalmouzia bazhanovi*, *Mistralestes arcensis*, and *Valentinella vitrollense* in the position of the posterior mental foramen below the ultimate premolar; differs from *Eozhelestes mangit*, *Parazhelestes* spp., *Aspanlestes aptap* Nesson, 1985a, *Zhelestes termirkazyk* Nesson, 1985a, *Eoungulatum kudukensis*, and *Zhalmouzia bazhanovi* by the ultimate molar hypoconulid being tall and

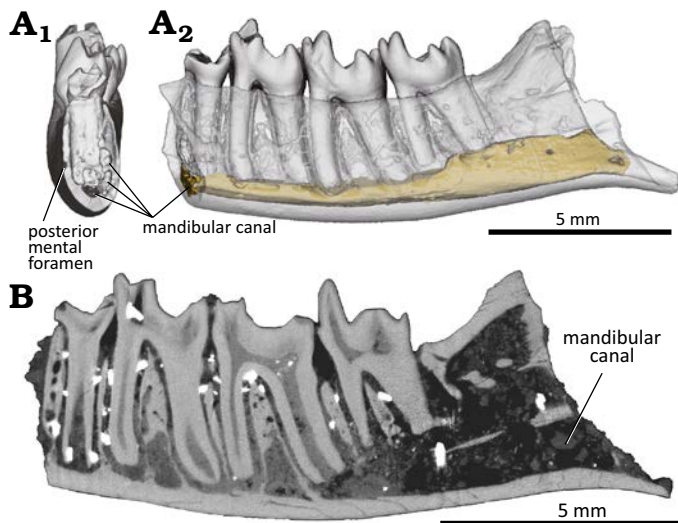


Fig 3. Digital reconstruction and CT image of the zhelestid eutherian mammal *Ravjaa ishiui* gen. et sp. nov. from the Upper Cretaceous Bayanshiree Formation, Dornogovi, Mongolia. **A.** 3D model in mesial (A_1) and lingual (A_2) views. **B.** Longitudinal section of mandible.

recurved; differs from *Eozhelestes mangit*, *Borisodon kara*, *Parazhelestes* spp., *Aspanlestes aptap*, *Zhelestes termirkazyk*, *Eoungulatum kudukensis*, and *Zhalmouzia bazhanovi* by the absence of the labial mandibular foramen; differs from *Eozhelestes mangit*, *Shikhdzheilia rezvyii*, *Borisodon kara*, *Lainodon orueetxebarriai*, *Aspanlestes aptap*, *Zhelestes termirkazyk*, *Eoungulatum kudukensis*, and *Zhalmouzia bazhanovi* by the compressed trigonid angle being less than 35° .

Description.—The new specimen, MPC-M 100s/001, is a partial right mandible retaining roughly the posterior half of the horizontal ramus, the base of the coronoid process, the anteroventral part of the ascending ramus, the talonid of the ultimate premolar, and m1–3 (Fig. 2). MPC-M 100s/001 represents an adult individual based on the presence of the m3 and the absence of replacement tooth, observed in micro CT images (Fig. 3). Despite the adult status, this specimen represents one of the smallest zhelestids reported to date, together with *Shikhdzheilia rezvyii* (Averianov and Archibald 2005) and *Aspanlestes aptap* (Archibald and Averianov 2012). The dorsoventral depth of the horizontal ramus is consistent throughout its preserved region. The alveolar margin of the ramus is slightly concave in the labial and lingual view, unlike the other zhelestids in which the margin is straight throughout its length (Fig. 2A, C). The dorsal margin of the ramus is higher lingually than labially (Fig. 2A₂). The ventral margin of the horizontal ramus is convex in labial and lingual views (Fig. 2A, C). Ventral to the ultimate premolar, the labial surface of the mandible bears a shallow sulcus that runs anteroposteriorly (Fig. 2A). The sulcus likely connects to a mental foramen anteriorly, as in *Borisodon kara*, *Parazhelestes* spp., and *Eoungulatum kudukensis* (Nessov et al. 1998; Archibald and Averianov 2012). The coronoid crest emerges as a low eminence, ventrally to the m2 talonid, at the level of the middle of the horizontal ramus (Fig. 2A₂).

Musculus buccinator is likely to insert anteromedial to the eminence. The crest extends posterodorsally and becomes prominent but is much less developed than the other zhelestids (e.g., *Borisodon kara*, *Aspanlestes aptap*, *Eoungulatum kudukensis*; Archibald and Averianov 2012), which may be suggestive of a reduced M. temporalis on this individual. The posterior surface of the coronoid crest is rugose and lacks the labial mandibular foramen (Fig. 2A₂). The coronoid crest defines the dorsal border of the masseteric fossa. The masseteric fossa is shallow, and its dorsal region likely provides the insertion for M. zygomaticomandibularis, whereas its ventral region is likely to be the insertion for M. masseter (Turnbull 1970). Ventral to the masseteric fossa, the ventral margin of the mandible, anterior to the angular process, is gently concave (Fig. 2A). Lingually, the ascending process lacks the coronoid facet, which can be observed in *Eoungulatum kudukensis* and *Parazhelestes* spp. (Archibald and Averianov 2012). The Meckelian sulcus is absent, as in all members of the Zhelestinae (Archibald and Averianov 2012). The CT images demonstrate the presence of the mandibular canal that runs anteroposteriorly throughout the mandible and the absence of replacement teeth (Fig. 3). Since no replacement teeth are present, all teeth observed in this specimen are considered to be permanent teeth.

MPC-M 100s/001 retains a partial ultimate premolar and all three molars (Fig. 3). While the anterior half of the ultimate premolar is severely damaged, the talonid and the posterior root are well preserved. Despite the damage, the anterior portion of the premolar width is clearly wider than the talonid. The talonid has a narrow and shallow basin anteriorly and has a single cusp posteriorly. All three molars are nearly complete, including two roots per molar, except for the apex of the m1 metaconid. The lower molars m1–3 are tribosphenic molars, and their shapes generally resemble each other. The overall sizes of the molars are subequal to each other, although the m3 is slightly smaller than the other two. The height of the m2 crown is 71% of the height of the horizontal ramus directly below it on labial view, which is markedly larger than in the other zhelestids (50–63%, with 56% on average).

The molars have moderately compressed trigonid with trigonid angles of 33° in m2 and m3 (Fig. 4C₁ and D₁). The maximum height of the trigonid is higher than that of the associated talonid but does not reach twice the talonid height (Fig. 4B₃–D₃). Their cusps are rounded as in *Parazhelestes* spp. and *Eoungulatum kudukensis* but unlike *Borisodon kara*. Among the trigonid cusps, paraconid is the smallest and is placed mesiolabially (Fig. 4B₁–D₁). This is also the only trigonid cusp demonstrating a clear sign of attrition, especially pronounced in the m1. The other two trigonid cusps, the protoconid and the metaconid, are subequal in height (Fig. 4B₂–D₂ and B₄–D₄). The protoconid is labially expanded at its base, making it the most robust cusp in the trigonid (Fig. 4B₂–D₂ and B₅–D₅). The anteroposterior length of the protoconid is the longest in the m1 and becomes successively thinner in the posterior molars (Fig. 4B₁–D₁).

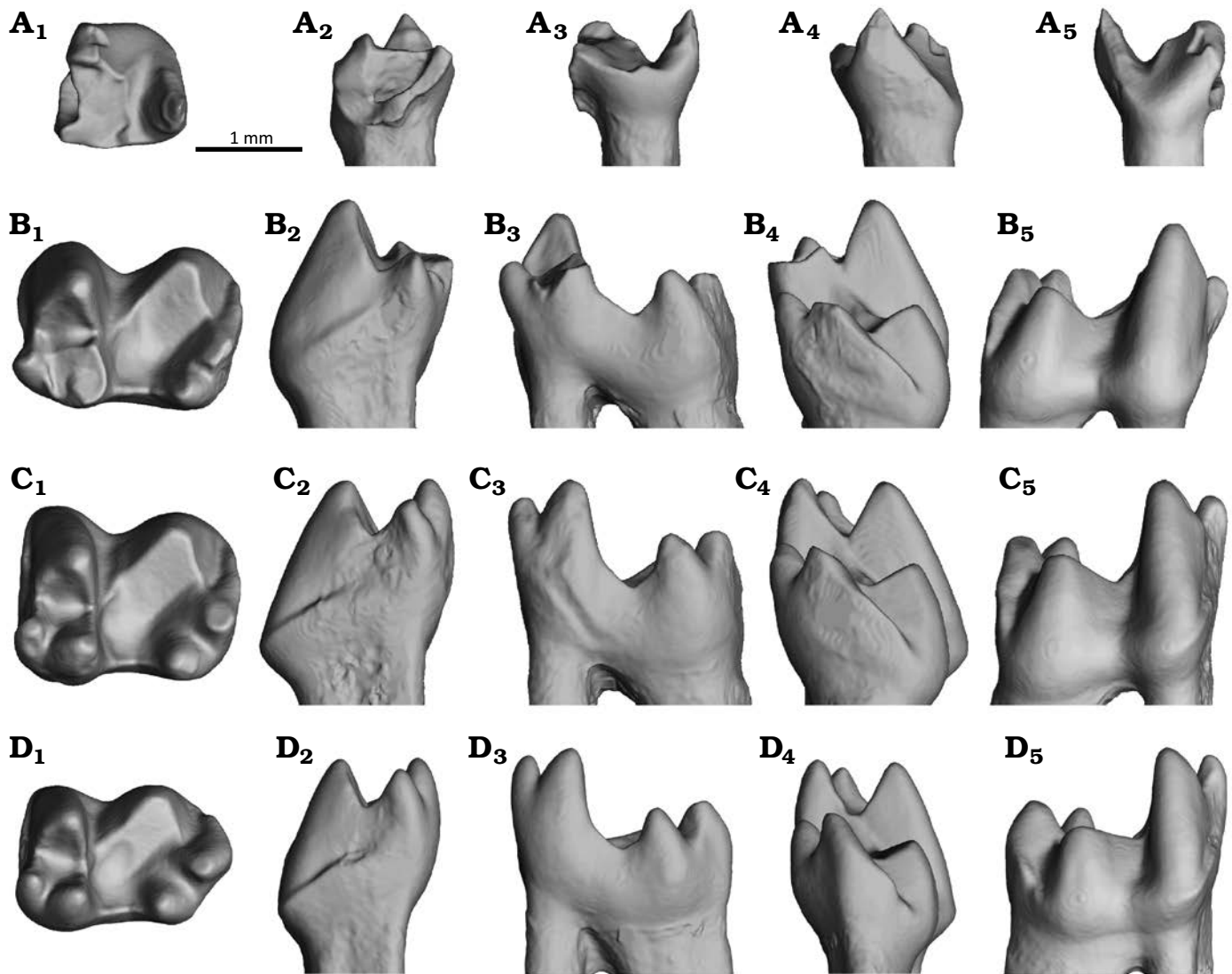


Fig 4. 3D model of ultimate premolar (A) and molars (B, m1; C, m2; D, m3) of the zhelestid eutherian mammal *Ravjaa ishiii* gen. et sp. nov. from the Upper Cretaceous Bayanshiree Formation, Dornogovi, Mongolia. Tooth in occlusal (A₁–D₁), mesial (A₂–D₂), lingual (A₃–D₃), distal (A₄–D₄), and labial (A₅–D₅) views.

Compared to the protoconid, the metaconid is slightly less developed and lacks lingual expansion at its base, though it is still more robust than the paraconid (Fig. 4B₃–D₃ and B₄–D₄). The paraconid and the metaconid are separated by a prominent gap (Fig. 4B₁–D₁), which tends to be more prominent in the posterior molars. The trigonid basin is V-shaped in anterior and posterior views (Fig. 4B₄–D₄). The protocristid is oriented transversely in the m2 and the m3, while it is slightly oblique in the m1 (Fig. 4B₁–D₁). The talonid width is subequal to the trigonid, and the talonid length accounts for more than half of the total length of the molar. Among the talonid cusps, the hypoconid is the most robust and the shortest cusp (Fig. 4B₁–D₁ and B₅–D₅). The attrition surface of the hypoconid is sloped lingually toward the talonid basin (Fig. 4B₄–D₄). The talonid basin of the m3 bears a faint spherical depression (Fig. 4D₁). The hypoconulid and the entoconid are in close approximation to each other (Fig.

4B₁–D₁), as in all other zhelestids (Archibald and Averianov 2012; Averianov et al. 2014). These two cusps are subequal in size, although the hypoconulids of the m1 and m2 are slightly worn mesiolabially (Fig. 4B₁–D₁). The hypoconulid of the m3 is more massive than that of the m1 and the m2 and markedly procurved (Fig. 4D₃ and D₅). The cristid obliqua emerges from the labial margin of the hypoconid and attaches slightly labially to the notch of the protocristid (Fig. 4B₁–D₁). The posterocristid is distinctly present between the hypoconid and the hypoconulid, while it is absent between the hypoconulid and the entoconid. The precingulid is present anterior to the trigonid in all molars and is more prominent in the posterior molars. The labial and lingual margins of the molars are smooth and lack a defined cingulid as in the other zhelestids except *Zhalmouzia bazhanovi*, in which the molar exhibits faint labial and lingual cingulids (Averianov et al. 2014). A prominent labial postcingulid is present posterolabial to

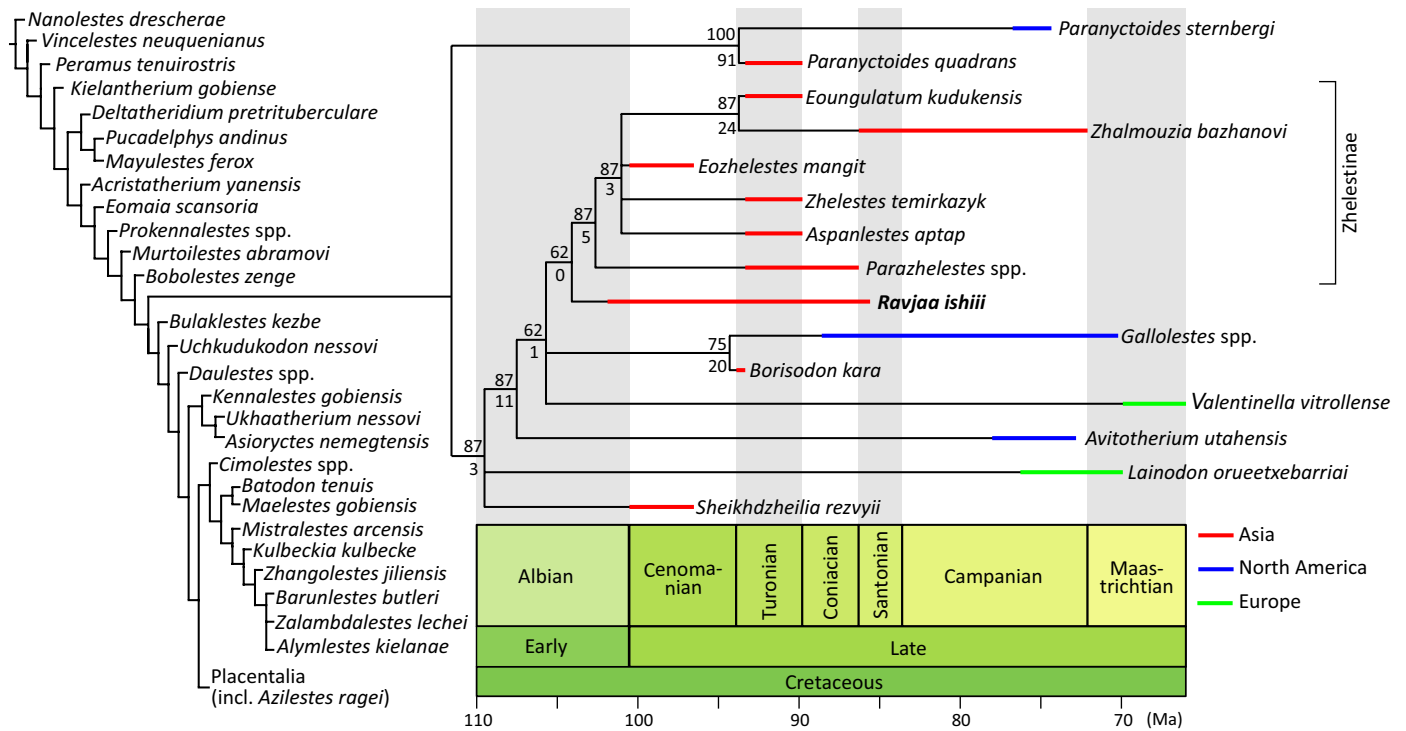


Fig 5. 50% majority-rule consensus tree of the phylogenetic analysis in this study. Numbers on nodes indicate the percentage of MPTs that contain a particular clade (above) and bootstrap supports (below). The figure provides a visualization of the fossil stratigraphic range within the Zhelestidae + *Paranyctoides* clade, with branching ages and branch lengths shown schematically. For further details see SOM 4.

the talonid and bears a cusp-like tubercle protruding from the hypoconulid base; this structure might be comparable to the hypocingulid, the term coined in Van Valen (1993).

Measurements (in mm).—m1: length 2.05; trigonid length 0.96, width 1.53; talonid length 1.09, width 1.55. m2: length 2.17; trigonid length 0.84, width 1.58; talonid length 1.33, width 1.59; distance between paraconid and metaconid 0.190. m3: length 2.09; trigonid length 0.76, width 1.43; talonid length 1.33, width 1.37; distance between paraconid and metaconid 0.19.

Stratigraphic and geographic range.—Type horizon and locality only.

Phylogenetic analysis

We conducted cladistic analyses to demonstrate the phylogenetic position of the new taxon within Zhelestidae (Fig. 5). According to Wible et al. (2009), which defined Zhelestidae as the clade formed by *Sheikhdzheilia rezvyii*, *Zhelestes termirkazyk*, and all their descendants, we treated this clade as the smallest clade containing *Sheikhdzheilia rezvyii* and *Zhelestes termirkazyk*. Additionally, Zhelestinae was treated as the smallest clade containing *Aspanlestes aptap*, *Parazhelestes* spp., and *Eoungulatum kudukensis*, according to Archibald and Averianov (2012), which listed these three genera as the included taxa of Zhelestinae. The phylogenetic analyses resulted in 64 of the most parsimoni-

ous trees (MPTs) with 2784 steps, each with a consistency index of 0.219 and a retention index of 0.570. The strict consensus tree is poorly resolved (SOM 2); early-branching eutherians such as zhelestids and some asioryctids are polyphyletic, as in the strict consensus trees of Tabuce et al. (2013) and Gheerbrant and Teodori (2021). The 50% majority-rule consensus tree demonstrates a better-resolved relationship, recovering the new taxon as the sister taxon of Zhelestinae within Zhelestidae (Fig. 5; see SOM 3 for the full majority consensus tree). *Paranyctoides sternbergi* Fox, 1979, and *Paranyctoides quadrans* Nessov, 1982, are recovered as a sister clade of Zhelestidae as in most recent works (e.g., Archibald and Averianov 2012; Gheerbrant and Teodori 2021) other than Averianov and Archibald (2016) in which *Paranyctoides* spp. was nested within Zhelestidae. It should be noted that two taxa previously reported as possible zhelestids, *Mistralestes arcensis* (Tabuce et al. 2013) and *Azilestes ragei* (Gheerbrant and Teodori 2021), are not recovered as zhelestids in the present analysis. Other than these differences, the topology within the Zhelestidae is congruent with the recent works (e.g., Archibald and Averianov 2012; Averianov et al. 2014). In our phylogenetic analyses, Zhelestidae is supported by six unambiguous synapomorphies, and the close approximation of the hypoconulid to the entoconid (120:3) is scored for the new taxon among the synapomorphies. The clade [Zhelestinae + *Ravjaa ishiii*] is supported by a single synapomorphy, absence of the “coronoid” facet (156:1), and *Ravjaa ishiii* sp. nov. has a single autapomorphy, the absence of the labial mandibular foramen

(139:0), which is broadly seen among placental mammals. *Eozhelestes mangit* has been suggested to be a non-zhelestine zhelestid in Averianov et al. (2014) but recovered as a member of Zhelestinae in our analysis. Zhelestinae is supported by two shared synapomorphies: the ultimate lower premolar talonid as wide as the anterior portion (57:0) and the short and erect hypoconulid of the ultimate molar (121:0).

Discussion

Our phylogenetic analyses recovered the new specimen within the Zhelestidae as the single sister taxon of the Zhelestinae (Fig. 5). Although the branch supports of the recovered tree are generally weak, the topology is well resolved compared to those of recent studies focused on zhelestid phylogeny (Tabuce et al. 2013; Gheerbrant and Teodori 2021), and attribution of the new specimen within Zhelestidae is supported by possession of a zhelestid synapomorphy (120:3). Among Zhelestidae, *Ravjaa ishiii* sp. nov. is unique in the large molars relative to the horizontal ramus of the dentary, the concave dorsal margin of the dentary, the absence of the labial mandibular foramen posterior to the coronoid crest of the dentary, and the undeveloped coronoid crest of the ascending ramus of the dentary. Together with the phylogenetic autapomorphy (139:0), the data provided here demonstrate that new specimens cannot be assigned to any known taxon and represent the new taxon within Zhelestidae.

The new taxon, *Ravjaa ishiii*, presented herein, is the first zhelestid from the Bayanshiree Formation and in fact the first discovered in Mongolia. The Upper Cretaceous strata in the Gobi Desert of Mongolia are known for extraordinarily well-preserved mammalian fossils and contribute to understanding the evolutionary history of the Asian Mesozoic mammals (Kielan-Jaworowska et al. 2000), but they have been mainly recovered from the overlying aeolian Djadokhta and Baruungoyot formations (Shuvalov 2000). Up to this study, a single dentary of a metatherian mammal *Tsagandelta dashzevegi* (Rougier et al. 2015) and a fragmentary skull of a eutherian mammal *Bayshinoryctes shuvalovi* (Lopatin and Averianov 2023) are the only known mammalian specimens from the Bayanshiree Formation. The occurrence of this new zhelestid further suggests the higher mammalian diversity in the Bayanshiree Formation than previously known.

The new occurrence of zhelestid reported here may suggest the environmental preference of this clade and/or ecological competition with the multituberculates as proposed in Nessov et al. (1998). Although only three mammalian taxa are known so far from the Bayanshiree Formation, none of the mammalian taxa from the Bayanshiree Formation is a multituberculate, which dominates the mammalian fauna in the overlying Djadokhta and Baruungoyot formations (Kielan-Jaworowska and Nessov 1992). The lack of zhelestids in the Djadokhta and Baruungoyot formations is in striking contrast with the mammalian fauna of southwestern Asia, which is characterized by abundant zhelestids

and rare multituberculates (Kielan-Jaworowska 1992). The localities of southwestern Asia, dominated by zhelestids, are mostly represented by the Bissekty and Aitym formations of Uzbekistan, deposited in a semi-humid coastal plain and marginal marine settings (Nessov et al. 1998; Redman and Leighton 2009). Contrarily, the multituberculate-dominated Djadokhta and Baruungoyot formations are interpreted to have been deposited under semi-arid to arid conditions in continental inland (Jerzykiewicz et al. 1993; Eberth 2018). The paleoenvironmental condition of the Bayanshiree Formation has been generally considered to be less arid than the aeolian-dominated Baruungoyot and Djadokhta formations (Jerzykiewicz and Russell 1991; Jerzykiewicz et al. 1993). This paleoenvironmental interpretation can be supported by the fluvial-dominated sequence described in this study, indicating the Bayan Shiree locality was generally in close proximity to a water resource compared to the aeolian-dominated Djadokhta and Baruungoyot localities. Thus, the new occurrence of zhelestid in the Bayanshiree Formation suggests that the paleoenvironmental condition allowed zhelestid to extend their habitat to the continental inland area.

Although the occurrence of zhelestid in the Bayanshiree Formation might support the habitat preference hypothesis of this clade, the anatomical feature of *Ravjaa ishiii* sp. nov. could also support the ecological competition hypothesis between zhelestids and multituberculates. The features such as the moderately compressed trigonid, the short protoconid, and more bulbous cusps seen on *Ravjaa ishiii* sp. nov. and most of the other zhelestids have been interpreted as reflecting more herbivorous tendency (Archibald 1996; Rose 2006) relative to the majority of Cretaceous therian mammals, which are generally considered to be insectivores with the developed trigonid, the tall protoconid, and the sharp cusps (e.g., Kielan-Jaworowska et al. 2004; Martin et al. 2020; Popowics and Mulimani 2023). This interpretation is further supported by morphometric analysis of the lower molar in early therian mammals from the K/Pg boundary, which suggests that zhelestid molar morphology reflects a frugivore/omnivore region, and it is noted that potentially contributes to increased functional diversity in the early Late Cretaceous (Grossnickle and Newham 2016). Since the majority of multituberculates have been generally thought to be herbivores (e.g., Wilson et al. 2012; Xu et al. 2015; Yuan et al. 2013), the inferred herbivorous tendency in *Ravjaa ishiii* sp. nov. suggests this taxon could have been an ecological competitor of multituberculates.

Ravjaa ishiii sp. nov. potentially represents the oldest member among zhelestids or as old as the currently known oldest zhelestids collected in Uzbekistan (Fig. 5), suggesting the emergence of this clade occurred around the Early/Late Cretaceous boundary to early Late Cretaceous. This period coincides with the early stage of the Cretaceous Terrestrial Revolution (KTR), a macroevolutionary event in the rapid diversification of various terrestrial animals in response to the radiation of angiosperms (e.g., Lloyd et al. 2008; Benton et al.

2022; Weaver et al. 2024). Through this event, angiosperms have been suggested to provide nutritious dietary resources to terrestrial ecosystems as leaves, flowers, fruits, and nuts (Weaver et al. 2024). The occurrence of potential angiosperm megafossils (Khand et al. 2000), including fruit fossils (Ksepka and Norell 2006) from a Bayanshirenian adjacent locality, suggests nutritious angiosperms indeed comprised the ecosystem that *Ravjaa ishiii* sp. nov. inhabited. Thus, the inferred herbivorous tendency in *Ravjaa ishiii* sp. nov. and zhelestids might have been an adaptation to an increase of angiosperms in the terrestrial ecosystem.

Conclusions

This study reports the new zhelestid specimen from the Bayanshiree Formation at Bayan Shiree. The zhelestid affinity of this species is supported by the subequally tall protoconid and metaconid, closely approximated hypoconulid and entoconid. Furthermore, this specimen shows unique characters among zhelestids, such as the faint coronoid crest, the concave dorsal alveolar margin, and the complete absence of the labial mandibular foramen. Thus, we erect a new taxon, *Ravjaa ishiii* based on the specimen. This species represents the first zhelestid from the Mongolian Late Cretaceous strata, suggesting a more prosperous mammalian diversity of the Bayanshiree formation than previously known and bringing potential paleobiogeographic and ecological insights of zhelestids. Zhelestids, including *Ravjaa ishiii* sp. nov., preferred relatively humid conditions, had an herbivorous tendency, and the evolution of this clade might have been tied to the KTR. These insights, however, should be thoroughly tested in the future, especially with more discoveries of mammalian fossils of the Mongolian Late Cretaceous strata, updated sedimentological data with the paleoenvironmental reconstruction of the Bayanshiree Formation, which is yet to be documented in detail, and thorough dietary reconstructions as performed in recent studies (Grossnickle and Newham 2016; Grossnickle et al. 2021).

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Appendix 1

Character scores for *Ravjaa ishiii* gen. et sp. nov., based on the matrix of Averianov and Archibald (2016).

1(0), 2(0), 57(0), 58(0), 61(1), 62(0), 63(0), 106(0), 107(0), 108(0), 109(0), 110(0), 111(2), 112(1), 113(1), 114(1), 115(1), 116(2), 117(1), 118, (2), 119(2), 120(3), 121(1), 122(2), 123(0), 124(0), 125(0), 126(1), 127(0), 130(2), 131(0), 132(1), 135(1), 136(1), 138(0), 139(0), 154(1), 156(1)