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Morphology of an Early Oligocene beaver *Propalaeocastor irtyshensis* and the status of the genus *Propalaeocastor*

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The Early to Late Oligocene *Propalaeocastor* is the earliest known beaver genus from Eurasia. Although many species of this genus have been described, these species are defined based on very fragmentary specimens. *Propalaeocastor irtyshensis* from the Early Oligocene Irtysh River Formation in northwestern Xinjiang, China is one of the earliest-known members of *Propalaeocastor*. This species is defined on a single maxillary fragment. We revise the diagnosis of *P. irtyshensis* and the genus *Propalaeocastor*, based on newly discovered specimens from the Irtysh River Formation. The dental morphology of *P. irtyshensis* is very similar to other early castorids. The caudal palatine foramen of *P. irtyshensis* is situated in the maxillary-palatine suture. This is a feature generally accepted as diagnostic character for the castorids. On the other hand, *P. irtyshensis* has two upper premolars, a rudimentarily developed sciuriform-like zygomatic plate, and a relatively large protrogomorph-like infraorbital foramen. Some previous researchers suggested that *Propalaeocastor* is a junior synonym of *Steneofiber*, while others took it as a valid genus. Our morphological comparison and phylogenetic analysis suggest that *Propalaeocastor* differs from *Steneofiber* and is a valid genus. We also suggest that *Agnotocastor aubekerovi*, *A. coloradensis*, *A. galushai*, *A. readingi*, *Oligotheriomys primus*, and “*Steneofiber aff. dehmi*” should be referred to *Propalaeocastor*. *Propalaeocastor* is the earliest and most basal beaver. The origin place of *Propalaeocastor* and castorids is uncertain. The Early Oligocene radiation of castorids probably is propelled by the global climate change during the Eocene-Oligocene transition.

1 Title:

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3 **genus *Propalaeocastor***

4

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21 ABSTRACT

22 The Early to Late Oligocene *Propalaeocastor* is the earliest known beaver genus from Eurasia.
23 Although many species of this genus have been described, these species are defined based on
24 very fragmentary specimens. *Propalaeocastor irtyshensis* from the Early Oligocene Irtys River
25 Formation in northwestern Xinjiang, China is one of the earliest-known members of
26 *Propalaeocastor*. This species is defined on a single maxillary fragment. We revise the diagnosis
27 of *P. irtyshensis* and the genus *Propalaeocastor*, based on newly discovered specimens from the
28 Irtys River Formation. The dental morphology of *P. irtyshensis* is very similar to other early
29 castorids. The caudal palatine foramen of *P. irtyshensis* is situated in the maxillary-palatine
30 suture. This is a feature generally accepted as a diagnostic character for castorids. On the other
31 hand, *P. irtyshensis* has two upper premolars, a rudimentarily developed sciuriform-like
32 zygomatic plate, and a relatively large protrogomorph-like infraorbital foramen. Some previous
33 researchers suggested that *Propalaeocastor* is a junior synonym of *Steneofiber*, while others
34 have taken it as a valid genus. Our morphological comparison and phylogenetic analysis suggest
35 that *Propalaeocastor* differs from *Steneofiber* and is a valid genus. We also suggest that
36 *Agnotocastor aubekerovi*, *A. coloradensis*, *A. galushai*, *A. readingi*, *Oligotheriomys primus*, and
37 “*Steneofiber aff. dehmi*” should be referred to *Propalaeocastor*. *Propalaeocastor* is the earliest
38 and most basal beaver. The place of origin of *Propalaeocastor* is uncertain, but the origin of the
39 castorids is likely to be North America. The Early Oligocene radiation of castorids was probably
40 propelled by the global climate change during the Eocene-Oligocene transition.

41

42 INTRODUCTION

43 Extant and fossil beavers are medium to large body-sized semi-aquatic, terrestrial or burrowing
44 rodents (Rybczynski, 2007; Flynn & Jacobs, 2008). Extant beavers include one genus and two
45 species (*Castor fiber* and *C. canadensis*). Fossil beavers are much more diverse, including at
46 least twenty seven genera and more than one hundred species (McKenna & Bell, 1997; Korth &
47 Samuels, 2015; Mörs, Tomida & Kalthoff, 2016; <https://www.paleobiodb.org/>). It is generally
48 accepted that all beavers represent a monophyletic family: Castoridae (McKenna & Bell, 1997;
49 Helgen, 2005; Rybczynski, 2007). Castoridae is closely related to the extinct family
50 Eutypomyidae, and the two families are usually referred to the superfamily Castoroidea
51 (Simpson, 1945; Wood, 1955, 1965; Hugueney, 1999; Flynn & Jacobs, 2008). Within crown
52 rodents, phylogenetic analyses based on molecular data and/or morphological data usually
53 support the sister-group relationship between the castorids and the geomyoids (a superfamily of
54 rodents that contains the pocket gophers, the kangaroo rats and mice (e.g., Douady et al., 2000;
55 Adkins et al., 2001; Adkins, Walton & Honeycutt, 2003; Murphy et al., 2001; Huchon et al.,
56 2002; Montgelard et al., 2002; Fabre et al., 2012).

57

58 The earliest-known castorid fossil, “*Agnotocastor*” *galushai*, was discovered from the South
59 Fork of Lone Tree Gulch of Wyoming (Emry, 1972). The age of the locality is middle to late
60 Chadronian of North American Land-Mammalian Ages (NALMA) within a precision $^{206}\text{Pb}/^{238}\text{U}$
61 zircon dates from 35.805 ± 0.076 Ma to 34.398 ± 0.022 Ma (Emry & Korth, 2012; Sahy et al.,

62 2015). The dental and cranial morphology of *Agnotocastor* shares many similarities with the
63 eutypomysid *Eutypomys* (Wilson, 1949a; Wood, 1965; Wahlert, 1977; Xu, 1995, 1996; Flynn &
64 Jacobs, 2008). The earliest-known beavers outside of the North America belong to the genus
65 *Propalaeocastor* Borissoglebskaya, 1967 (Misonne, 1957; Borissoglebskaya, 1967; Lytschev,
66 1970; Kretzoi, 1974; Bendukidze, 1993; Lytschev & Shevyreva, 1994; Wu et al., 2004;
67 Bendukidze et al., 2009).

68 The validity of *Propalaeocastor* is debatable. The type species, *P. kazakhstanicus*, is from
69 the Early Oligocene of Kyzylkak, Dzhezkazgan, Kazakhstan (Borissoglebskaya, 1967). Lytschev
70 & Shevyreva (1994), and Lopatin (2003, 2004) considered *Propalaeocastor* as a junior synonym
71 of *Steneofiber* Geoffroy Saint-Hilaire, 1833. Some other researchers did not agree and suggested
72 that *Propalaeocastor* is different from *Steneofiber* and is a valid genus (McKenna & Bell, 1997;
73 Korth, 2002; Wu et al., 2004). Kretzoi (1974) referred “*Steneofiber*” *butselensis* Misonne, 1957
74 to a new genus “*Asteneofiber*”. However, the validity of *Asteneofiber* was not widely recognized.
75 Some researchers considered “*Asteneofiber*” as the junior synonym of *Steneofiber* (McKenna &
76 Bell, 1997; Korth, 2002), while Wu et al. (2004) regarded “*Asteneofiber*” as a junior synonym of
77 *Propalaeocastor*.

78 There are quite a few species attributed to *Propalaeocastor*, but the species attribution of
79 this genus is ambiguous, because all of the species are represented by isolated teeth and/or jaw
80 fragments. Besides the type species *Propalaeocastor kazakhstanicus*, Borissoglebskaya (1967)
81 also named *P. habilis* in the same paper. In their study of beaver remains from Maylibay of
82 Zaissan (or Zaysan) Basin, Kazakhstan, Lytschev & Shevyreva (1994) synonymized *P. habilis*
83 with *P. kazakhstanicus* and reported another three species: *P. shevyrevae*, *P. aff. shevyrevae* and
84 *P. zaissanensis*. Wu et al. (2004) recognized *P. butselensis*, *P. shevyrevae*, *P. sp. aff. P.*
85 *shevyrevae*, *P. zaissanensis*, *P. kazakhstanicus*, and named the species *P. irtyshensis*. Lopatin
86 (2003) suggested that “*Capacikala sajakensis*” is the junior synonym of “*Steneofiber*”
87 *kumbulakensis*. Bendukidze et al. (2009) synonymized “*Capacikala sajakensis*” to “*Capatanca*”
88 *schokensis*, and transferred “*Capatanca*” *schokensis* Bendukidze, 1993 and “*Steneofiber*”
89 *kumbulakensis* Lytschev, 1970 to *Propalaeocastor*.

90 Because of the impoverishment of specimens and ambiguous generic diagnosis, the
91 systematic position of *Propalaeocastor* is also in doubt. It has been assigned to the tribe
92 Anchitheriomini by Korth (2001), the subfamily Anchitheriomyinae by Korth (2004) and tribe
93 Minocastorini by Mörs et al. (2016). The handful of dental specimens of *Propalaeocastor* exhibit
94 a pattern resembling both *Agnotocastor* and *Eutypomys*. For instance, one of the
95 *Propalaeocastor* species (*P. kumbulakensis* Lytschev, 1970) was even considered a member of
96 *Eutypomys* (Xu, 1996).

97 To clarify the validity and species attribution of *Propalaeocastor*, we report a few newly
98 discovered specimens of *P. irtyshensis* from the Early Oligocene Irtys River Formation in
99 Xinjiang, China. These specimens make *P. irtyshensis* the best-known species of
100 *Propalaeocastor*. We examine the dental features of most of the castorid genera, and develop a
101 data matrix for phylogenetic analysis. Based on the newly collected specimens and the results of
102 our phylogenetic analysis on castorids, we are able to emend the generic diagnosis of

103 *Propalaeocastor* and clarify the phylogenetic relationships among *Propalaeocastor*,
104 *Agnotocastor*, *Eutypomys* and other early beavers.

105

106 GEOLOGIC SETTING

107 Cenozoic sediments are widely exposed in the drainage area of the Irtysh (=Ertix) River in
108 Burqin-Jeminay region in northwestern Xinjiang of China (Figs. 1A & B). *Propalaeocastor*
109 *irtyshensis* was discovered from the lower portion of the Early Oligocene Irtysh River Formation
110 at the XJ200203 locality in the Burqin-Jeminay region (Fig. 1B) (Wu et al., 2004; Stidham et al.,
111 2015). Only upper dentition was previously known. The new specimens of *P. irtyshensis*
112 reported here were discovered from a new fossiliferous locality of the lower Irtysh River
113 Formation about 50 km southwest to the XJ200203 locality. The Irtysh River Formation is a set
114 of fluviolacustrine mudstone, siltstone, sandstone and thick conglomerate. The fossiliferous layer
115 of the Irtysh River formation is dated as 32.0 Ma (Sun et al., 2014). The same fossiliferous layer
116 at the XJ200203 locality can be traced to the new locality despite the long distance between the
117 two localities. This fossiliferous layer at the new fossil locality is an approximately 5-meter thick
118 bed of grey greenish and light brown-reddish mudstone with rich calcareous nodules (Fig. 1C).
119 The new *P. irtyshensis* remains include a fragmentary maxilla, several incomplete jaws and
120 isolated cheek teeth. The small mammals associated with these new beaver fossils include
121 *Cricetops dormitor*, *Parasminthus tangingoli*, *Cyclomytus lohensis*, and *Prosciurus* sp. These
122 small mammals are also present at the XJ200203 locality (Ni et al., 2007; Sun et al., 2014).

123

124 MATERIALS, METHODS AND ABBREVIATIONS

125 The new materials include a broken maxilla preserving P4-M1, two isolated upper cheek teeth
126 and three mandibular fragments. The holotype of *Propalaeocastor irtyshensis* (IVPP V 13690) is
127 re-described. All fossils are housed at the Institute of Vertebrate Paleontology and
128 Paleoanthropology, Chinese Academy of Sciences, Beijing. The specimens were CT-scanned
129 using the 225 kV Micro-CT at the Key Laboratory of Vertebrate Evolution and Human Origins,
130 Chinese academy of Sciences. Segmentations and 3D virtual reconstructions were made
131 following the standard procedure introduced by Ni et al. (2012). Specimens were measured using
132 an Olympus SZX7 microscope and mandibles by vernier caliper both with a precision of 0.01
133 mm. The length is defined as the mesiodistal chord. The width is defined along the chord
134 perpendicular to the length. For incisors, the same standard is used to define the length and width.

135 The dental terminology (Figs. 2, 3) is modified from Stirton (1935), Huguency (1975, 1999),
136 Lopatin (2003), and Wu et al. (2004). We use “-loph” and “-lophid” for the major ridges or crests,
137 and “-lophule” and “-lophulid” for the thin, short spur-like ridges that are developed from the
138 lophs and lophids. The major change is that we abandon the use of terms “mesoloph” and
139 “mesolophid” in castorids. The mesoloph and mesolophid are usually defined as “crest from
140 mesocone(id) toward the lingual or buccal side of the tooth.” (Wood & Wilson, 1936). The
141 mesocone and mesoconid are distinctly present in *Eutypomys*, and the mesoloph and mesolophid
142 are clearly derived from the mesocone and mesoconid, respectively. In beavers, however, the
143 mesocone and mesoconid are absent. The so-called “mesoloph(id)” is derived from the posterior

144 arm of the protocone(id). Here we treat the so-called “mesoloph” and “mesolophid” as protoloph
145 II and metalophid II, respectively. The dental cusp-ridge connections of the *Eutypomys*,
146 *Agnotocastor*, *Propaleocastor*, and other early beavers are very complicated, i.e. their ridges are
147 normally irregular and wrinkled with variable valleys or enamel islands. We use the term “mass”
148 to describe this complex status, including paracone mass, metacone mass, metaconid mass, and
149 entoconid mass. The suffixes flexus/flexid, fossette/fossettid and stria/striid are used for
150 describing the valleys between two loph/lophids or between two cusps. Flexus and flexid are
151 used when the valleys are open to the tooth sides, usually in relatively unworn specimens. Stria
152 and striid refer to the notches running down the tooth crown in buccal or lingual view. These
153 notches are the buccal or lingual openings of the valleys. As the tooth wear deepens, the flexus
154 or flexid will be gradually closed near the tooth sides. These closed flexus or flexids are called
155 fossettes or fossettids. Paraflexid/fossettid/striid and metaflexid/fossettid/striid were often used
156 for the mesial and distal valleys respectively (Stirton, 1935; Hugueney, 1975, 1999; Wu et al.,
157 2004). Here we followed Lopatin (2003) by using metaflexid/fossettid/striid for the mesial valley
158 and entoflexid/fossettid/striid for the distal flexid. We use premetafossettid instead of
159 proparafossettid (Hugueney, 1999) or parafossettid (Lopatin, 2003) to describe the small fossa
160 enclosed between anterolophid and metalophid I.

161 We developed a data matrix including 145 characters scored for 42 taxa. The 145 characters
162 comprise 120 dental and 25 cranial characters. *Marmota monax*, *Keramidomys fahlbuschi* and
163 *Eutypomys inexpectatus* were selected as outgroup taxa. Eutypomyids are widely considered as
164 the sister group of castorids (Korth, 1994; Rybczynski, 2007; Flynn & Jacobs, 2008)). *Marmota*
165 and *Keramidomys* have the same dentition formula as that in castorids, but the phylogenetic
166 relationship between these two taxa and castorids is probably further than that between castorids
167 and eutypomyids. The ingroup comprises 39 taxa, of which, only *Castor canadensis* is an extant
168 species. The data matrix was edited in Mesquite v3.2 software (Maddison & Maddison, 2017)
169 and saved in the NEXUS format. The scored specimens, and the definition and arguments for the
170 characters are listed in the NEXUS file (see Supplementary Information). Parsimony analysis
171 was undertaken using TNT, Tree analysis using New Technology, a parsimony analysis program
172 subsidized by the Willi Hennig Society (Goloboff et al., 2008). We ran multiple replications,
173 using sectorial searches, drifting, ratchet and fusing combined. Random sectorial search,
174 constraint sectorial search and exclusive sectorial search were used. Ten cycles of tree drifting,
175 10 cycles of ratchet and 10 cycles of tree fusing were performed in the search. Default parameter
176 settings for random sectorial search, constraint sectorial search, exclusive sectorial search, tree
177 drifting, ratchet and fusing were used. The search level was set as 10 for 42 taxa. Optimal scores
178 were searched with 10000 replications. Twenty-four characters are set as “ordered” (listed in the
179 Supplementary Information). The outgroups were not used as reference for ordering the
180 character states. We hypothesized that the states of these characters are addable. These addable
181 states can be observed in some chronologically succeeding castorid taxa. All characters have
182 equal weight. We used absolute Bremer Support and relative Bremer Support (Bremer, 1994;
183 Goloboff et al., 2001), calculated in TNT, to describe the stability of the phylogenetic result.
184 TNT script for running multiple replications, using sectorial searches, drifting, ratchet and fusing

185 combined, and script for calculating the Bremer Supports and Relative Bremer Supports were
186 adopted from Ni et al. (2013).

187 **Abbreviations:** **AMNH**, American Museum of Natural History; **CSC**, Chadron State
188 College; **FAM**, Frick American Mammals, Department of Vertebrate Paleontology, the
189 American Museum of Natural History; **IVPP**, Institute of Vertebrate Paleontology and
190 Paleoanthropology, Chinese Academy of Sciences; **UCM**, University of Colorado Museum; **XJ**,
191 prefix to Xijiang, field localities of the IVPP.

192

193 RESULTS

194

195 Systematic Paleontology

196 **Order** Rodentia, Bowdich, 1821

197 **Family** Castoridae Hemprich, 1920

198 **Genus** *Propalaeocastor* Borissoglebskaya, 1967

199 **Synonym.** *Asteneofiber* Kretzoi, 1974: p.427; *Oligotheriomys* Korth, 1998: p.127

200 **Type Species.** *Propalaeocastor kazachstanicus* (including *P. habilis*) Borissoglebskaya,
201 1967.

202 **Included Species.** *P. coloradensis* (Wilson, 1949b); *P. butselensis* (Misonne, 1957), *P.*
203 *kumbulakensis* Lytshev, 1970, *P. galushai* (Emry, 1972), “*Steneofiber* aff. *dehmi*” (in Hugueneu,
204 1975), *P. aubekerovi* (Lytshev, 1978), *P. readingi* (Korth, 1988), *P. schokensis* (Bendukidzes,
205 1993), *P. shevyreva* (Lytshev & Shevyreva, 1994), *P. sp. aff. P. shevyreva* (Lytshev &
206 Shevyreva, 1994), *P. zaissanensis* (Lytshev & Shevyreva, 1994), *P. primus* (Korth, 1998) , and *P.*
207 *irtyshensis* Wu et al., 2004.

208 **Distribution.** Early to Late Oligocene, Eurasia; Late Eocene to Early Oligocene, North
209 America.

210 **Emended Diagnosis.** A small-sized castorid. Dental formula: 1/1, 0/0, 2/1, 3/3. Zygomatic
211 process of maxilla forming a sloping surface. Infraorbital foramen large. Infraorbital canal short.
212 Sciurognathous lower jaw. Digastric eminence present in some advanced species. Lower incisor
213 enamel surface smooth, mediolaterally convex, and lacking enamel ornamentation. Lower
214 incisor root terminating in a lateral capsule. Wide space present between lower tooth row and
215 vertical ramus. Cheek teeth unilaterally mesodont. Upper cheek tooth crown nearly quadrate. P3
216 present. P4 slightly larger than M1 and M2. M3 being the smallest. Upper cheek teeth presenting
217 complicated paracone mass and metacone mass. Premesoflexus and postmesoflexus always
218 present. Metaflexus buccally open. p4 mesiodistally elongated. Lower molar crown rectangular.
219 p4 larger than molars. m3 being the narrowest. Lower cheek teeth having complex metaconid
220 mass and entoconid mass. Premesofossettid present in some species. Postmesoflexid always
221 present. Metastylid crest present. Crown (Coronal) cementum absent.

222

223 *Propalaeocastor irtyshensis* Wu et al., 2004

224 (Figs. 4-7; Tables 1 & 2)

225 **Holotype.** IVPP V 13690, a right maxillary fragment preserving P4-M3. Locality XJ200203,

226 northwest of Burqin, Xinjiang. The Irtysh Formation, Early Oligocene.

227 **Referred specimens.** IVPP V 23138.1, a right maxillary fragment preserving P4-M1, IVPP
228 V 23138.2, an isolated left P4, and IVPP V 23138.3, an isolated left M1, probably belong to the
229 same individual; IVPP V 23139, a right dentary fragment preserving p4-m3; IVPP V 23140, a
230 right dentary fragment preserving p4-m1; IVPP V 23141, a right dentary fragment preserving p4.

231 **Localities and Horizon.** Northeast of Jeminay County, Junggar Basin, Xinjiang (Fig. 1B).
232 Irtysh River Formation, Early Oligocene.

233 **Emended Diagnosis.** P3 present. Infraorbital foramen large, infraorbital canal short.
234 Differing from *P. kazachstanicus* in having greater mandibular depth beneath p4, complete
235 endoloph and open postmesoflexus on P4, two premesofossettids and more transverse
236 mesoflexid on lower cheek teeth, and in lacking digastric eminence. Different from *P.*
237 *butselensis* in having more complicated septa or spurs in buccal premesoflexus, metaflexus and
238 premesofossettid, more distally extending mesoflexus. Different from *P. kumbulakensis* in
239 having smaller size, lower tooth crown, less distally extended mesoflexus, closed postmesoflexus
240 on P4, and two premesoflexids on p4. Differing from *P. zaissanensis* in having separated
241 hypoflexus and mesoflexus on M3. Differing from *P. schokensis* in having less massive paracone
242 mass and metacone mass, and in lacking metalophule I on upper cheek teeth. Differing from *P.*
243 *aubekerovi* by lacking digastric eminence and having greater mandibular depth beneath p4.
244 Differing from *P. readingi* in having more transversely expanded m1 and m2. Differing from *P.*
245 *shevyrevae* in having lower tooth crown, less folded inner surface of enamel islets, and in lacking
246 premetafossettids and having double premesofossettids on p4, and less elongated m3 lacking
247 septum in entofossettids. Differing from *P. primus* in having smaller size and lower tooth crown.

248 **Measurements.** See Tables 1 & 2.

249 **Description.** The two maxillary fragments (V 13690, holotype and V 23138.1) preserve a
250 part of the palatine process, a part of the alveolar process, and a part of the zygomatic process.
251 The alveolar process forms the tooth sockets and holds the teeth. The dorsal side of the alveolar
252 process is flat and smooth. It does not show any bulges for the expansion of the tooth roots. On
253 its dorsal-medial side above the M2, it presents the opening of the caudal palatine foramen
254 (=dorsal palatine foramen), which leads to a canal running in the maxillary-palatine suture (Figs.
255 4A1-2). The preserved palatine process is very small. On V 13690, only the major palatine
256 foramen is well preserved. It is an oval and oblique opening situated between M1 and M2, and in
257 the suture between the palatine process of the maxilla and the palatine bone (Figs. 5A1-2). On V
258 23138.1, the broken surface shows that the major and minor palatine foramina (=paired posterior
259 palatine foramina) lead to short canals and meet at the caudal palatine foramen (Fig. 4A1). The
260 preserved zygomatic process of the maxilla is quite long. It extends dorsolaterally from a place at
261 the level of the mesial root of P4. The mesial surface of the zygomatic process slopes
262 rostrorodorsally, indicating that a narrow zygomatic plate probably is present (Figs. 5A1-2). No
263 masseteric tubercle for the superficial masseter is present on the root of the zygomatic process.
264 Dorsal to the zygomatic process, a round and smooth surface indicates that the infraorbital
265 foramen is probably large and round, and the infraorbital canal is very short (Figs. 5A3, B2).
266 Dorsoventrally, the infraorbital foramen and infraorbital canal are at the level of the tooth roots,

267 a situation as in extant protrogomorphous and sciurumorphous rodents.

268 On both V 13690 and V 23138.1, there is a small semi-cylindrical depression mesial to the
269 mesial roots of P4 (Figs. 5A1-2 & B1). This depression indicates the presence of a small single-
270 rooted P3. Because the M3 of both specimens were already erupted and moderately worn, this
271 small depression cannot be for the deciduous tooth. For a dP3, it should have more than one root.
272 On the mesial surface of the P4, no obvious contacting facet is present. It is probably because the
273 crown of P3 is very small and low, and has no tight contact with P4.

274 The lingual side of the upper cheek tooth crown is higher than the buccal side (Figs. 4, 5B1;
275 Table 1). From the mesial side to the distal side of the tooth row, the tooth size decreases
276 gradually. The lingual tooth cusps, namely protocone and hypocone, are distolingually expanded
277 and form two fold-like structures on each tooth. The buccal cusps (paracone and metacone) and
278 their accessory ridges form the complex paracone mass and metacone mass.

279 The P4 (Figs. 4A, B; Fig. 5A1) is the largest of the upper cheek teeth. Its occlusal surface
280 has an inverted trapezoid outline with its mesial side is wider than its distal side. The tooth can
281 be roughly divided into four regions: the protocone region on the mesiolingual side, the paracone
282 mass on the mesiobuccal side, the hypocone region on the distolingual side and the metacone
283 mass on the distobuccal side. The lingual sides of the protocone region and hypocone region are
284 separated by the deep and mesiobuccally directed hypoflexus. The buccal sides of those two
285 regions are connected by the strong and oblique endoloph. The paracone mass is separated from
286 the metacone mass by the deep mesoflexus. The protocone distolingual side is expanded and
287 forms a fold-like structure. The buccal side of the protocone has two arms, the mesial protocone
288 arm and the distal protocone arm, which merge with the anteroloph and endoloph respectively.
289 The parastyle is a very small cusp. It is well delimited as a small node situated mesial to the
290 paracone on a slightly worn specimens (V 23138.1-2). In the moderately worn specimen (V
291 13690), the parastyle is merged with the anteroloph. The paracone mass includes the paracone
292 and two protoloph. The lingual side of the paracone smoothly extends into the protoloph I
293 (mesial protoloph). In the less worn specimen (V 23138.1), the lingual end of the protoloph I
294 does not join the protocone and is separated from the latter by a shallow groove. In the slightly
295 more deeply worn specimens (V 13690, V 23138.2), the lingual end of the protoloph I connects
296 the mesial arm of protocone through the short protolophule I. The protoloph II (distal protoloph)
297 is a long and curved crest. Its buccal end extends to the distal side of the paracone (V 23138.1-2)
298 or merges with paracone (V 13690). Its lingual end connects the endoloph in two specimens (V
299 23138.1-2) through strong protolophule II, but is separated from the endoloph in the holotype (V
300 13690). On V13690, an extra fold is present at the distolingual side of protoloph II. Mesocone
301 and mesoloph are absent. Three small fossae/flexi are present in the paracone mass: including
302 paraflexus, lingual premesofossette and buccal premesofossette. Paraflexus is enclosed by
303 anteroloph and protolophI. Lingual premesoflexus is enclosed by protoloph I and protoloph II.
304 Buccal premesoflexus is enclosed by protoloph II and postparacrista. In the hypocone region, the
305 hypocone forms a fold, which is smaller than the protocone. The mesial and distal arms of
306 hypocone are smoothly merged with the endoloph and the posteroloph respectively. The
307 metacone mass normally develops three ridges including double metalophs (metaloph I and

308 metaloph II) and an extra mesial short ridge. The extra mesial ridge is the shortest, and
309 mesio buccally extends towards the protoloph II. The longest ridge is the metaloph I, which
310 transversely connects the metacone and hypocone. Distal to the metaloph I, there is a long ridge
311 referred as metaloph II here. This ridge extends distobuccally and always connects the
312 posteroloph via a short and thin ridge (metalophule II). The postmesoflexus, which lies between
313 the metaloph I and metaloph II, is always buccally open. The metaflexus lying in between the
314 metaloph II and posteroloph is divided into two or three fossae by small ridges. From the buccal
315 view, the tooth has 3 deep grooves, which are collectively called buccal striae. From the mesial
316 to the distal, the three buccal striae are named as the parastria, the mesostria and the metastria.
317 From the lingual view, the only deep groove generated by the hyoflexus is the hypostria. The
318 hypostria is the longest. It extends nearly two third of the crown height. The mesostria is the
319 second deepest groove that reaches about a half of the crown height. The parastria and the
320 opening of postmesoflexus are very short. P4 has a strong lingual root and two slim buccal roots.

321 The M1 (Figs. 4A, C; Fig. 5A1) has a more rectangular crown than the P4, but both teeth
322 have very similar cusp-ridge pattern. The width of M1 is larger than the length. In the paracone
323 mass, two protolophes are present. The paraflexus and the groove between the protoloph I and
324 protoloph II are worn into four enamel islets. The mesoflexus is a straight groove in V 23138.1
325 and V 23138.2. In the slightly more deeply worn specimen (V 13690) the groove is divided into
326 two parts by a longitudinal ridge. In the metacone mass, it develops three or four ridges. The
327 grooves between those ridges are divided into three or four fossae. As in the P4, the lingual side
328 of M1 has one deep groove (hypostria), and the buccal side of M1 has one (mesostria) or two
329 (mesostria and metastria) shallow grooves on moderately worn specimens (V 23138.1, .3) and
330 lacks a groove on the deeply worn specimen (V 13690). The M1 has one strong lingual root and
331 two slim buccal roots.

332 The M2 (Fig. 5A1) is very similar to the M1 in both size and cusp-ridge pattern. In its
333 paracone mass, there are two protolophes, two opened grooves and one enclosed enamel islet. The
334 mesial groove is the long and narrow paraflexus. The distal groove is the premesoflexus. A short
335 ridge divided the premesoflexus into an open groove on the buccal side and a small enamel islet
336 on the lingual side. The mesoflexus is a long and curved groove separating the paracone mass
337 and the metacone mass. In the metacone mass, there are also two metalophes. The metaloph I has
338 an indentation in its middle part. This indentation joins the postmesofossette with the fossa
339 between the metalophes. The metaloph II is a complete ridge that connects the metacone and
340 hypocone. From the distolingual part of the metaloph II, it develops a short spur protruding into
341 the metaflexus. The posteroloph is a strong ridge as a buccal extension of the distal arm of the
342 hypocone. The development of the striae on the lingual and buccal sides of M2 is identical to
343 those in M1 of the same dentition (V 13690).

344 M3 (Fig. 5A1) has a narrower distal edge than in the M1-2. The hypocone of M3 is
345 relatively small and the posteroloph is reduced. The paracone mass is almost identical to those in
346 M1 and M2, while the metacone mass is proportionally smaller.

347 The newly collected specimens include three mandibular fragments (Fig. 6). Two of the
348 three specimens preserve most of the horizontal ramus and a portion of the vertical ramus (V

349 23139 and V 23141). The other specimen preserves only a small part of the horizontal ramus (V
350 23140). The horizontal ramus of the mandible can further be divided into two parts: the part that
351 bears the incisor and the part that bears the premolar and molars. The part that bears the incisor
352 contains a long incisor alveolus, which runs beneath the premolar and molars and extends
353 distally and buccally to a point lateral and above the level of tooth crown. The tooth roots show
354 bulges on the lingual side of the mandible, and form the alveolar juga. The buccal surface of the
355 mandible is smooth. A large and round mental foramen is present at a place ventral to the p4
356 (Figs. 6A3, B3, C3). On the mesiolingual surface of the horizontal ramus of the mandible, an
357 oval rugose region mesioventral to the alveolus of p4 is identified as the caudoventral expansion
358 of the mandibular symphysis (Figs. 6A1, C1). Ventral to this rugose region, no digastric
359 eminence is present. On the ventral portion of the lingual side of the mandible, there are many
360 nutrient foramina. A small portion of the angular process of the mandible is preserved in two
361 specimens, and it extends caudoventrally (Fig. 6A1). On the lingual side of the vertical ramus,
362 the medial pterygoid muscle fossa is very deep. On the buccal surface of the vertical ramus, the
363 masseteric fossa is well defined by the masseteric crest. The dorsal and ventral branches of
364 masseteric crest are convergent nearly at a right angle, and extend to a point ventral to the m1.
365 The coronoid process of the vertical ramus arises lateral to the m1. It includes a lateral bulge that
366 contains the most distal extension of the incisor root. On the medial side of the coronoid process,
367 there is a well-developed ridge (Figs. 6A2, C2). This ridge probably marks the inferior limit for
368 the lateral pterygoid muscle. The space between the tooth row and the vertical ramus of the
369 mandible is broad.

370 The lower incisor is only preserved in one specimen (V 23141). The cross-section of this
371 lower incisor is in a rounded triangular shape. The pulp cavity is large and round. The enamel
372 band of the incisor is smooth and buccoventrally convex (Fig. 7D).

373 The buccal sides of the lower cheek tooth crowns are slightly higher than the lingual crown
374 side (Table 1). From p4 to m3, the sizes are gradually reduced. On all the cheek teeth, the
375 protoconid and hypoconid are large and mesiobuccally protruding. The metaconid and entoconid
376 and the ridges associated with them from the complicated metaconid mass and entoconid mass.

377 All the three mandibles preserve the fourth premolar (Figs. 6-7). The crown of the p4 (Figs.
378 6A2, B2, C2) has a trapezoid outline with its mesial side narrower than its distal side. The
379 hypoflexid and mesoflexid (=mesofossettid when its lingual side is closed) form a waist that
380 divides the tooth crown into mesial and distal lobes. The protoconid, the anterolophid and the
381 mesial part of the ectolophid are merged into a strong curved ridge that defines the buccal margin
382 of mesial lobe. The metaconid, the lingual part of the metalophid II, the metastylid and the
383 metastylid crest are fused into another curved ridge that forms the lingual margin of the mesial
384 lobe. In less worn individuals (V 23139, 23140), the cusps and ridges in the mesial lobe enclose
385 three fossae (Figs. 6A2, B2). In a deeply worn individual (V 23141), only one fossa is left (Fig.
386 5C2). The metaflexid (=metafossettid when its lingual side is closed), which is enclosed between
387 the anterolophid and metalophid II, is a long and curved groove (or fossa). The metalophid I is
388 present as a spur derived from the anterolophid and extends into the metafossettid. Between the
389 metalophid II and the metastylid crest, two fossae are present, namely the buccal

390 premesofossettid and the lingual premesofossettid (Figs. 6A2, B2). The distal side of the lingual
391 premesofossettid is open in one specimen (Fig. 6B2). The two fossae disappear in the heavily
392 worn specimen (V 23141, Fig. 6C2). The mesoflexid is a long and deep groove that extends
393 transversely across more than half of the crown width. The hypoflexid on the buccal tooth side
394 has a broad opening. It extends distolingually to the mesiolingual side of the hypoconid. The
395 hypoflexid and mesoflexid are separated by the ectolophid. The ectolophid also connects the
396 mesial and distal lobes. The distal lobe is formed by the hypoconid, entoconid and the ridges and
397 arms associated with those two cusps. The hypoconid is very large and forms the buccal half of
398 the distal lobe. The posterolophid, the entoconid, the hypolophids and the distal part of
399 ectolophid form the lingual half of the distal lobe. The mesial hypolophid (hypolophid I) and the
400 small postmesofossettid are present in the less worn specimen (Figs. 6B2). The entoflexid is
401 present as long groove between the distal hypolophid (hypolophid II) and posterolophid.
402 Complicated enamel folds developed from the hypolophid II and posterolophid protrude into the
403 entoflexid. In the deeply worn specimen (V 23141), these folds connect to each other and divide
404 the entoflexid into 3 enamel islets. Two broad roots are present on p4 (Figs. 7A2, B2, C2).

405 The m1 (Figs. 6A2, B2) is preserved on two specimens (V 23139, V 23140). Both of them
406 are heavily worn. The m1 has a rectangular crown, with its width larger than its length. As in the
407 p4, the conspicuously deep mesoflexid and hypoflexid form a waist and divide the tooth into
408 mesial and distal lobes. The protoconid, metaconid and the ridges associated with the mesial lobe
409 tend to merge together. One or two enamel islets are enclosed in the mesial lobe. It is hard to
410 deduce whether they homologize with the metafossettid or with the premesofossettid. The
411 mesoflexid is lingually open on V 23140 but closed on V 23139. The hypoflexid of m1 is
412 narrower than that of the p4. The distal lobe of m1 is slightly broader than the mesial lobe. In the
413 slightly worn specimen (V 23140), a small enamel islet is identified as the postmesofossettid. A
414 transverse curved groove is the entoflexid. In the heavily worn specimen (V 23139), the
415 hypoconid, the entoconid, the hypolophids, and the posterolophid completely merge. The tooth
416 has three roots, including two slim mesial roots and one broad distal root (Figs. 7A2, B2, C2).

417 Only one specimen (V 23139) preserves m2 and m3. The m2 (Fig. 6A2) is very similar to
418 the m1. A shallow oval fossa in the middle of mesial lobe can be identified as the metafossettid.
419 The mesoflexid between the mesial lobe and distal lobe is lingually closed. In the distal lobe, the
420 transverse fossa is identified as the entofossettid. As in m1, m2 also has two slim mesial roots
421 and one broad distal (Figs. 7A2, C2).

422 The m3 (Fig. 6A2) is very similar to m1 and m2, but is slightly longer and narrower. Its
423 mesial lobe has two fossae. The large buccal one is identified as the metafossettid. The tiny
424 lingual fossa is identified as the premesofossettid. As in m2, the mesoflexid is lingually closed.
425 In the distal lobe, the large and oblique entoflexid is preserved. The tooth has three roots as those
426 of m1 and m2 (Fig. 7C2).

427

428 **Phylogenetic Analysis**

429 The parsimony search of our phylogenetic analysis provided 6 most parsimonious trees. Each has
430 a best score of 543 steps (CI = 0.3554 and RI = 0.6625). The majority-rule consensus shows that

431 most clades have 100% consensus (Fig. 8). The inner group (castorids) is a monophyletic group
432 with robust absolute and relative Bremer Supports. Character-state optimization using the
433 Accelerated transformation (ACCTRAN) criterion shows that the inner group is supported by 21
434 dental and 2 cranial synapomorphies (Table 3). Six species of *Propalaeocastor* (*P. schokensis*, *P.*
435 *butselensis*, *P. kazachstanicus*, *P. kumbulakensis*, *P. irtyshensis*, *P. shevyrevae*), four species
436 previously referred to *Agnotocastor* (*P. galushai*, *P. readingi*, *P. coloradensis*, *P. aubekerovi*), *P.*
437 *primus* and “*Steneofiber* aff. *S. dehmi*” form a monophyletic group. The absolute and relative
438 Bremer Supports show that the monophyly of this group is quite robust. This result supports our
439 systematic revision of *Propalaeocastor*. Character-state optimization shows that
440 *Propalaeocastor* clade is supported by 7 dental and 2 cranial synapomorphies (Table 4).

441

442 DISCUSSION

443

444 **Comparisons.** Many researchers suggested that *Propalaeocastor* is similar to *Steneofiber*
445 (Lytschev, 1970; Lytschev & Shevyreva, 1994; Lopatin, 2003, 2004; Bendukidze et al., 2009).
446 Wu et al. (2004) also listed seven characters shared by the two genera. *Steneofiber* was
447 established by Geoffroy Saint-Hilaire (1833) for the beavers fossils discovered at Langy (Allier)
448 in the basin of Saint-Gérard-le-Puy, France. Its type species is *S. castorinus* identified by Pomel
449 (1846) (see Stirton, 1935). The new Jeminay specimens reported here show that *Propalaeocastor*
450 differs from *Steneofiber* by presenting a P3, and in having a larger P4 and p4 relative to the
451 molars, a mesiodistally more elongated P4 and p4, relatively wider molars, and more
452 complicated ridge-fossa pattern. In *Propalaeocastor*, the metalophs on the upper teeth and the
453 hypolophids on the lower teeth are divided to two or three branches. The upper teeth and the
454 lower teeth usually have a premesofossette and postmesofossette, and a premesofossettid and
455 postmesofossettid respectively. The mesoflexus and mesoflexid are more transversely orientated.
456 In the narrower flexures and fossae of *Propalaeocastor*, many crenulated enamel folds usually
457 develop from the adjacent lophs or ridges. In a sharp contrast, *Steneofiber* has a relatively much
458 simpler and less crenulated ridge-fossa pattern.

459 “*Steneofiber* aff. *dehmi*” from the early Oligocene Saint-Martin-de-Castillon of France
460 (Huguency, 1975) was treated as a member of *Propalaeocastor* by Wu et al. (2004). Here we
461 follow their assignment. As in other *Propalaeocastor* specimens, “*Steneofiber* aff. *dehmi*” has
462 premesofossettes and postmesofossettes on the upper cheek teeth, and has premesofossettids and
463 postmesofossettids on the lower cheek teeth. Compared to *P. irtyshensis*, “*Steneofiber* aff. *dehmi*”
464 is larger. The mesoflexus on the upper cheek teeth are more distally extended due to lacking a
465 metalophule I. The lower cheek teeth are more slender and have the metastylid crests.

466 *Propalaeocastor* shares many similarities with the North American late Eocene to early
467 Oligocene *Agnotocastor*, which is widely regarded as the oldest castorid genus (Korth, 1994; Xu,
468 1995, 1996; Flynn & Jacobs, 2008). As in *Propalaeocastor*, a single-rooted P3 is also present in
469 *Agnotocastor*. Previously six species were included in this genus. Four of them, namely the type
470 species *A. praetereadens*, “*A.*” *coloradensis*, “*A.*” *galushai* and “*A.*” *readingi*, are from North
471 America. Two species, “*A.*” *aubekerovi* and *A. devius*, are from Kazakhstan of Asia (Stirton,

472 1935; Wilson, 1949b; Emry, 1972; Lytshev, 1978; Korth, 1988; Lytshev & Shevyreva, 1994).
473 Based on the dental morphology and our phylogenetic analysis, we transfer four species
474 (*coloradensis*, *galushai*, *readingi* and *aubekerovi*) to *Propalaeocastor*, and reserve only *A.*
475 *praetereadens* and *A. devius*, in *Agnotocastor*. *A. praetereadens* is from the White River
476 Formation of South Dakota, USA, and is represented by a skull (AMNH 1428). As in *P.*
477 *irtyshensis*, P3 is also present in *A. praetereadens* and *A. devius*. *A. praetereadens* differs from *P.*
478 *irtyshensis* in having simpler dental morphology that lacks premesofossettes and
479 postmesofossettes on upper cheek teeth. *A. devius* from Mayliaby of Zaissan Basin (Lytshev &
480 Shevyreva, 1994) also has a distinctly simpler dental morphology. It differs from *P. irtysheensis*
481 in having smaller tooth size, shallower mandibular depth beneath the p4, and more caudodorsally
482 extending angular process of the mandible.

483 *P. coloradensis*, *P. galushai*, and *P. aubekerovi* include only lower jaw fragments and lower
484 teeth. They all have distinct postmesofossettids on their lower cheek teeth. This is the diagnostic
485 feature of *Propalaeocastor*. Furthermore, the position of the mental foramen of these three
486 species is also beneath the anterior root of p4. *P. readingi* from the Orella Member of Brule
487 Formation of Dawes County in Nebraska was named based on a mandibular fragment preserving
488 p4-m2 (CSC 80-1; Korth, 1988). Later, Korth (1996a) described additional specimens of this
489 species and emended its diagnostic features. Its dental morphology displays a complicated
490 pattern, such as presenting the premesofossette and postmesofossette on the upper cheek teeth,
491 and the postmesofossettids on the lower cheek teeth. These features are typically seen in
492 *Propalaeocastor*.

493 *P. coloradensis* from the Brule Formation of Logan County in Colorado (Wilson, 1949b)
494 differs from *P. irtysheensis* in having greater tooth size, lower tooth crown, deeper mandibular
495 depth beneath p4 (Table 2), and in presenting a digastric eminence and distinct metastylid crests
496 on the lower cheek teeth. *P. galushai* from the South Fork of Lone Tree Gulch in Wyoming
497 (Emry, 1972) is similar to *P. irtysheensis* in size (Table 2). *P. galushai* has a stronger digastric
498 eminence and lower tooth crowns. Its p4 metaconid mass and entoconid mass show weaker
499 connections to the protoconid and the hypoconid respectively than in *P. irtysheensis*. *P. readingi*
500 is slightly larger than *P. irtysheensis* (Table 2). *P. irtysheensis* differs from *P. readingi* in having
501 more transversely expanded m1 and m2. Given the very wide geographic separation, the minor
502 difference between *P. readingi* and *P. irtysheensis* is remarkable. Compared to *P. aubekerovi*
503 from Tort-Molla, Ulutau, Dzhezkazgan Province in Kazakhstan (Lytshev, 1978), *P. irtysheensis* is
504 different by lacking the digastric eminence and presenting much thicker mandibular depth
505 beneath p4 (Table 2).

506 *Propalaeocastor primus* from the Brule Formation of Fitterer Ranch in North Dakota, USA
507 was raised as the type species of *Oligotheriomys* (Korth, 1998). Here we take *Oligotheriomys* as
508 the junior synonym of *Propalaeocastor*. *P. primus* has only one right maxilla preserving M1-2
509 (FAM 64016). The preserved alveolus indicates that the P3 is present. The molar morphology of
510 this species is complicated. As in other species of *Propalaeocastor* but different from other basal
511 castorids, the paracone and metacone and the ridges associated with these two cusps form
512 complex paracone mass and metacone mass. The premesofossette and postmesofossette are

513 clearly present. *P. primus* differs from *P. irtyshensis* by its distinctly larger size, higher crown
514 and much shallower hypoflexus and mesoflexus.

515 The type species *Propaleocastor kazachstanicus* was discovered from Kyzylkak,
516 Dzhezkazgan and Kazakhstan (Borissoglebskaya, 1967). Compared to *P. kazachstanicus*, *P.*
517 *irtyshensis* has a relatively deeper mandibular depth beneath the p4 (Table 2). Caudoventral to
518 the mandibular symphysis, a small digastric eminence is present in *P. kazachstanicus*, but not in
519 *P. irtyshensis*. The preserved part of the angular process in *P. irtyshensis* shows that the angular
520 process probably is more caudoventrally directing than that in *P. kazachstanicus*. *P. irtyshensis*
521 has more transverse mesoflexids on the lower cheek teeth than those in *P. kazachstanicus*.
522 Lytshev & Shevyreva (1994) referred nine isolated cheek teeth discovered from Maylibay of
523 Zaissan Basin to *P. kazachstanicus* (fig 2 in Lytshev & Shevyreva, 1994). These teeth differ
524 from *P. irtyshensis* by having narrower crowns, and by having more distally extended
525 mesoflexus on M1-2 and only one premesofossettid on p4.

526 Compared to *P. butselensis* from the Hoogbustsel-Hoeleden in Belgium (Misonne, 1957), *P.*
527 *irtyshensis* has a more complicated dental structure. The premesofossette, metaflexus and
528 premesofossettid in *P. irtyshensis* are usually divided by extra septa or spurs. The mesoflexus in
529 *P. irtyshensis* is more distally extending, while in *P. butselensis* it is nearly transverse.
530 “*Steneofiber* cf. *S. butselensis*” from the Buran Svita of Podorozhnik, locality K15, south of Lake
531 Zaissan (Emry et al., 1998) was also regarded as a member of *Propalaeocastor* by Wu et al.
532 (2004). These specimens are very similar to *P. irtyshensis*. They have a slightly smaller tooth
533 size and relatively narrower m1-2 than *P. irtyshensis*.

534 *P. kumbulakensis* was discovered from the Kumbulak cliffs, the loc. Altyn Schokysu, the
535 loc. Akotau, the loc. Akespe, and the loc. Sayaken near the Aral Sea (Lytshev, 1970; Lopatin,
536 2003; Lopatin, 2004; Bendukidze et al., 2009). It is much larger and more robust than *P.*
537 *irtyshensis*. The upper teeth of *P. kumbulakensis* have premesofossettes, postmesofossettes and
538 double metalophs. The lower teeth have the postmesofossettids and double hypolophids. These
539 features are similar to those in *P. irtyshensis*. Similar to *P. irtyshensis*, *P. kumbulakensis* does not
540 have a digastric eminence, and its angular process extends caudoventrally. The p4 of *P.*
541 *kumbulakensis* has a single premesofossettid, and a large groove merged by mesoflexid and
542 metaflexid. The hypoflexid in *P. kumbulakensis* is very deep and extends lingually on the p4-m1.
543 The postmesofossettid is absent on the p4, but is present on the m1. The tooth crown of the m1 in
544 *P. irtyshensis* is mesial-distally more compressed and buccal-lingually wider than in *P.*
545 *kumbulakensis*.

546 *P. schokensis* from the Altyn Schokysu of Kazakhstan (Bendukidze, 1993) is larger than *P.*
547 *irtyshensis* (Table 2). It differs from *P. irtyshensis* in having much more massive paracone and
548 metacone masses on upper cheek teeth but with simpler metaconid mass on the p4 (see
549 Bendukidze et al., 2009).

550 Compared to *P. irtyshensis*, *P. shevyrevae* from Talagay in the Zaissan Basin (Lytshev &
551 Shevyreva, 1994) has relatively lower tooth crowns, less folded inner surfaces of enamel islets,
552 smaller p4 with a more rounded protoconid and a less projected hypoconid. The lower cheek
553 teeth of *P. shevyrevae* have premetafossettids and single premesofossettids. The m3 is more

554 elongated and has two metafossettids. *Propalaeocastor* aff. *P. shevyreva* from the Podorozhnik
555 and the Novei Podorozhnik in the Zaissan Basin (Lytshev & Shevyreva, 1994) is similar to *P.*
556 *irtyshensis* in overall morphology. The P4 of *Propalaeocastor* aff. *P. shevyreva* is slightly
557 larger and more slender than that of *P. irtyshensis*. It differs from *P. irtyshensis* in having more
558 tortuous enamel folds that protrude into the fossae on upper teeth, and in having one
559 premesofossettid on p4.

560 *P. zaissanensis* from the Talagay in the Zaissan Basin (Lytshev & Shevyreva, 1994) is
561 very close to *P. irtyshensis* in both tooth size and morphology. *P. zaissanensis* differs from *P.*
562 *irtyshensis* in having a relatively narrow p4, and a hypoflexus transversely confluent with the
563 mesoflexus on M3.

564 Some other basal castorid genera including *Miotheriomys*, *Microtheriomys*, *Minocastor* and
565 *Neatocastor* were regarded as close relatives of *Propalaeocastor* (Korth, 1996b, 2004; Korth &
566 Samuels, 2015; Mörs et al., 2016). All these genera include their type species only. Korth (1996b)
567 dumped “*Steneofiber*” *hesperus* Douglass, 1901, “*S.*” *complexus* Douglass, 1901 and “*S.*”
568 *montanus* Scott, 1893 into one species (“*S.*” *hesperus*) and established a new genus (*Neatocastor*)
569 for it. The type specimens of *Neatocastor hesperus* was from the Arikarean (late Oligocene) of
570 the Blacktail Deer Creek of Beaverhead County in Montana. It has a dP3 and relatively
571 complicate upper dental morphology, but with relatively simple lower teeth similar to that of
572 *Steneofiber*. *N. hesperus* differs from *Propalaeocastor* in having more convex lower incisor
573 enamel surface and weakly developed endolophs on the upper cheek teeth, and in lacking the
574 postmesofossettes on the upper cheek teeth and the premesofossettids and the postmesofossettids
575 on the lower cheek teeth. *Miotheriomys stenodon* is from the Runningwater Formation (Early
576 Hemingfordian, Early Miocene) of western Nebraska (Korth, 2004). It differs from
577 *Propalaeocastor* in lacking the premesofossettids and the postmesofossettids on the lower cheek
578 teeth. *Microtheriomys brevirhinus* is from the John Day Formation (early Early Arikarean, late
579 Early Oligocene) in Oregon (Korth & Samuels, 2015). It is different from *Propalaeocastor* by
580 lacking the P3, lacking the premesofossettids and the postmesofossettids on the lower cheek
581 teeth, and presenting the dorsal palatine foramen entirely within the palatine bone. *Minocastor*
582 *godai* is from the lower Miocene of the Kani Basin in central Japan (Mörs et al., 2016). It is
583 distinctly larger than the all the species of *Propalaeocastor*. The enamel surface of its lower
584 incisor is more convex than that of *Propalaeocastor*. Its lower cheek teeth are more *Steneofiber*-
585 like by presenting very reduced presmesofossettids and postmesofossettids. Its upper cheek teeth
586 display a relatively complicated dental pattern as in *Propalaeocastor*, but without the
587 postmesofossette.

588 The new *Propalaeocastor irtyshensis* specimens reported here show that the dental
589 morphology of this species is similar to other early castorids, such as *Agnotocastor*, and
590 *Neatocastor* and *Microtheriomys*. On the other hand, *P. irtyshensis* also possesses some features
591 that superficially similar to the eutypomyids, such as two upper premolars and complicate cusp
592 and ridge patterns. Among castorids, it is known that two upper premolars are present in
593 *Agnotocastor devius* (Lytshev & Shevyreva, 1994), and some North American early castorids,
594 such as *Agnotocastor*, *Neatocastor* and “*Oligotheriomys*” (which is sunk into *Propalaeocastor*

595 here) of North America (Stirton, 1935; Korth, 1996b, 1998).

596 Extant and fossil castorid skulls clearly exhibit the sciuriform skull pattern, while the
597 sister-group of castorids, the eutypomyids, show the protrogomorphous morphology (Wood,
598 1965). In basal castorids, it was not clear whether they have the protrogomorphous pattern or the
599 sciuriform pattern. The zygomatic process of maxilla of *P. irtyshensis* displays a
600 conspicuous mesiodorsally-distoventrally oblique surface. In protrogomorph skulls the
601 zygomatic root ventral to the infraorbital foramen has an oval roughened scar for the attachment
602 of the anterior part of the deep masseter and the superficial masseter. No such a scar is present in
603 *P. irtyshensis*. A sloping zygomatic process of maxilla without the oval scar indicates that a
604 rudimentary sciuriform-like zygomatic plate probably is present (Figs. 5A1-2). Medial to this
605 rudimentary zygomatic plate and dorsal to the zygomatic root of the maxilla, it presents a smooth
606 and round surface. This surface indicates that the infraorbital foramen is large and round, and the
607 infraorbital canal is short. The rudimentarily developed zygomatic plate coupled with a large
608 infraorbital foramen and canal probably is the plesiomorphic feature for all castorids. In extant
609 beavers, the infraorbital foramen is small, the infraorbital canal is long, and the sciuriform
610 zygomatic plate forms a deep fossa locating lateral to the infraorbital canal (Cox & Baverstock,
611 2016). More derived fossil beavers, such as *Monosaulax*, *Eucastor*, *Procastoroides* etc., all have
612 the sciuriform-like zygomatic plate with a deep fossa. In myomorphous rodents, the zygomatic
613 plate is present and the infraorbital foramen is large. Different from the protrogomorphous
614 rodents, the large infraorbital foramen in the myomorphous rodents is mediolaterally compressed.
615 The large infraorbital foramen in *P. irtyshensis* does not show any sign of compression.

616 Xu (1996) argued that *Propalaeocastor kumbulakensis* should be assigned to the
617 eutypomyid genus *Eutypomys* because the lower jaw of *P. kumbulakensis* does not have a
618 digastric eminence, and its angular process extends caudoventrally. We re-examined the
619 mandibular specimens of *Propalaeocastor* and found that the digastric eminence is variably
620 present in different species. In *P. kumbulakensis*, *P. irtyshensis*, *P. readingi* and *P. devius*, the
621 digastric eminence is absent, while in some other species, such as *P. coloradensis*, *P. galushai*, *P.*
622 *aubekerovi* and *P. kazachstanicus* it is well-developed. In *P. irtyshensis*, the articular facet of the
623 mandibular symphysis has a large expansion beneath the genial fossa. The presence of this
624 enlargement strengthens the mandibular symphysis. In all the castorids with genial region
625 preserved, the articular facet of the mandibular symphysis all has this ventral expansion. When
626 the digastric eminence is present, the articular facet always extends onto it. The so-called
627 digastric eminence probably is a part of articular expansion related to the strengthening of the
628 mandibular symphysis, not just for providing the arising places for the digastric muscles. In that
629 sense, the expansion of the articular facet of the mandibular symphysis is associated with the
630 appearance of digastric eminence, therefore should be regarded as a feature shared by all
631 castorids.

632 The angular process of mandible is also variably present in different species of
633 *Propalaeocastor* and other basal castorids. In some species preserving that part, such as *P.*
634 *kumbulakensis*, *P. irtyshensis*, *P. aubekerovi*, and *P. galushai*, the angular process of the
635 mandible extends caudoventrally, while in *P. kazachstanicus*, the angular process shows a

636 tendency of caudodorsal extension (Fig. 9). It is likely that the direction of the angular process is
637 related to the development of the medial pterygoid muscle, and probably also superficial
638 masseter.

639 Korth (1994), Rybczynski (2007), and Flynn & Jacobs (2008) enumerated many features of
640 Castoridae that are different from Eutypomidae, such as the relatively high rostrum cross-
641 sectional shape, wider nasals, the small and mediolaterally compressed infraorbital foramen, the
642 long infraorbital canals, the distinctive chin process (symphyseal flange, or mandibular
643 eminence), and the base of lower incisor terminating in a lateral bulbous expansion etc.. Xu
644 (1996) once defined the castorids as “the rodents that have sciuriformous masseter
645 arrangement on the skull and a derived mandible here termed the beaver-pattern mandible.” His
646 “beaver-pattern mandible” is referred to a mandible presenting “digastric eminence” and “the
647 angle extending up posteriorly”. Eutypomyidae is characterized by presenting a narrow
648 zygomatic plate, a large and round infraorbital foramen, a short infraorbital canal, two upper
649 premolars, and a lower jaw lacking the digastric eminence and having a caudoventrally
650 extending angular process (e.g., Wahlert, 1977; Korth, 1994). Obviously all these features are
651 cranial features. Our phylogenetic analysis is based on a data matrix that includes mainly dental
652 features. It is not possible to evaluate all the differences between castorids and eutypomyids
653 mentioned above, but our analysis does discover that presence of digastric eminence and large
654 capsular process are synapomorphies of all castorids. The caudal palatine foramen situated in
655 maxillary-palatine suture is a feature generally accepted as a diagnostic character for castorids
656 (Korth, 2002). In our analysis, this feature is a synapomorphy of *Propalaeocastor*, but not the
657 synapomorphy of castorids (Tables 3 & 4). On the other hand, our phylogenetic analysis suggests
658 that dental features are also important for distinguishing the castorids and eutypomyids. Twenty-
659 one dental features are synapomorphies of the castorid clade (Table 4).

660

661 **Phylogeny and Applications.** It is generally believed that *Agnotocastor* and *Propalaeocastor*
662 are close to each other (Korth, 2002, 2004; Korth & Samuels, 2015; Mörs et al., 2016). Our
663 phylogenetic analysis suggests that some species of “*Agnotocastor*”, namely of *P. galushai*, *P.*
664 *readingi*, *P. coloradensis* and *P. aubekerovi*, should be reassigned to *Propalaeocastor*.
665 “*Steneofiber* aff. *S. dehmi*” from the Saint-Martin-de-Castillon in France (Huguenev, 1975) is
666 morphologically more similar to *Propalaeocastor* than to *Steneofiber*. Wu et al. (2004) assigned
667 this species to *Propalaeocastor* but did not give a new name to it. The result of our analysis
668 indicates that “*Steneofiber* aff. *S. dehmi*” and three North American species (*P. galushai*, *P.*
669 *readingi* and *P. primus*) form a monophyletic group. This result is consistent with our
670 comparisons and that of Wu et al. (2004). *P. primus* was described as a new species based on the
671 comparison with *Anchitheriomys* (Korth, 1998). Our result suggests that *P. primus* is the sister
672 group of *P. readingi*, deeply nesting in the monophyletic clade of *Propalaeocastor*. To keep the
673 monophyly of *Propalaeocastor*, we should sink *Oligotheriomys* to *Propalaeocastor*. The type
674 species of *Agnotocastor* (*A. praetereadens*) and *A. devius* (Stirton, 1935; Lytschev & Shevyreva,
675 1994) form a monophyletic group with high Bremer Support. They are not the sister group of
676 *Propalaeocastor*, but stem taxa that eventually lead to the crown castoroid group.

677 *Steneofiber* was suggested to be very close to *Propalaeocastor*. Lytschev & Shevyreva
678 (1994), and Lopatin (2003, 2004) even suggest that *Propalaeocastor* is a junior synonym of
679 *Steneofiber*. Some species, such as *P. butselensis*, *P. kumbulakensis* and *P. schokensis*, were
680 referred to *Steneofiber* (Hugueney, 1975; Lytschev & Shevyreva, 1994; Lopatin, 2003, 2004),
681 while Wu et al. (2004) and Bendukidze et al. (2009) referred them to *Propalaeocastor*. Our
682 phylogenetic analysis indicates that *Steneofiber* is a polyphyletic group. The type species,
683 *Steneofiber castorinus*, is the sister group of *Chalicomys* + *Castor*, suggesting that *Steneofiber* is
684 far more derived than the basal castorid *Propalaeocastor*.

685 Korth (2001) believed that *Propalaeocastor* is close to *Oligotheriomys* and *Anchitheriomys*,
686 and assigned these genera to the Tribe Anchitheriomyini of the Subfamily Agnotocastorinae.
687 Later, Korth (2004) named *Miotheriomys* and elevated the Tribe Anchitheriomyini into the
688 Subfamily Anchitheriomyinae to include *Propalaeocastor*, *Oligotheriomys*, *Anchitheriomys* and
689 *Miotheriomys*. Korth & Samuels (2015) named *Microtheriomys* and also include it into the
690 Subfamily Anchitheriomyinae. Mörs et al. (2016) named *Minocastor* and raised a tribe (Tribe
691 Minocastorini) of the Subfamily Anchitheriomyinae to include *Minocastor*, *Microtheriomys*,
692 *Miotheriomys*, *Oligotheriomys* and *Propalaeocastor*. Our phylogenetic analysis indicates that
693 *Oligotheriomys* is nested in the species of *Propalaeocastor*, and we synonymize *Oligotheriomys*
694 to *Propalaeocastor* to reflect this result. In our phylogenetic analysis, we discovered that
695 *Anchitheriomys*, *Minocastor* and *Miotheriomys* are close to each other, but form a paraphyletic
696 group. *Microtheriomys* takes a more basal position than those three genera.

697 The late Eocene *Propalaeocastor galushai* is the oldest-known castorid. It possesses many
698 plesiomorphic features, such as the persistence of P3, the angular process of the mandible
699 extending caudoventrally, and the complicate dental pattern. These features are present in most
700 of the species of *Propalaeocastor*, and they are also present in the eutypomyids, which are
701 widely considered as the sister group of castorids. Therefore, these features are likely
702 plesiomorphic for all castorids. However, our phylogenetic analysis shows that *P. galushai* is not
703 the most basal castorid, not even the most basal *Propalaeocastor* (Fig. 8). This result would
704 suggest that the diversification of *Propalaeocastor* is before the late Eocene.

705 It was suggested that castorids originated in North America, and probably dispersed into
706 Asia during the Early Oligocene (Lytschev, 1978; Lytschev & Shevyreva, 1994; Xu, 1995; Korth,
707 2002; Rybczynski, 2007). This hypothesis is supported by our phylogenetic analysis. The place
708 of origin of *Propalaeocastor* is uncertain. Based on the result of our phylogenetic analysis, it is
709 equally parsimonious to predict an Asian origin, a European origin or a North American origin of
710 castorids (Fig. 8). A castorid earlier than *P. galushai* and more primitive than *P. irtyschensis* and
711 *P. butselensis* is yet to be discovered. The rapid radiation of castorids in the early Oligocene
712 probably is propelled by the global climate changes during the Eocene-Oligocene transition
713 (EOT). Dramatic sea level drop during the EOT probably produced multiple passages enabling
714 the dispersal of *Propalaeocastor*-like basal castorids across the northern continents. These basal
715 castorids then quickly became diversified and adaptive to new ecological niches.

716

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724

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Figure 1(on next page)

Jeminay and Burqin *Propalaeocastor irtyshensis* fossil localities in the Irtys River drainage area in northwestern Xinjiang, China (modified from Stidham & Ni, 2014).

A: Map showing the location of the *P. irtyshensis* localities in the Irtys River region within China adjacent to several other countries; **B:** Detailed map showing the border region between Xinjiang and Kazakhstan and the localities of *P. irtyshensis*; **C:** Panoramic view of the fossiliferous profile of the Irtys River Formation that produced the additional material of *P. irtyshensis*.

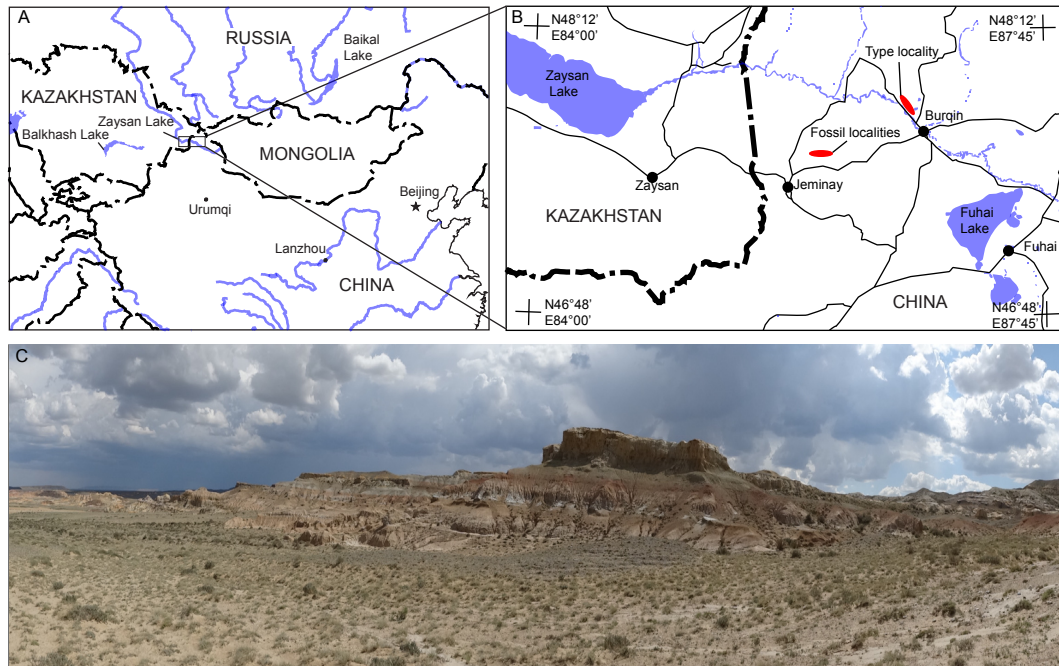


Figure 2(on next page)

The upper dental structure of the Castoridae after the example of the moderately worn fourth premolars of *Propalaeocastor irtyschensis*.

A: *Propalaeocastor*; **B:** *Steneofiber*; **C:** *Castor*; **D:** *Dipoides*. From left to right: lingual view, occlusal view, buccal view. Enamel=white, dentine=dark grey, cement=light grey, -fossette=-flexus=-stria. Modified from Stirton, 1935; Hugueney, 1975, 1999; Lopatin, 2003 and Wu et al., 2004.

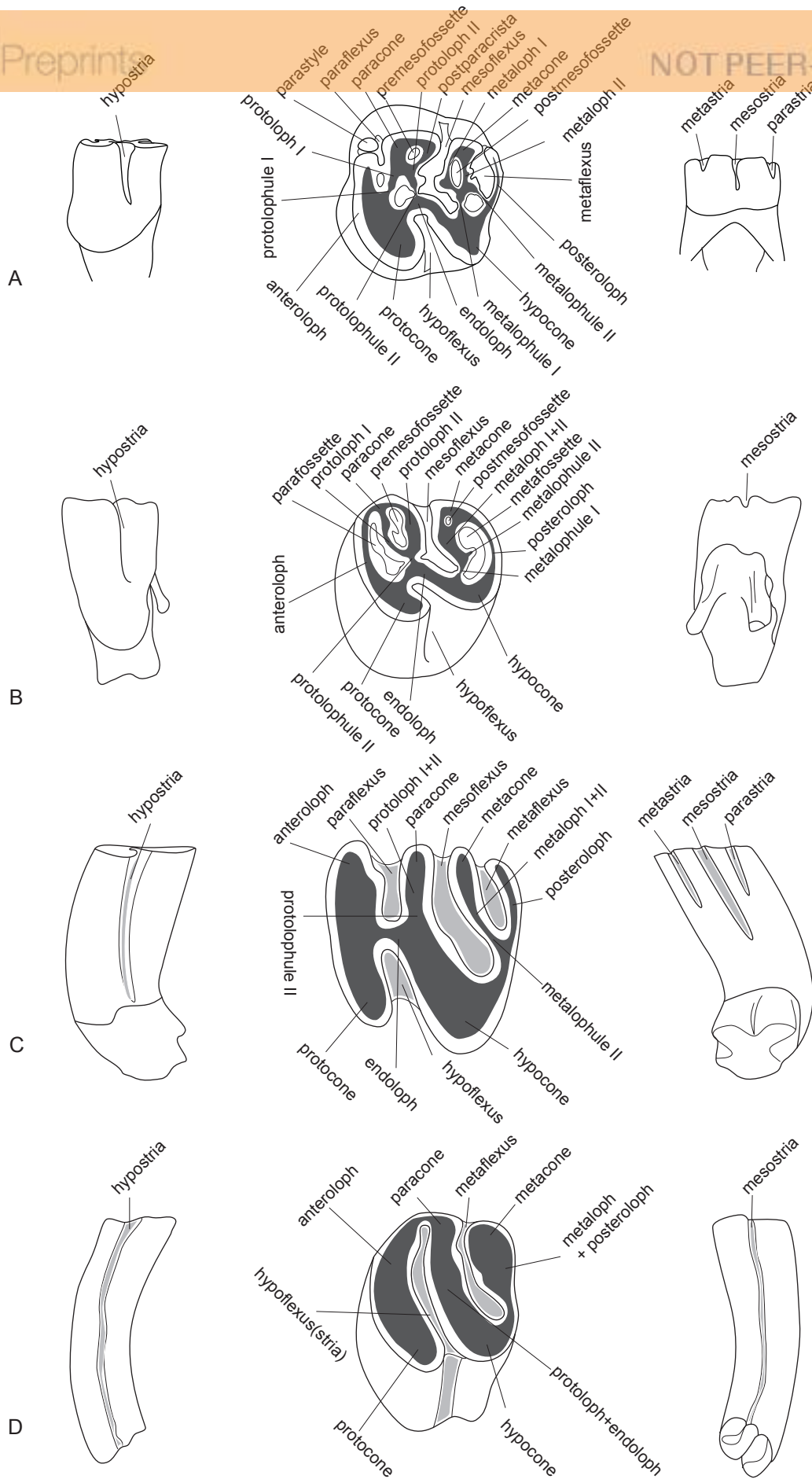


Figure 3(on next page)

The lower dental structure of the Castoridae after the example of the moderately worn fourth premolars of *Propalaeocastor irtyschensis*.

A: *Propalaeocastor*; **B:** *Steneofiber*; **C:** *Castor*; **D:** *Dipoides*. From left to right: lingual view, occlusal view, buccal view. Enamel=white, dentine=dark grey, cement=light grey, -fossettid=-flexid=-striid. Modified from Stirton, 1935; Huguene, 1975, 1999; Lopatin, 2003 and Wu et al., 2004.

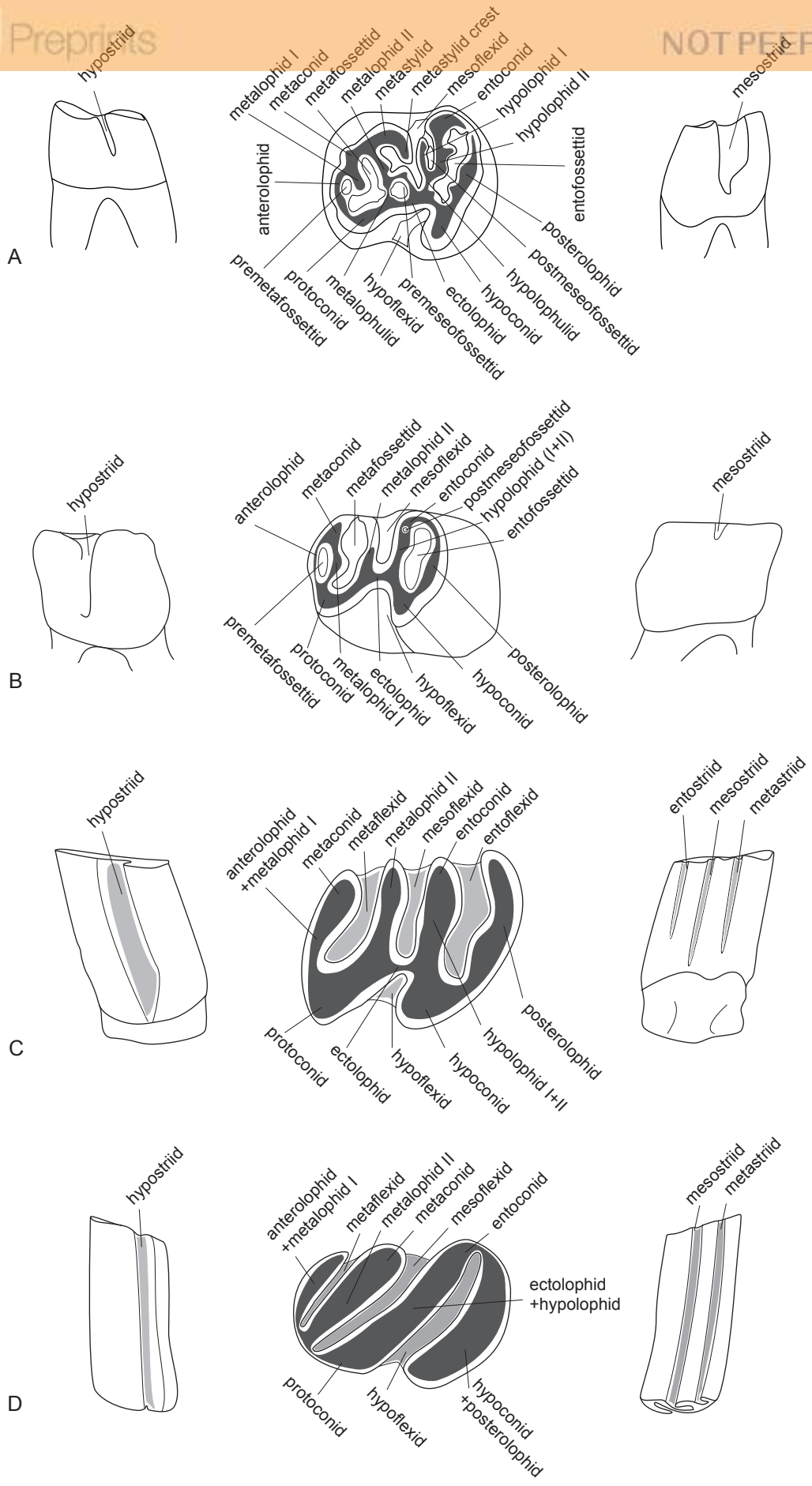


Figure 4(on next page)

Maxilla and isolated upper cheek teeth referred to *Propalaeocastor irtyshensis* from Jeminay area, northwestern Xinjiang, China.

Yellow shadow showing the divergence of palatine nerve. **A1-3**: broken maxilla with right P4-M1 (IVPP V 23138.1); **B1-3**: left P4 (IVPP V 23138.2); **C1-3**: left M1 (IVPP V 23138.3). **A1, B1, C1**: lingual; **A2, B2, C2**: buccal; **A3, B3, C3**: occlusal. All in same scale.

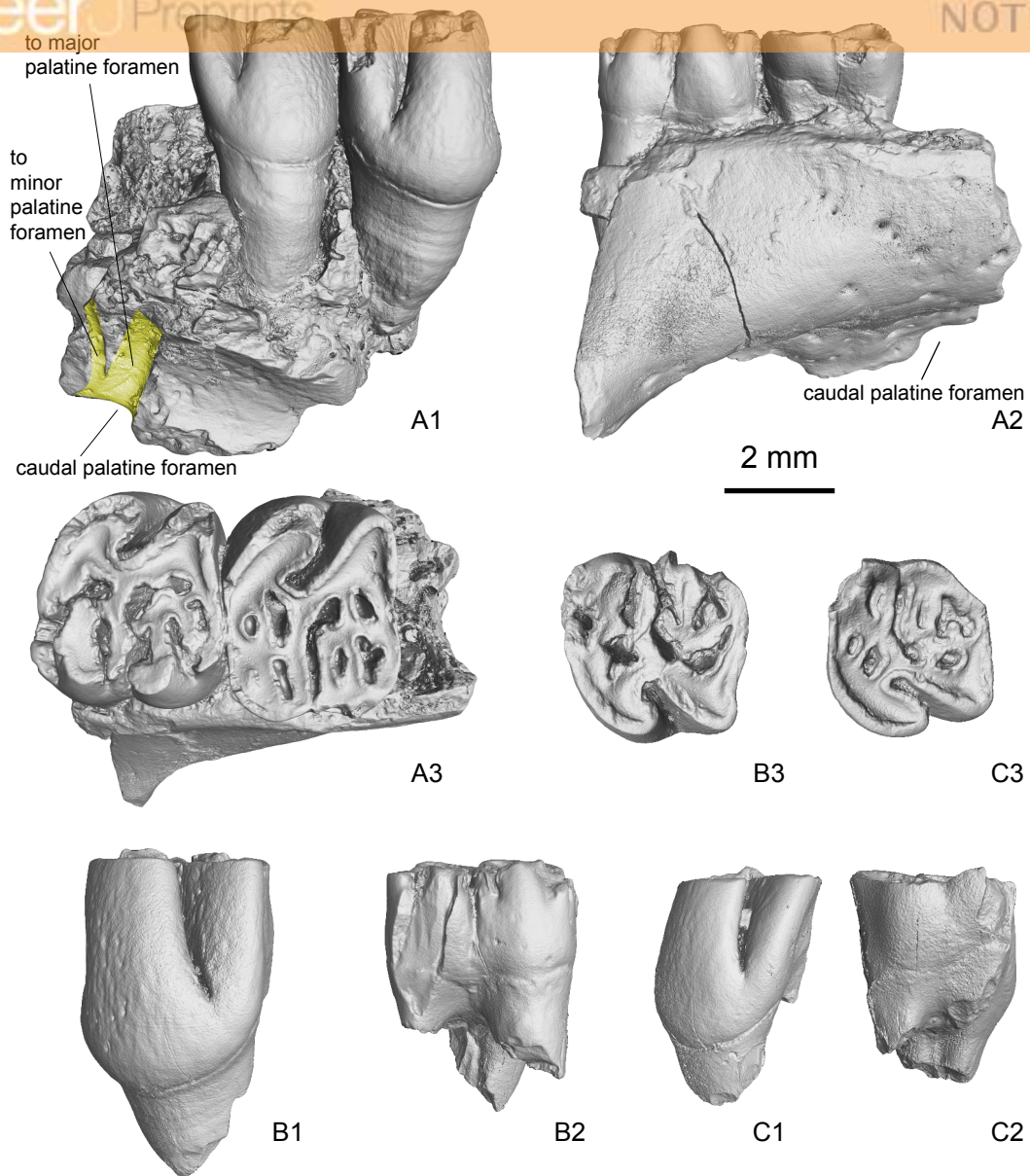


Figure 5(on next page)

3D virtual reconstruction of the maxillae of *Propalaeocastor irtyshensis* by the X-ray computed tomography.

Red shadow showing a residual P3 alveolus mesial to the mesial roots of P4; dashed cycle displaying a relative large and round infraorbital foramen dorsal to the zygomatic arch root preserved in both holotype of Burqin (**A1-3**: IVPP V 13690) and additional specimen of Jeminay (**B1-2**: IVPP V 23138.1).

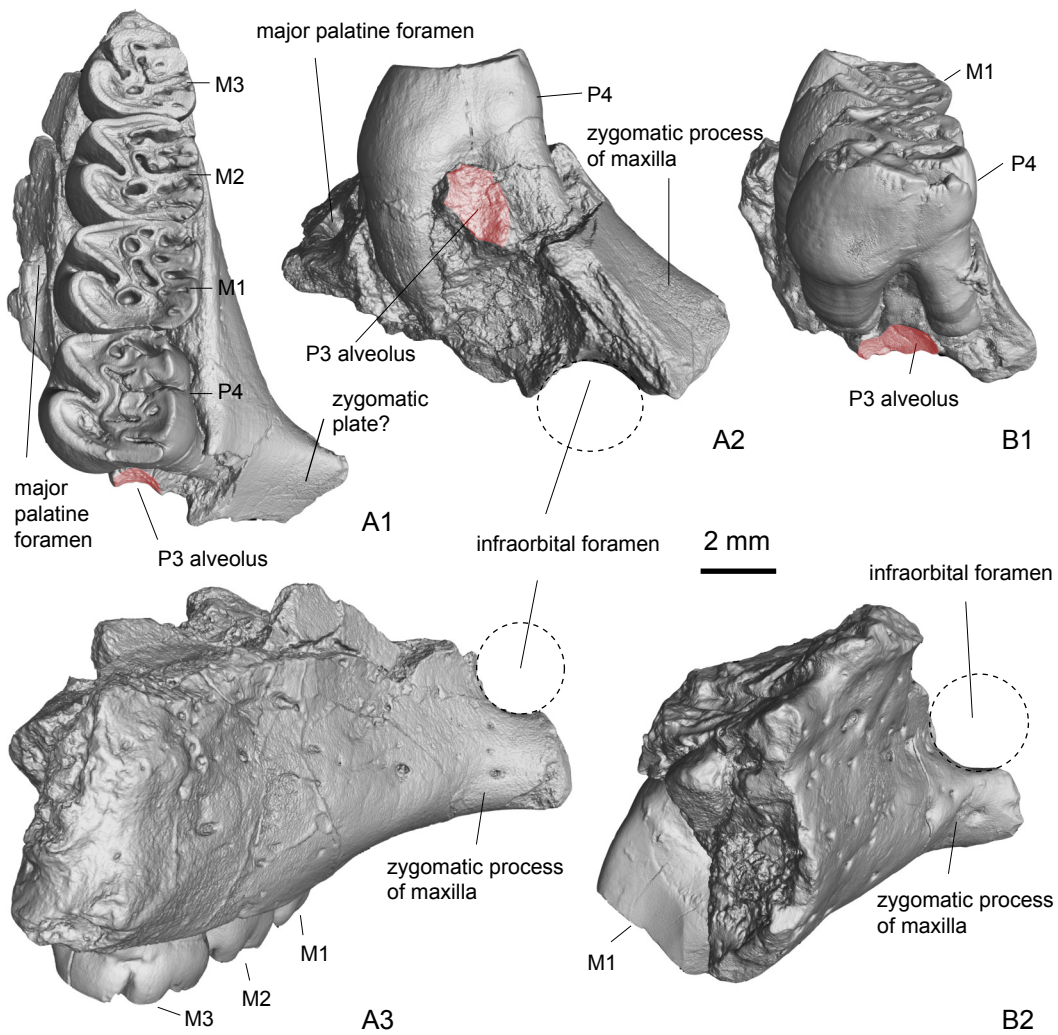


Figure 6(on next page)

Fragmentary dentaries of *Propalaeocastor irtyshensis* from Jeminay, Xinjiang.

Red shadow displaying articular facet of mandibular symphysis. **A1-3**: fragmentary right dentary with broken p4-m3 (IVPP V 23139); **B1-3**: broken right dentary with p4-m1 (IVPP V 23140); **C1-3**: broken right dentary with p4 (IVPP V 23141). **A1, B1, C1**: lingual; **A2, B2, C2**: occlusal; **A3, B3, C3**: buccal. All in same scale.

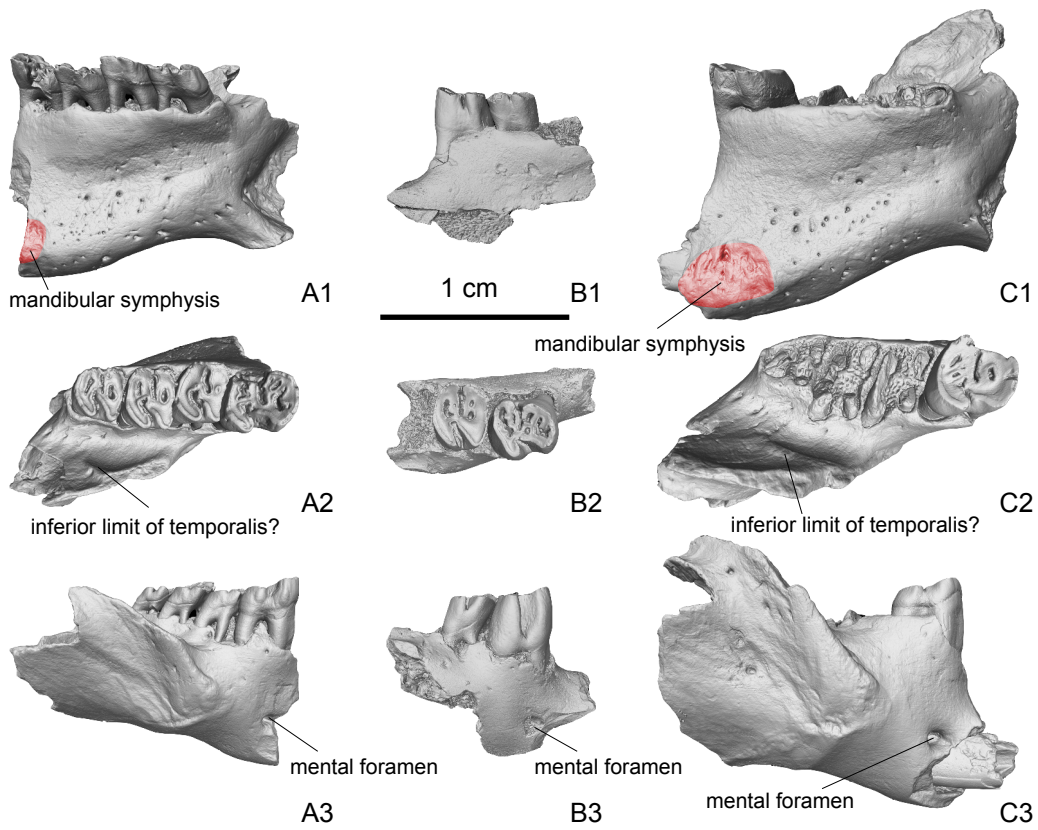


Figure 7 (on next page)

Transverse and sagittal sections of dentaries and transverse section of lower incisor of *Propalaeocastor irtyshensis* of Jeminay, Xinjiang.

Showing convex enamel surface of lower incisor, permanent fourth premolar and root number of lower cheek teeth (p4:m1:m2:m3=2:3:3:3). **A1-2**: fragmentary right dentary with p4 (IVPP V 23141); **B1-2**: broken right mandible with p4-m1 (IVPP V 23140); **C1-2**: fragmentary right dentary with p4-m3 (IVPP V 23139). **D**: lower incisor (IVPP V 231411). **A1**, **B1**, **C1**: sagittal section; **A2**, **B2**, **C2**, **D**: transverse section.

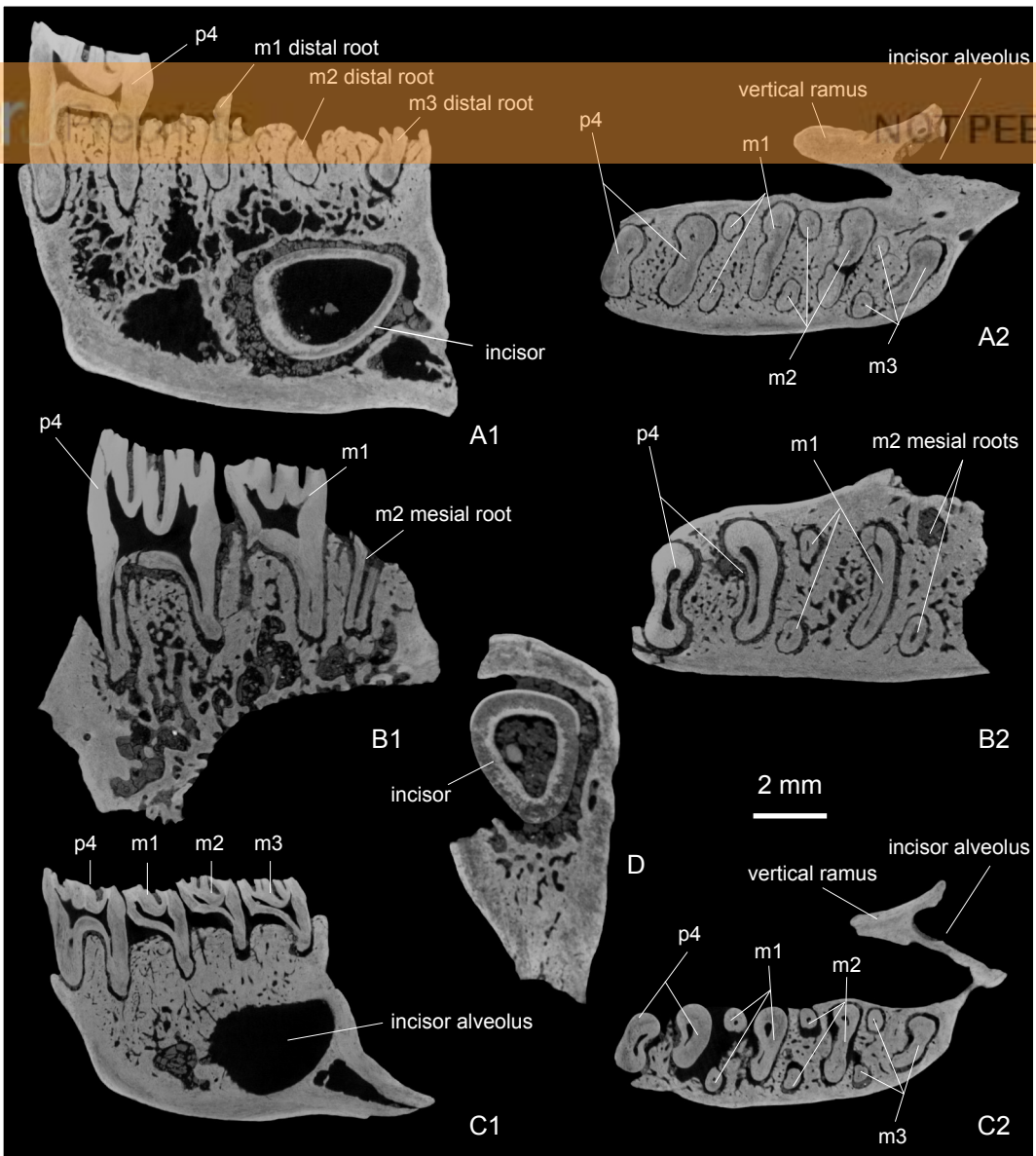


Figure 8(on next page)

Majority-rule consensus of 6 most parsimonious trees.

Parsimony analysis is based on a data matrix including 145 characters scored for 42 taxa. *Marmota monax*, *Keramidomys fahlbuschi* and *Eutypomys inexpectatus* were selected as outgroup taxa. Numbers before the slashes at the internodes are the absolute Bremer Support values; numbers after the slashes are Relative Bremer Support values; numbers after the comma are percentage of consensus. Internodes without the percentage of consensus show the topologies that are present in all the 6 most parsimonious trees. The geographic distribution of all the taxa was mapped on the majority-rule consensus tree and the ancestral states were reconstructed using the parsimony criterion in Mesquite 3.2 (Maddison & Maddison, 2017). Red clades represent Asian origin, blue clades represent European origin, and black clades represent North American origin. Clades in dashed line indicate equally-parsimonious or ambiguous Asian, European or North American origins. Scale bar equals 20 character changes.

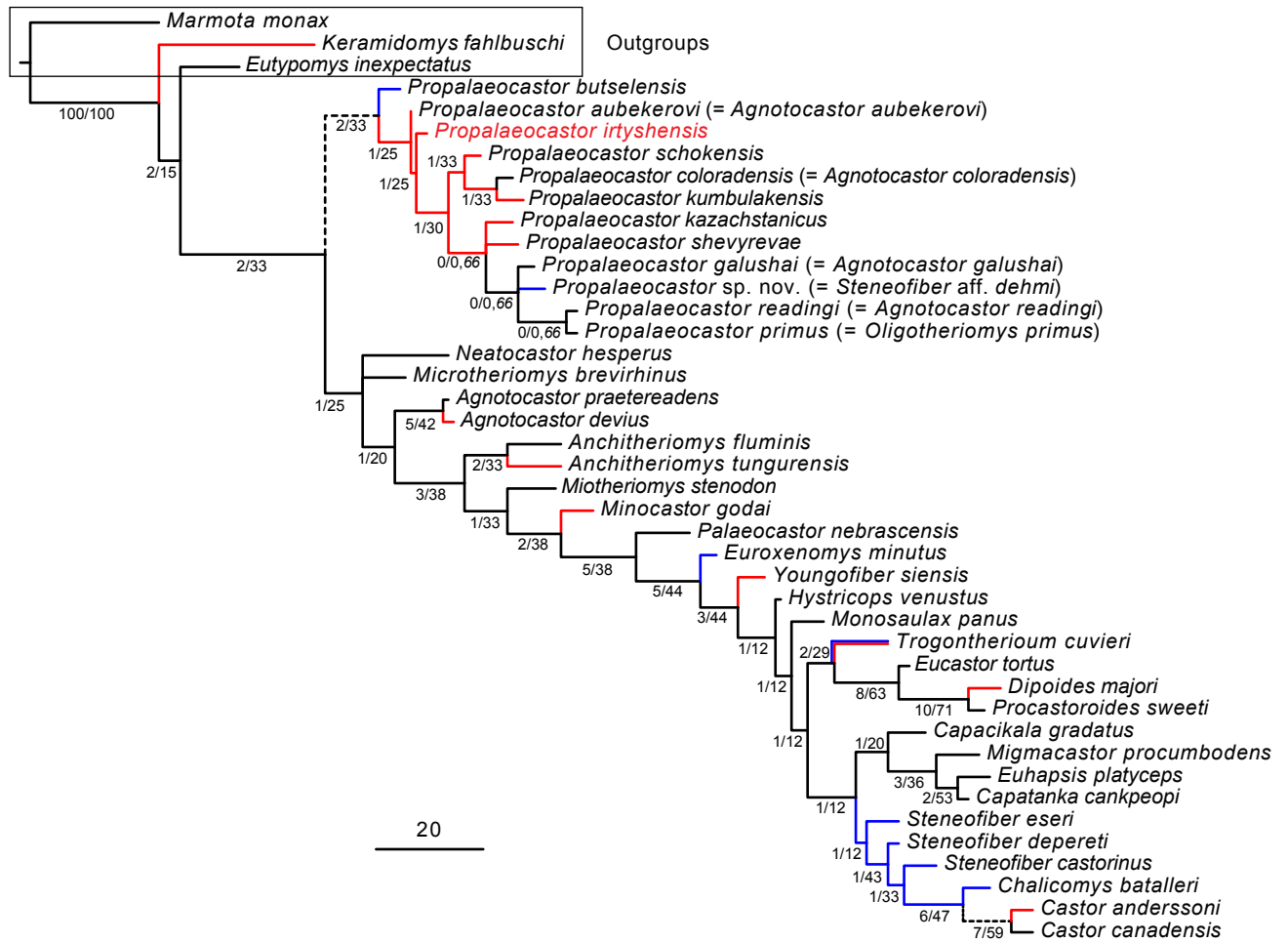


Figure 9 (on next page)

Chronologic and geographic distribution of *Propalaeocastor* and *Agnotocastor*, and comparisons of dentary and dental patterns.

Displaying the developments of digastric eminence and angular process of the mandible extending orientations of their mandibles and similarities of dental constructions. Asterisk showing the type species of *Agnotocastor* and *Propalaeocastor*. Except for the figures of *Propalaeocastor irtyschensis* (dentary, IVPP V 23141; lower dentition, IVPP V 23139, upper dentition, IVPP V 13690), the illustrations in right column are facsimiles of their original figures (Stirton, 1935; Wilson, 1949b; Borissoglebskaya, 1967; Lytschev, 1970; Emry, 1972; Hugueney, 1975; Lytschev, 1978; Korth, 1988, 1996, 1998; Lytschev & Shevyreva, 1994). Abbreviations used in left column are biochrons of North American Land-Mammal Ages (NALMA): Ch-1=Early Chadronian; Ch2-3=Middle Chadronian; Ch4=Late Chadronian; Or1-Or4=Orellan; Wh1-Wh2=Whitneyan (see Flynn & Jacobs, 2008). Dentaries and dentitions are in same scales, respectively.

	Eocene		Oligocene		Epoch	Taxa	Dentaries	Lower dentitions	Upper dentitions	
	Late	Early	Late				1 cm	5 mm	5 mm	
	Ch-1	Ch-2	Ch-3	Ch-4	Or-1	Or-2	Or-3	Or-4	Wh-1	Wh-2
Europe										
						<i>Propalaeocastor butselensis</i>				
						" <i>Steneofiber</i> aff. <i>S. dehmi</i> "				
North America						<i>Agnotocastor praetereadens*</i>				
						<i>Propalaeocastor primus</i>				
						<i>Propalaeocastor coloradensis</i>				
						<i>Propalaeocastor galushai</i>				
						<i>Propalaeocastor readingi</i>				
Asia						<i>Propalaeocastor aubekerovi</i>				
						<i>Agnotocastor devius</i>				
						<i>Propalaeocastor kazachstanicus*</i>				
						<i>Propalaeocastor ityshensis</i>				
						<i>Propalaeocastor kumbulakensis</i>				

Table 1 (on next page)

Measurements (in mm) of incisor and cheek teeth of *Propalaeocastor irtyshensis* from Jeminay, Xinjiang (L., length; W., width).

1

Inventory numbers	Tooth	Occlusal L. \times W.	Base L. \times W.	Buccal height	Lingual height	Mesostria(id) height	Hypostria(id) height	Maximum height/maximum length indices
V 23138.1	P4	3.53 \times 3.47	3.63 \times 4.67	2.09	3.49	0.49	1.37	0.96
V 23138.1	M1	3.15 \times 3.79	3.15 \times 4.81	1.56	2.43	-	0.84	0.77
V 23138.2	P4	3.46 \times 3.48	3.66 \times 4.38	1.97	3.91	0.92	1.74	1.06
V 23138.3	M1	3.04 \times 3.16	3.27 \times 3.69	1.87	3.38	-	1.64	1.03
V 23139	p4	3.79 \times 3.29	4.10 \times 3.52	2.06	1.97	-	1.14	0.50
V 23139	m1	3.12 \times 3.35	3.26 \times 3.62	1.39	1.35	0.2	0.29	0.43
V 23139	m2	3.08 \times 3.58	3.48 \times 3.71	1.53	1.42	-	0.34	0.44
V 23139	m3	3.05 \times 3.04	3.64 \times 3.22	1.44	1.45	-	0.52	0.40
V 23140	p4	3.38 \times 3.02	4.12 \times 3.54	3.83	2.60	0.82	2.27	0.93
V 23140	m1	3.05 \times 3.39	3.54 \times 3.98	2.49	2.07	0.49	0.9	0.70
V 23141	i1	3.3 \times 3.70						
V 23141	p4	4.31 \times 4.18	4.65 \times 4.41	1.83	1.54	-	0.42	0.39

2

Table 2 (on next page)

Measurements comparison of mandibles among *Propalaeocastor irtyshensis* and other taxa of *Propalaeocastor* and *Agnotocastor devius*.

Asterisk numbers are re-measured from their originally illustrations i.e. Borissoglebskaya, 1967; Lytshev, 1970; Lytshev & Shevyreva, 1994.

1

Taxa	Inventory numbers	Mandibular depth beneath p4	p4-m3, mesiodistal length	p4-m2, mesiodistal length	p4-m1, mesiodistal length	p4, mesiodistal length
<i>P. irtyshensis</i>	V 23139	11.8	12.4	9.2	6.2	3.79-4.10
<i>P. irtyshensis</i>	V 23140	-	-	-	12.0	3.38-4.12
<i>P. irtyshensis</i>	V 23141	13.2	-	-	-	4.31-4.65
<i>P. kazachstanicus</i>	No. 2259-322	9.5*	-	8.4*	6.1*	3.5*
<i>P. aubekerovi</i>	M-2041/74	9.1	11.6	9.0	6.0-7.0	3.2-3.7
<i>P. schokensis</i>	No. 15/48	-	15.7	-	-	-
<i>P. schokensis</i>	No. 15/48	-	16.4	-	-	-
<i>A. devius</i>	No. 3463-4	7.0*	10.0*	7.7*	5.1*	3.2*
<i>P. kumbulakensis</i>	M-2020/66	11.4*	19.8*	16.1*	10.9*	6.3*
<i>P. readingi</i>	CSC 80-1	11.0	-	10.6		3.35
<i>P. coloradensis</i>	UCM 19809	14.6	-	11.9		4.1
<i>P. galushai</i>	FAM 79310	10.1	11.7	9.1	-	3.4

2

3

Table 3 (on next page)

Synapomorphy list for *Propalaeocastor*.

Tree description were undertaken in PAUP* Version 4.0b10. The majority-rule consensus tree was rooted using outgroup method. Character-state optimization is done under Accelerated Transformation (ACCTRAN) model. The double arrow "=>" represents unambiguous changes, and the single arrow "-->" represents ambiguous changes.

1

Character No.	Characters	CI	State changes		
2	Upper P4 size relative to upper M1	0.200	subequal	==>	large
33	Upper P4 postmesofossette shape in metacone mass in medium wear stage	0.250	transversely long valley	==>	small enamel island-like fossa(e)
54	Upper P4 postcingulum buccal end fused with the metacone in moderate wear	0.333	present, the metaflexus is closed	-->	absent, the metaflexus is open
55	Upper M1-2 postcingulum buccal end fused with the metacone in moderate wear	0.333	present, the metaflexus is closed	-->	absent, the metaflexus is open
79	Lower p4 metastylid crest	0.250	absent	==>	present
106	Lower m1-2 metastylid crest	0.200	absent	==>	present
108	Lower m1-2 premesofossettid	0.250	absent	==>	present
122	Korth 2002. posterior palatine foramina in palatine-maxillary suture	0.250	no	-->	yes
145	Mandible, space between the lower tooth row and vertical ramus	0.143	narrow	-->	broad

2

Table 4(on next page)

Synapomorphy list for castorids.

Tree description were undertaken in PAUP* Version 4.0b10. The majority-rule consensus tree was rooted using outgroup method. Character-state optimization is done under Accelerated Transformation (ACCTRAN) model. The double arrow "=>" represents unambiguous changes, and the single arrow "-->" represents ambiguous changes.

Character No.	Characters	CI	State changes		
11	Cheek teeth crown height	0.333	brachydont	==>	mesodont
12	Upper teeth lingual higher than buccal or lower teeth buccal higher than lingual	0.500	absent	-->	present
13	Cheek teeth crown structure	0.500	bunodont-lophodont	-->	lophodont
16	Upper incisor buccal surface flatness	0.250	very convex	-->	slightly convex
17	Lower incisor buccal surface flatness	0.250	very convex	==>	slightly convex
44	Upper M1-2 preprotocrista buccal end height relative to the paracone mass or paracone	0.200	lower	-->	subequal
46	Upper M1-2 preprotocrista buccal end (parastyle) fused with the paracone mass or paracone	0.250	absent	==>	present
65	Upper cheek teeth mesocone	0.500	present	-->	absent
66	Upper cheek teeth mesoloph	0.500	present	-->	absent
67	Upper cheek teeth mesostyle	1.000	present	==>	absent
68	Lower p4 size relative to m1	0.250	smaller	-->	larger
69	Lower p4 anteroconid	1.000	present	==>	absent
70	Lower p4 anterolophid (=paracristid)	0.333	absent	-->	present
71	Lower p4 postprotocristid	0.500	absent	-->	present
82	Lower p4 mesoconid	0.500	present	-->	absent
83	Lower p4 mesolophid	0.500	present	-->	absent
93	Lower cheek teeth hypoflexid extension	0.667	shallow	-->	medium
95	Lower m1-2 anterior cingulid	0.500	present	-->	absent
103	Lower m1-2 premetafossettoid enclosing by the metalophid I and the paracristid (anterolophid)	0.167	absent	==>	present
109	Lower m1-2 mesoconid	0.500	present	-->	absent
110	Lower m1-2 mesolophid	0.500	present	-->	absent
130	Xu 1995 C2. digastric eminence on mandible	0.200	absent	-->	present
142	Mandible capsular process	0.333	weak	==>	large