

Neogene amphibians and reptiles (Caudata, Anura, Gekkota, Lacertilia, Testudines) from south of Western Siberia, Russia and Northeastern Kazakhstan (#12606)

1

Third revision

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




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



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



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3



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Organize by importance of the issues, and number your points

1. Your most important issue
2. The next most important item
3. ...
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Line 56: Note that experimental data on sprawling animals needs to be updated. Line 66: Please consider exchanging "modern" with "cursorial".

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I commend the authors for their extensive data set, compiled over many years of detailed fieldwork. In addition, the manuscript is clearly written in professional, unambiguous language. If there is a weakness, it is in the statistical analysis (as I have noted above) which should be improved upon before Acceptance.

Neogene amphibians and reptiles (Caudata, Anura, Gekkota, Lacertilia, Testudines) from south of Western Siberia, Russia and Northeastern Kazakhstan

Davit Vasilyan ^{Corresp., 1,2,3}, Vladimir S. Zazhigin ⁴, Madelaine Böhme ^{1,5}

¹ Department of Geosciences, Eberhard-Karls-University Tübingen, Tübingen, Germany

² JURASSICA Museum, Porrentruy, Switzerland

³ Department of Geosciences, University of Fribourg, Fribourg, Switzerland

⁴ Institute of Geology, Russian Academy of Sciences, Moscow, Russia

⁵ Senckenberg Center for Human Evolution and Palaeoecology, Tübingen, Germany

Corresponding Author: Davit Vasilyan

Email address: davit.vasilyan@jurassica.ch

Background. The present-day amphibian and reptile fauna of Western Siberia is the least diverse of the Palearctic Realm, as a consequence of the unfavourable climatic conditions that predominate in this region. The origin and emergence of these herpetofaunal groups are poorly understood. Aside from the better-explored European Neogene localities yielding amphibian and reptile fossil remains, the Neogene herpetofauna of Western Asia is understudied. The few available data need critical reviews and new interpretations, taking into account the more recent records of the European herpetofauna. The comparison of this previous data with that of the European fossil record would provide data on palaeobiogeographic affiliations of the region as well as on the origin and emergence of the present-day fauna of Western Siberia. An overview of the earliest occurrences of certain amphibian lineages is still needed. In addition, studies that address such knowledge gaps can be useful for molecular biologists in their calibration of molecular clocks.

Methods and Results. In this study, we considered a critically reviewed available data from amphibian and reptile fauna from over 40 Western Siberian, Russian and Northeastern Kazakhstan localities, ranging from the Middle Miocene to Early Pleistocene. Herein, we provided new interpretations that arose from our assessment of the previously published and new data. More than 50 amphibian and reptile taxa were identified belonging to families Hynobiidae, Cryptobranchidae, Salamandridae, Palaeobatrachidae, Bombinatoridae, Pelobatidae, Hylidae, Bufonidae, Ranidae, Gekkonidae, Lacertidae and Emydidae. Palaeobiogeographic analyses were performed for these groups and palaeoprecipitation values were estimated for 12 localities, using the bioclimatic analysis of herpetofaunal assemblages.

Conclusions. The Neogene assemblage of Western Siberia was found to be dominated by groups of European affinities, such as Palaeobatrachidae, *Bombina*, *Hyla*, *Bufo bufo*, and a small part of this assemblage included Eastern Palearctic taxa (e.g. *Salamandrella*, *Tylototriton*, *Bufo viridis*). For several taxa (e.g. *Mioproteus*, *Hyla*, *Bombina*, *Rana temporaria*), the Western Siberian occurrences represented their most eastern Eurasian records. The most diverse collection of fossil remains was found in the Middle Miocene. Less diversity has been registered towards the Early Pleistocene, potentially due to the progressive cooling of the climate in the Northern Hemisphere. The results of our study showed higher-amplitude changes of precipitation development in Western Siberia from the Early Miocene to the Pliocene, than previously assumed.

1 **Neogene amphibians and reptiles (Caudata, Anura,**
2 **Gekkota, Lacertilia, Testudines) from south of Western**
3 **Siberia, Russia and Northeastern Kazakhstan**

4

5 Davit Vasilyan^{1, 2, 3}, Vladimir S. Zazhigin⁴, and Madelaine Böhme^{1,5}

6

7 ¹ Department of Geosciences, Eberhard-Karls-University Tübingen, Sigwartstraße 10, 72076

8 Tübingen, Germany.

9 ² JURASSICA Museum, Route de Fontenais 21, 2900 Porrentruy, Switzerland,

10 davit.vasilyan@jurassica.ch.

11 ³ Department of Geosciences, University of Fribourg, Chemin du musée 6, 1700 Fribourg,

12 Switzerland.

13 ⁴ Institute of Geology, Russian Academy of Sciences, Pyzhevsky per. 7, 119017 Moscow, Russia,

14 zazhvol@gmail.com.

15 ⁵ Senckenberg Center for Human Evolution and Palaeoecology, Sigwartstraße 10, 72076

16 Tübingen, Germany, m.boehme@uni-tuebingen.de.

17

18 Corresponding author:

19 Davit Vasilyan

20 Current address: JURASSICA Museum, Route de Fontenais 21, 2900 Porrentruy, Switzerland

21 Email address: davit.vasilyan@jurassica.ch

22

23 **Abstract**

24 **Background.** The present-day amphibian and reptile fauna of Western Siberia is the least
25 diverse of the Palaearctic Realm, as a consequence of the unfavourable climatic conditions that
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28 and reptile fossil remains, the Neogene herpetofauna of Western Asia is understudied. The few
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30 recent records of the European herpetofauna. The comparison of this previous data with that
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40 new data. More than 50 amphibian and reptile taxa were identified belonging to families
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42 Pelobatidae, Hylidae, Bufonidae, Ranidae, Gekkonidae, Lacertidae and Emydidae.
43 Palaeobiogeographic analyses were performed for these groups and palaeoprecipitation values
44 were estimated for 12 localities, using the bioclimatic analysis of herpetofaunal assemblages.

45 **Conclusions.** The Neogene assemblage of Western Siberia was found to be dominated by
46 groups of European affinities, such as Palaeobatrachidae, *Bombina*, *Hyla*, *Bufo bufo*, and a small
47 part of this assemblage included Eastern Palearctic taxa (e.g. *Salamandrella*, *Tylototriton*,
48 *Bufo viridis*). For several taxa (e.g. *Mioproteus*, *Hyla*, *Bombina*, *Rana temporaria*), the
49 Western Siberian occurrences represented their most eastern Eurasian records. The most
50 diverse collection of fossil remains was found in the Middle Miocene. Less diversity has been
51 registered towards the Early Pleistocene, potentially due to the progressive cooling of the
52 climate in the Northern Hemisphere. The results of our study showed higher-amplitude changes
53 of precipitation development in Western Siberia from the Early Miocene to the Pliocene, than
54 previously assumed.

55

56 **Introduction**

57 Western Siberia is a geographic region restricted to the territories of Russia and parts of
58 northern Kazakhstan. It includes the region between the Ural Mountains in the west, Central
59 Siberian Plateau in the east, and the Kazakh Plain and Altay Mountains, including the Zaisan
60 Lake in the south (Fig. 1). Western Siberia region incorporates the drainage basin of the major
61 Siberian rivers such as the Irtysh and Ob rivers, both flowing into the Kara Sea of the Arctic
62 Ocean. The region is characterised by a highly continental climate, under the influence of the
63 Westerlies (winds). The mean annual precipitation (MAP) is relatively uniform and varies from
64 400 mm in the north (415 mm at Omsk) to 200 mm in the south (255 mm at Pavlodar). The
65 region has a high relative humidity in summer due to labile convective heating and frequent
66 torrential rainfalls. The mean annual range of temperature reaches 4 °C and more (Omsk: cold

67 month temperature – CMT -19 °C, warm month temperature – WMT 20 °C, mean annual
68 temperature – MAT 0.4 °C; Semipalatinsk: CMT -16 °C, WMT 22 °C, MAT 3.1 °C; Lake Zaisan:
69 CMT up to -27 °C, WMT 23 °C; after *Müller & Hennings* (2000). The area is covered by diverse
70 biomes, namely the tundra ('cold steppe') and taiga (coniferous forests) biomes, which are
71 replaced by open landscapes in the north (tundra) and in the south (steppe). The region that
72 contains the studied Neogene outcrops is located in the transitional zone between the dry and
73 the more humid temperate biomes, where taiga, forest-steppe and steppe biomes are
74 distributed (*Ravkin et al., 2008*).

75 Due to the strong continental climate, the present-day herpetofauna in the territory of Western
76 Siberia is comparatively far less diverse, represented only by six to ten amphibian species and
77 seven reptile species (Table 1). It is assumed that the present distribution of amphibians and
78 reptiles in Western Siberia was strongly influenced by Quaternary climatic fluctuation (*Ravkin,*
79 *Bogomolova & Chesnokova, 2010*). According to *Borkin* (*Borkin, 1999*), the present-day
80 amphibian fauna of Western Siberia belongs to the Siberian region of amphibian distribution in
81 the Palaearctic Realm. According to different authors (e.g. *Kuzmin, 1995; Amphibiaweb, 2016*),
82 the region is inhabited by few amphibians, namely two species of salamanders and four to eight
83 species of anurans, belonging to five genera and five families (Table 1). This is the poorest
84 regional diversity of fauna in the Palaearctic Realm, without any endemic species. Only
85 *Salamandrella keyserlingii* and *Rana amurensis* are characteristic of the territory, but they are
86 widely distributed and are also found in smaller areas in the neighbouring regions (*Borkin,*
87 1999). The Western Siberian reptile fauna listing includes few species: *Natrix natrix*, *Elaphe*

88 *dione*, *Vipera berus*, *Vipera renardi*, *Gloydus halys*, *Zootoca vivipara*, *Lacerta agilis* and *Eremias*
89 *arguta* (Ananjeva et al., 2006; Ravkin, Bogomolova & Chesnokova, 2010).

90

91 **Geology and stratigraphy**

92 The Neogene sediments in Western Siberia have a wide distribution. Over many decades,
93 through systematic palaeontological studies and research in the Neogene and Quaternary
94 sediments of this area, rich fossil deposits of molluscan and small and large mammalian faunas
95 have been discovered (e.g. Zykin, 1979; Zykin & Zazhigin, 2008; Zykin, 2012). Based on the
96 studies of the small fossil mammals, the Neogene stratigraphy of the area is complemented
97 with biochronologic data. Continental sedimentation in the western part of the Siberian Plain
98 began in the Oligocene, after regression of the Turgai Strait in the Late Eocene, and continued
99 until the Quaternary period (e.g. Chkhikvadze, 1984, 1989; Tleuberdina et al., 1993; Malakhov,
100 2005). The sedimentary basin is surrounded by the Ural Mountains in the west, the Central
101 Kazakh Steppe and Altai-Sayan Mountains in the south, and the western margin of the Siberian
102 Plateau in the east. The surrounding regions deliver clastic material to the basin. Some
103 researchers include the Zaisan Basin, located to the west of the Altai-Sayan Mountains in this
104 territory (Borisov, 1963). The Neogene sediments are represented by lacustrine, fluvial, alluvial
105 and other continental depositions, overlying marine Eocene sediments. The thickest section
106 (300 m) of the Neogene and early Quaternary sediments occurs in the Omsk Basin. Neogene
107 strata outcrops are mainly found in the interfluves of the Irtysh and Ishim rivers (Gnibitenko,
108 2006; Zykin, 2012). All these sediments are terrestrial (fluvial and alluvial facies) and have
109 produced rich fossil layers of vertebrate fauna (Zykin, 2012). The vertebrate-bearing Neogene

110 sediments are found in several areas along the Irtysh River and its tributaries – Petropavlovsk-
111 Ishim (e.g. Petropavlovsk 1, Biteke 1A), Omsk (e.g. Novaya Stanitsa 1, Cherlak), Pavlodar (e.g.
112 Pavlodar, Baikadam) and the Novosibirsk areas (e.g. Kamen-na-Obi) (Fig. 1). Detailed geological
113 descriptions of the stratigraphic sections and fossil localities are summarised in *Zykin* (1979);
114 *Zykin & Zazhigin* (2004); *Gnibitenko* (2006); *Zykin* (2012).

115 The stratigraphic subdivision is based mainly on the Russian concept of svitas. A svita has
116 lithologic, biochronologic and genetic (sedimentologic) significance and has no precise
117 equivalent in Western stratigraphic theory and terminology (*Lucas et al.*, 2012). The
118 stratigraphy of Neogene sediments in Western Siberia is supported by magnetostratigraphic
119 investigations (e.g. *Gnibitenko*, 2006; *Gnibidenko et al.*, 2011), in which the recovered polarity
120 signals are combined with biochronologic data and correlated to the geomagnetic polarity time
121 scale (GPTS) (*Fejfar et al.*, 1997; *Vangengeim, Pevzner & Tesakov*, 2005; *Zykin, Zykina &*
122 *Zazhigin*, 2007). The biozonation is based on fast-evolving lineages of small mammals, mainly
123 jerboas (Dipodidae), hamsters (Cricetidae) and voles (Arvicolidae). Owing to these bio-
124 magnetostratigraphic data the mean temporal resolution of the late Neogene faunal record
125 from the Ob-Irtysh Interfluve is estimated to be approximately 200 kyr (Fig. 2, Table S1, Data
126 S2). The main sections of these vertebrate fossil localities are referred to certain svitas (e.g.
127 Kalkaman, Pavlodar, Irtysh Svitas), however, the stratigraphic assignment of three localities
128 Olkhovka 1A, 1B, 1C to svitas is not available (Fig. 2, Table S1). No fossils are available in the
129 initial deposits of the early Late Miocene.

130

131 **State-of-art in palaeoherpetological studies in Western Siberia**

132 The fossil record of amphibians and reptiles in Western Siberia, including the Zaisan Basin
133 record, remain largely unknown. There are very few works devoted to the studies of the
134 Western Siberian late Paleogene and Neogene herpetofaunal assemblages (e.g. *Chkhikvadze*,
135 1984, 1989; *Tleuberdina et al.*, 1993; *Malakhov*, 2005). The vast majority of data on fossil
136 amphibians and reptiles are represented as short notes or are mentioned in faunal lists (e.g.
137 *Bendukidze & Chkhikvadze*, 1976; *Chkhikvadze*, 1985; *Malakhov*, 2005). In this present
138 contribution we analysed the available data from specimens described below and from new
139 generated data as well.

140 The earliest report on Neogene fossil amphibians was compiled by *Iskakova* (1969), wherein
141 she described amphibian faunas from two Priirtyshian localities, Gusiniy Perelet and Karashigar.
142 Gusiniy Perelet is a well-renovated Late Miocene vertebrate fossil locality, situated on the
143 riverbank of the Irtysh River, within the town of Pavlodar. The sedimentary sequence in this
144 locality contains layers of different ages from the late Late Miocene until the late Early Pliocene.
145 Three localities (also ‘horizons’) within the town of Pavlodar (Pavlodar 1A, 1B, 3B) are grouped
146 into several svitas and can be distinguished from the Gusiniy Perelet vertebrate locality. The
147 fossil content of the Gusiniy Perelet locality comes from the lower horizon – Pavlodar 1A.
148 *Iskakova* (1969) described an amphibian fauna from this layer.

149 The age of the Karashigar locality is unclear. In a study by *Tleuberdina et al.* (1989), this locality
150 has been estimated to date back to the Late Oligocene; however, *Lychev* (1990) placed it in the
151 Middle Miocene, Kalkaman Svita (the list of the small mammal fauna; see Data S2). The
152 amphibian taxa described by *Iskakova* (1969) in the Priirtyshian localities (*Bombina* cf.
153 *bombina*, *Pelobates* cf. *fuscus*, *Bufo* cf. *viridis*, *Bufo* cf. *bufo*, *Rana* cf. *ridibunda*, *Rana* cf.

154 *temporaria*) were identified based mainly on the vertebrae (cervical, dorsal and sacral)
155 morphology, which is not diagnostic in frogs at that taxonomic level. *Chkhikvadze* (1984)
156 restudied the material from the Pavlodar 1A (= Gusiniy Perelet) locality and identified *Bufo* cf.
157 *raddei*, *Bufo* sp., *Pelophylax* cf. *ridibundus*, *Eremias* sp., and *Coluber* sp. In this study, we did not,
158 however; assess the material from the above-mentioned works in order to verify Chkhikvadze
159 (1984) taxonomic identifications. Our sample from this locality (Pavlodar 1A) (Table S1), did not
160 reveal any element listed in these earlier studies (*Chkhikvadze*, 1984).

161 *Chkhikvadze* (1984) summarised all known fossil amphibians and reptiles from the former
162 Union of Soviet Socialist Republics (USSR), including those from Western Siberia. Accurate
163 descriptions are not yet available for many of these species. The Middle Miocene Kalkaman
164 locality (*Tleuberdina*, 1993), presently known as Malyi Kalkaman 1 (*Zykin*, 2012), has provided a
165 diverse record of fossil herpetofauna. The fossil record of this locality was partially restudied
166 and amended by us, which included the collection of new material.

167 Over the last decade, fresh attempts has been made to study the herpetofauna from the
168 Western Siberian localities (*Malakhov*, 2003, 2004, 2005, 2009). In the resultant works,
169 undescribed material from several Neogene localities of Kazakhstan were summarised, revised,
170 and studied, thereby providing critical overviews. In spite of the advances of the recent years,
171 however, the Neogene herpetofauna from Western Asia remains largely unknown, with
172 available fossil material continuing to be insufficiently studied. The main goals of the present
173 study were, therefore, to assess the descriptions and taxonomic classifications of the new
174 amphibian and reptile fossil material collected by Vladimir Zazhigin (co-author), as well as
175 already published data so as to provide a comprehensive faunistic analysis and

176 palaeobiogeographic and environmental interpretations. To avoid confusion around the names
177 used by different authors in the Russian literature to describe the localities, we have provided
178 all known names for these studied fossil localities.

179

180 **Materials & Methods**

181 The new materials used in the present study were collected by V. Zazhigin (co-author) using the
182 screen-washing technique during his long-term excavations in different Western Siberian
183 localities from the 1960's to 2008. These localities outcrop along the riverbanks of the Irtysh,
184 Ishim and Ob rivers. This fossil material is stored in the Institute of Geology, Russian Academy
185 of Sciences under the collection numbers:

186 GIN 950/2001 (Baikadam), GIN 1107/1001 (Malyi Kalkaman 1), GIN 1107/2001 (Malyi Kalkaman
187 2), GIN 1106/1001 (Shet Irgyz 1) GIN 952/1001 (Petrovsk 1), GIN 1109/1001 (Znamenka),
188 GIN 640/5001 (Pavlodar 1A), GIN 951/1001 (Selety 1A), GIN 951/2001 (Kedey), GIN 948/2001
189 (Novaya Stanitsa 1A), GIN 1115/1001 (Borki 1A), GIN 1110/2001 (Cherlak), GIN 945/2001
190 (Beteke 1A), GIN 640/6001 (Pavlodar 1B), GIN 1130/1001 (Lezhanka 2A), GIN 1130/2001
191 (Lezhanka 2B), GIN 1111/1001 (Olkhovka 1A), GIN 1111/2001 (Olkhovka 1B), GIN 1111/3001
192 (Olkhovka 1C), GIN 1118/3001 (Peshniovo 3), GIN 1131/2001 (Isakovka 2), GIN 1131/1001
193 (Isakovka 1A), GIN 1131/3001 (Isakovka 1B), GIN 1117/1001 (Kamyshlovo), GIN 945/2001
194 (Beteke 1B), GIN 945/3001 (Beteke 1C), GIN 1112/1001 (Andreievka – Speranskoe), GIN
195 1108/2001 (Pavlodar 2B), GIN 1112/2001 (Andreievka 1), GIN 1129/2001 (Livenka), GIN
196 1129/1001 (Lezhanka 1), GIN 1108/3001 (Pavlodar 3A), GIN 950/3001 (Lebiazhie 1A), GIN

197 950/4001 (Lebiashie 1B), GIN 950/5001 (Podpusk 1), GIN 945/60001 (Beteke 2), GIN 946/2001
198 (Kamen-na-Obi), GIN 945/8001 (Beteke 4), GIN 664/2001 (Razdole).

199 Various groups of amphibians and reptiles are represented in the available material. A report of
200 part of this material, i.e. of the snakes and anguine lizards, has been published in a separate
201 paper (e.g. *Vasilyan, Böhme & Klembara*, 2016). The present study included an assessment of
202 the materials collected from four fossil sites in Kazakhstan: Akyspe (also known as Agyspe), Aral
203 Horizon, leg. by Bendukidze in 1977; Kentyubek, Turgai Basin; Ryzhaya II (Ryzhaya Sopka),
204 Zaisan Svita, Zaisan Basin, leg. in 1970; Ayakoz (known also as Ayaguz), Zaisan Basin, leg. in
205 1970-1971; Petropavlovsk 1/2¹, leg. 1972 (Table S1). In addition, the few available data from
206 the literature were included in this study (after critical revision) to amend the record of
207 herpetofaunal assemblages of some localities as well as to reassign and revise the stratigraphic
208 position of these localities using biochronologic information of small and large mammalian
209 fauna (see full list in the Datas S2, S3).

210 The photographs of the fossil material were taken using a digital microscope Leica DVM5000
211 (Tübingen) and inspected with a scanning electron microscope, FEI Inspect S (Madrid). The
212 figures and tables were produced using Adobe Photoshop and Illustrator programs. The
213 osteological nomenclature of this study followed that of *Vasilyan et al.* (2013) for the
214 salamander remains, that of *Sanchíz* (1998a) for frogs, that of *Daza, Aurich & Bauer* (2011) and
215 *Daza & Bauer* (2010) was used for Gekkota, and the lepidosaurian terminology of *Evans* (2008).

¹ In the town of Petropavlovsk two fossil sites (Petropavlovsk 1 (MN12) and Petropavlovsk 2 (MN14)) having different ages are known, see *Zykin* (2012). Since the enclosed collection label to the material indicates only 'locality Petropavlovsk, leg. 1972' any stratigraphic allocation of the fossils to one of those layers is impossible.

216 Based on the herpetofaunal assemblages, the palaeoprecipitation values for the fossil localities
217 were estimated using the method of bioclimatic analysis of the ecophysiological groups of
218 amphibian and reptile taxa (Böhme et al., 2006). For the localities considered to be 'poor' in
219 amphibian and reptile taxa, the range-through approach (Barry et al., 2002) was used, in which
220 the faunas of two or more localities with age differences less than 100 kyr and/or belonging to
221 a single stratigraphic unit – svita, were considered as one. The taxa, that were added to the
222 herpetofaunal assemblage using the range-through approach, are indicated in grey in Table S4.

223

224 **Institutional/collection abbreviations.**

225 **GPIT:** Paläontologische Sammlung der Universität Tübingen, Tübingen, Germany; **HC:** collection
226 of Marcela Hodrova (Prague University), now stored in GPIT; **MNCN:** Museo Nacional de
227 Ciencias Naturales, Madrid, Spain; **NMNHK:** National Museum of Natural History, Kiev, Ukraine;
228 **PIN:** Palaeontological Institute, Russian Academy of Sciences, Moscow, Russia; **GNM:** National
229 Museum of Georgia, Tbilisi, Georgia; **GIN:** Geologic Institute, National Academy of Russia,
230 Moscow, Russia.

231

232 **Anatomical abbreviations.**

233 **ao:** antrum olfactorium; **alo:** antrum pro lobo olfactorio; **dl:** dental lamina; **ds:** dental shelf; **hl:**
234 horizontal lamella; **is:** incisura semielliptical; **ff:** frontoparietal facet; **fcpr:** facial process of
235 maxilla; **fMx5:** foramina for mandibular division of the fifth cranial (trigeminal) nerve; **hfr:**
236 haemal foramen; **hl:** horizontal lamella; **lf:** lacrimal facet; **lg:** longitudinal groove; **lh:** lamina
237 horizontalis; **lp:** lateral processes; **ls:** lamina supraorbitalis; **mc:** Meckelian canal; **na:** neural

238 arch; **nc**: neural canal; **nf**: nasal facet; **onf**: orbitonasal foramina; **olf**: olfactory foramina; **pf**:
239 parasphenoid facet; **pfc**: palatine facet; **ph**: paries horizontalis; **prz**: prezygapophysis; **psz**:
240 postzygapophysis; **pv**: paries verticalis; **pxp**: premaxillary process; **pyp**: pterygapophysis; **sac**:
241 opening of superior alveolar canal; **sg**: symphyseal groove; **sf**: splenial facet; **tpr**: transverse
242 process.

243

244 **RESULTS**

245 **Systematic palaeontology**

246 Class **Amphibia** *Gray, 1825*

247 Order **Caudata** *Scopoli, 1777*

248 Family **Hynobiidae** *Cope, 1859*

249 Genus ***Salamandrella*** *Dybowski, 1870*

250 ***Salamandrella*** sp.

251

252 (Fig. 3D-3G)

253 **Localities and material examined.** Malyi Kalkamana 1, GIN 1107/1001-AM12, 1 right femur;

254 Selety 1A, GIN 951/1001-AM01 – -AM03, 3 trunk and GIN 951/1001-AM04, 1 caudal vertebra;

255 GIN 951/1001-AM05, 1 distal end of bone (humerus?); Novaya Stanitsa 1A, GIN 948/2001-

256 AM01 – -AM11, 11 trunk vertebrae; Lezhanka 2A, GIN 1130/1001-AM01 – -AM26, 26 trunk and

257 GIN 1130/1001-AM27 – -AM28, 2 caudal vertebrae; Cherlak, GIN 1110/2001-AM01 – -AM12, 12

258 trunk vertebrae; Lezhanka 2B, GIN 1130/2001-AM01, 1 trunk vertebra, GIN 1130/2001-AM02, 1

259 extremity bone; Olkhovka 1B, GIN 1111/2001-AM01, 1 trunk vertebra; Iskakovka 2A, GIN

260 1131/2001-AM01, 1 trunk vertebra; Andreievka – Speransko, GIN 1112/1001-AM01, 1 trunk
261 vertebra; Lezhanka 1, GIN 1129/1001-AM01 – -AM02, 2 trunk and GIN 1129/1001-AM03, 1
262 caudal vertebrae; Beteke 1C, GIN 945/3001-AM01 – -AM02, 2 trunk vertebrae.

263 **Description and comments.** The vertebrae have an elongated to nearly slender form. The
264 vertebral centrum is amphicoelous. The basapophyses at the vertebral centrum are either
265 absent or are present in the form of a small protuberance at the laterodorsal corners of the
266 anterior portion of the vertebral centrum (Fig. 3G). A pair of subcentral foramina is situated at
267 the basis of the transverse processes. The neural arch is tall in lateral view (Fig. 3F) and
268 relatively broad in dorsal view (Fig. 3D). The posterior edge of the pterygapophysis is
269 bifurcated. Sometimes the neural spine is present but in general the dorsal surface of the
270 neural arch is flat. The pre- and postzygapophyses have an elongated oval shape. In anterior
271 view, the neural canal has an outline of a regular pentagon. The transverse process is
272 uncapitate (Fig. 3D, 3G). The anterior and posterior alar processes are absent. The vertebrae
273 can be assigned to the family Hynobiidae based on: (1) the small size and their amphicoelous
274 centrum with circular articular surfaces; (2) the lack of or being weakly pronounced
275 basapophyses; (3) the lack of neural spine; (4) the notch on the posterior margin of neural arch;
276 (5) the fused rib-bearers; and (6) the intervertebrally exiting spinal nerve in both trunk and
277 caudal vertebrae (e.g. *Edwards*, 1976; *Venczel*, 1999a, 1999b). Further, characteristic features
278 can be observed on the vertebrae of representatives of the genus *Salamandrella*, namely the
279 absence of the subcentral foramen and the concave anterior margin of the neural arch that
280 reaches the middle part of the prezygapophyseal articular facets (*Venczel*, 1999b; *Ratnikov* &
281 *Litvinchuk*, 2009; *Syromyatnikova*, 2014) (Fig. 3D-3G). The detailed description of hynobiid

282 material from the Western Siberian localities and comparison with recent and fossil hynobiids is
283 provided in a forthcoming paper.

284

285 Family **Cryptobranchidae** *Fitzinger, 1826*

286 **Cryptobranchidae** indet.

287

288 (Fig. 3A-3C)

289 **Localities and material examined.** Pavlodar 1A (=Gusiniy Perelet), 1 fragmentary right dentary
290 and 2 fragments of jaw bones, for details about the stratigraphic allocation see the section
291 'Cryptobranchidae', unnr. PIN specimen.

292 **Description and comments.** Among the fragments, a posterodorsal portion of a large right
293 dentary, 27 mm in length, is present. In lingual view, the pars dentalis is composed entirely of
294 dental lamina and the subdental lamina is present, but reduced. The pars dentalis possesses 30
295 pedicels of pleurodont teeth. The subdental shelf inclines slightly ventrally. The lamina
296 horizontalis is prominent. The corpus dentalis above the Meckelian groove has a concave
297 surface. Ventrally, this surface possesses a ridge running parallel to the lamina horizontalis. The
298 cross section of the dentary shows a relatively low portion of cancellous bone and a dominance
299 of compact bone. The size of the bones, the form and structure of the pars dentalis and the
300 cross section of the bone are characteristic of giant salamanders (*Vasilyan et al., 2013*).

301

302 Family **Proteidae** *Gray, 1825*

303 Genus **Mioproteus** *Estes & Darevsky, 1977*

304 *Mioproteus* sp.

305

306 (Fig. 3H-3S)

307 **Localities and material examined.** Ryzhaya II (known also Ryzhaya Sopka), GNM unnr.

308 specimen, 2 trunk vertebrae; Malyi Kalkaman 2, GIN 1107/2001-AM01, 1 right premaxilla; Borki

309 1A, GIN 1115/1001-AM01, 1 trunk vertebra; Ayakoz, GNM unnr. specimen, 1 trunk vertebra;

310 Akespe, unnr. HC specimens, 3 vertebrae; Petropavlovsk 1/2, GNM unnr. specimen, 22

311 vertebrae.

312 **Description.** The preserved left premaxilla is fragmentary (Fig. 3P-3Q) and the posterior process

313 is broken off. In ventral view, the bone has a rough surface. The pars dentalis of the premaxilla

314 is located on the anterior side of the bone. The crowns of pleurodont teeth are missing and

315 only their pedicellar portions are preserved. The bone surface is slightly rough in dorsal view.

316 The lamelliform anterolateral ridge of the posterior process is high at the middle part of the

317 bone. The amphicoelous vertebrae are flat and wide. The centrum is dumb-bell in shape and

318 narrows to the middle region. The basapophyses, if present, are small and weakly developed.

319 Two subcentral foramina are present at the central part of the vertebral centrum. In lateral

320 view, the vertebra is low; the anterior and posterior zygapophyseal crests are pointed, forming

321 the dorsal border of the deep depressions anteriorly and posteriorly to the transverse process.

322 The middle part of the neural arch is lower than its cranial and caudal margins. The posterior

323 edge of the neural arch is forked (Fig. 3H) (not visible at Fig. 3M). The neural spine extends as

324 far as the preserved anterior margins of the neural arch, whereas posteriorly, it terminates

325 before the posterior margin of the neural arch. The preserved right pre- and postzygapophyseal
326 articular facets are ellipsoid.

327 **Comparison and comments.** A direct comparison with *Mioproteus* specimens from previous
328 reports was not possible due to the extremely scarce description of the skull elements
329 attributed to this taxon (e.g. *Estes & Darevsky, 1977; Miklas, 2002*). We therefore used the
330 material of *Mioproteus* sp. from the Grytsiv locality (Ukraine, earliest Late Miocene) (Fig. 3R-3S)
331 for the taxonomic identification of the fossil premaxilla from Malyi Kalkaman 2 (Fig. 3P-3Q). Our
332 comparison finds no differences in the premaxilla morphology between the Kazakhstan and
333 Ukrainian *Mioproteus* sp. The vertebrae from the Borki 1A and Ayakoz localities can be easily
334 assigned to the genus *Mioproteus* based on following characters: (1) robust vertebra with an
335 amphicoelous centrum; (2) a tall cranial margin of the neural arch; (3) the presence of the
336 basapophyses; (4) a distinct wide depression at the anterior base of the transverse process; (5)
337 intervertebrally exiting spinal nerves; and (6) a forked neural spine (*Edwards, 1976; Estes &*
338 *Darevsky, 1977; Ivanov, 2008*).

339

340 Family **Salamandridae** *Goldfuss, 1820*

341 Subfamily **Pleurodelinae** *Tschudi, 1838*

342 Genus **Chelotriton** *Pomel, 1853*

343 **Chelotriton** sp.

344

345 (Fig. 3T-3Y)

346 **Localities and material examined.** Malyi Kalkaman 1, GNM unnr. specimen, 1 trunk vertebra;
347 Ayakoz, GNM unnr. specimen, 1 trunk vertebra.

348 **Description.** The single fragmentary trunk vertebra of *Chelotriton* from the Malyi Kalkaman 1
349 locality has been scantily described (*Tleuberdina* et al., 1993, pp. 133-134). The centrum of the
350 vertebra is ophistocoelous and dorsally curved. Both the posterior one-third of the vertebra
351 and cotyle are broken. The condyle is dorsoventrally slightly compressed and oval in shape. The
352 middle part of the ventral surface of the centrum bears a pair of the foramina subcentrale. The
353 ventral bases of both transverse processes are pierced by a foramen (potentially the ventral
354 foramen for the spinal nerve).

355 The neural spine is tall, long, and almost equal in length to the vertebral centrum. The dorsal
356 surface of the neural spine has the form of an elongated isosceles triangle and it is covered by a
357 distinct pustular sculpture. The anterior margin of the neural spine is concave in outline. The
358 posterior half of the spine is wider than the anterior one (Fig. 3Y). In anterior view, the neural
359 arch and the neural canal have a triangular form. The roof of the neural canal is flat, on both
360 sides of the neural spine.

361 The pre- and postzygapophyses are damaged. The anterior portion of the left postzygapophysis
362 is present and it shows a horizontal surface. The anterior bases of both prezygapophyses at the
363 contact with the centrum possess small subprezygapophyseal foramina. Behind the left
364 prezygapophysis, the accessory alar process exhibits a marked step (Fig. 3Y), projects
365 posteroventrally and connects caudally with the anterior alar process. The contact point of the
366 accessory and anterior alar processes probably corresponds to the base of the parapophysis.
367 Both transverse processes are broken, but the bases are preserved. Apparently, two rounded

368 upper and lower prominences, seen in left lateral view, correspond to the dia- and
369 parapophysis. The parapophysis is located anteriorly and dorsally to the level of the
370 diapophysis; thus, the transverse process becomes a bent projection. The arterial canal runs
371 behind the base of the transverse process. Anteriorly, its dorsal and ventral walls are built by
372 the accessory and anterior alar processes.

373 The vertebra from the Ayakoz locality (Fig. 3T-3X) is fragmentary, its neural arch and left
374 transverse process are lost, the centrum is compact, short and wide, and it possesses an
375 elliptical central foramen. The diapophysis of the preserved right transverse process is broken,
376 but it can be assumed that the dia- and parapophysis were separated from each other. The
377 accessory alar process runs from the prezygapophysis to the dorsal edge of the diapophysis.
378 The posterior and anterior alar processes run from the cotyle and condyle straight along the
379 transverse process to the parapophysis. This morphology is characteristic of the first trunk
380 vertebrae.

381 **Comparison and comments.** This vertebra was previously described by *Tleuberdina et al.*
382 (1993). Here we have assigned this specimen to the genus *Chelotriton* owing to the presence of
383 a triangular and well-sculptured plate on the top of the neurapophysis. This character,
384 however, is not a unique feature of *Chelotriton* and is also seen in other salamanders, e.g.
385 Recent species of *Tylotriton* and *Echinotriton*, and in *Cynops pyrrhogaster*, *Lissotriton boscai*
386 (unnr. GPIT specimen), *Paramesotriton* (MNCN 23557, 13645), as well as the fossil taxa
387 *Archaeotriton* (Böhme, 1998), aff. *Tylotriton* sp. (Baikadam locality, this paper),
388 *Carpathotriton* (Venczel, 2008). The vertebra from the Malyi Kalkaman 1 resembles the species
389 of *Chelotriton*, *Paramesotriton*, *Tylotriton*, *Echinotriton*, *Cynops pyrrhogaster*, and

390 *Carpathotriton* in their mutual presence of a subprezygapophyseal foramen. The vertebra can,
391 however, be justified as *Chelotriton* sp. and distinguished from other salamanders by: (1) its
392 longer length (vs. *Echinotriton*, *Cynops* and *Carpathotriton*); (2) a longer neural spine with a
393 rugose sculptured and triangular dorsal surface (vs. aff. *Tylototriton* sp., Baikadam locality, this
394 paper); and (3) a well-pronounced accessory alar process (vs. *Tylototriton*).

395 The fragmentary vertebra from the Ayakoz locality can be assigned also to this group because
396 of the presence of massive rib-bearers and large dimensions (*Ivanov*, 2008). Its vertebra is
397 identical to that of vertebra of *Chelotriton* sp. type II described from the Mokrá-Western
398 Quarry, 2/2003 Reptile Joint (Early Miocene, Czech Republic) (*Ivanov*, 2008).

399 The abundant European Cenozoic record of the genus *Chelotriton*, however, showed that
400 vertebral morphology is insufficient for taxonomic identification as *Chelotriton* (*Böhme*, 2008).
401 This genus has an unknown higher diversity, which can be uncovered by the study of complete
402 skeletons of those species. We hence classified the vertebrae from studied localities as aff.

403 *Chelotriton* sp.

404

405 Genus *Tylototriton* Anderson, 1871 (*Anderson*, 1871)

406 aff. *Tylototriton* sp.

407

408 (Fig. 4A-4K)

409 **Locality and material examined.** Baikadam, GIN 950/2001-AM01, -A14–A17 5 trunk vertebrae;

410 Ayakoz, GNM unnr. specimen, 2 trunk vertebrae.

411 **Description.** All preserved vertebrae are opisthocoelous. The condyle and cotyle are
412 dorsoventrally compressed. The vertebrae are slender, slightly narrow, and high. The neural
413 canal is round, but in anterior view, the ventral margin of the neural canal is flat. The same
414 occurs with the dorsal wall of the vertebral centrum. The centrum is dorsally curved in lateral
415 view (Fig. 4A, 4F, 4G). The neural spine was most probably high but does not reach the level of
416 the pustular region. The neural spine begins behind the cranial margin of the neural arch. The
417 neural arch is tilted dorsally and does not extend beyond the posterior edge of the
418 postzygapophysis. The dorsal plate of the neural spine is short, poorly developed, and covered
419 with rugosities. It has the form of an isosceles triangle. Due to the concave shape of the
420 posterior margin of the caudal border, we suggest that the neural spine was probably
421 bifurcated. The length of the neural spine, without the sculptured structure, is the same in all
422 preserved vertebrae and corresponds nearly to almost half of the entire vertebral length (Fig.
423 4A, 4F, 4G).

424 The pre- and postzygapophyses are horizontal and almost at the same level (e.g. Fig. 4A). The
425 pre- and postzygapophyseal articular facets are oval in shape. Small subprezygapophyseal
426 foramina are present at the level of the connection between the anterior bases of both
427 prezygapophyses with the vertebral centrum. The posterolaterally directed transverse process
428 is horizontally flattened and displays a bicapitate articulation surface with the rib. The
429 diapophysis and parapophysis are separated, with the former being smaller than the latter. A
430 low and moderately deep notch is developed at the posterior edge of the neural arch. The
431 transverse process has an anterior (accessory alar process) and posterior laminar edges (i.e. the
432 posterior alar process and dorsal lamina). The straight, posteroventrally directed accessory alar

433 process connects the prezygapophysis caudally with the base of the parapophysis (e.g. 4F). The
434 dorsal lamina starts from the diapophysis and extends to the postzygapophyses, whereas the
435 lamelliform posterior alar process starts at the parapophysis and terminates directly before the
436 cotyle. Subparallel to the accessory alar process, a thin anterior alar process runs along the
437 cranial half of the centrum. Behind and in front of the transverse process two 'cavities' (a
438 shallow anterior and a deep posterior) are present. These 'cavities' are connected by a canal
439 (possibly an arterial canal), that runs through the transverse process. In ventral view, the
440 vertebral centrum does not possess a ventral keel. The centrum is flattened and nearly plane in
441 the middle portion. Its surface is rough and pierced by numerous foramina. Two large
442 subcentral foramina are located at the posterior corner between the centrum and transverse
443 process (Fig. 4C, 4L).

444 **Comparison and comments.** The vertebrae resemble the morphology of pleurodeline
445 salamanders *Echinotriton*, *Tylotriton*, *Cynops*, *Chelotriton*, *Paramesotriton* and *Tylotriton*
446 and *Carpathotriton* in characteristics such as: (1) the presence of rugosities on the neural arch;
447 (2) the connection of the prezygapophysis and parapophysis with the accessory alar process,
448 except in *Carpathotriton*, *Cynops* and cf. *Tylotriton* sp. from Möhren 13 (*Böhme*, 2010: p. 11,
449 fig. 6f), where this process connects prezygapophysis with diapophysis; (3) a moderately
450 developed posterior 'cavity' behind the transverse process; and (4) the presence of
451 subprezygapophyseal foramen (for collection references see subsection 'Comparison' of
452 *Chelotriton* sp. in this report). In terms of the general morphology, the vertebrae mainly
453 resemble the genus *Tylotriton* and differ from the compared genera in having: (1) a low,
454 elongate, narrow and lesser flattened vertebrae; (2) a weakly developed pustular structure of

455 the neural arch (similar character as seen in *Paramesotriton*); (3) a low and long neural spine
456 without the sculptured structure; (4) a dorsoventrally compressed cotyle and condyle; (5) a
457 deep posterior 'cavity' behind the transverse process, and an extended dorsal lamina and
458 posterior alar process; (6) a low and shallow posterior notch of the neural arch; and (7) in
459 having an accessory alar process that reaches the parapophysis, which differs from specimens
460 of the genus *Cynops* wherein it reaches the diapophysis. The Siberian *Tylototriton* differs from
461 the European Oligocene cf. *Tylototriton* (see *Böhme*, 2010: p. 11, fig. 6f) by having: (1) a
462 ventrally deflected accessory alar process that terminates ventrally to the parapophysis; (2) a
463 shorter and lower neural spine; and (3) a shorter dorsal plate of the neural spine.
464 Taking into account the above-mentioned differences, we suggest that the described vertebrae
465 should be assigned to a new pleurodeline salamander genus that shows affinities with the
466 genus *Tylototriton*. However, we do not consider it reasonable to describe a new form unless
467 cranial material of this salamander is available.

468

469 Order **Anura** *Fischer von Waldheim*, 1813470 Family **Palaeobatrachidae** (*Cope*, 1865)471 **Palaeobatrachidae** sp. indet.

472

473 (Fig. 5A-5D)

474 **Locality and material examined.** Novaya Stanitsa 1A, GIN 948/2001-AM12, 1 sphenethmoid.

475 **Description.** This specimen is represented by a very robust sphenethmoid that lacks the
476 posterior region. The two anterior cavities corresponding to the antrum olfactorium are

477 anteroposteriorly shallow. The posterior cavity, antrum pro lobo olfactorio, is deep and narrow
478 (Fig. 5A, 5B). The olfactory foramen is larger than the orbitonasal foramen (Fig. 5C). The
479 processus rostralis is elongated and projects anteriorly. Anteriorly, dorsal face of the bone, two
480 sharply marked crescentic depressions (nasal facets) correspond to the contacts with the nasal
481 bones (Fig. 5A). In dorsal view, the frontoparietal facet (contacting with the frontoparietal
482 cranial bones) shows a slightly striated surface. The lateral processes protrude laterally. The
483 lamina supraorbitalis is well developed. The most anterior part of the incisura semielliptical is
484 preserved on the specimen. The remaining part of this structure demonstrates that it
485 approaches cranially to the anterior border of the bone. The ventral face of the sphenethmoid
486 possesses a narrow and long depression corresponding to the contact area with the cultriform
487 process of the parasphenoid (the parasphenoid facet) (Fig. 5B).

488 **Comparison and comments.** The bone has strong similarities to that of palaeobatrachids in
489 having: (1) a long sphenethmoid with a frontoparietal fenestra corresponding to more than half
490 of the bone length; (2) the articulation area of the parasphenoid delimited by two parallel
491 ridges, in ventral view; (3) a very short septum nasi and lateral process (*Vergnaud-Grazzini &*
492 *Młynarski, 1969; Sanchíz & Młynarski, 1979*). The palaeobatrachid from the Novaya Stanitsa 1A
493 locality exhibits all these characters aside from the short septum nasi, which is longer in the
494 fossil bone. We presume that the frontoparietal fenestra was longer more than half of the
495 sphenethmoid length because the overall length of the frontoparietal and nasal facets has
496 similar proportions as these seen in other palaeobatrachids. Furthermore, according to *Venczel,*
497 *Codrea & Fărcaş (2012)*, the sphenethmoidal ossification forms the anterior margin of

498 frontoparietal fontanelle in palaeobatrachid frogs (*Palaeobatrachus* + *Albionbatrachus*), which
499 can also be observed in the studied specimen.

500

501 Family **Bombinatoridae** Gray, 1825

502 Genus ***Bombina*** Oken, 1816

503 ***Bombina*** sp. / ***Bombina*** cf. ***bombina*** (Linnaeus, 1761)

504

505 (Fig. 6A-6F)

506 **Localities and material examined.** Malyi Kalkaman 2, GIN 1107/2001-AM02, 1 ilium; Selety 1A,
507 GIN 1107/2001-AM06, 1 ilium; Cherkal, GIN 1107/2001-AM06, 1 ilium.

508 **Description.** The bone description is based on the ilium from the Selety 1A locality, since the
509 specimens from the Malyi Kalkaman 2 and Cherkal localities are greatly damaged. In lateral
510 view, the iliac shaft is almost straight and lacks the dorsal crest. The tuber superior is a weakly
511 pronounced tubercle. In dorsal view, a spiral groove is observable and continues on the medial
512 surface of the shaft. The acetabulum is round and strongly extended (Fig. 6A). The junction
513 between the iliac shaft and corpus ossi is slightly constricted and the ventral base of the corpus
514 ossi possesses a preacetabular fossa. The ventral ridge of the acetabulum is high. In lateral and
515 posterior views, the pars descendens is reduced and wide, whereas the pars ascendens is high
516 but narrow (Fig. 6A, 6B). In ventral view, the pars descendens is broad and nearly flat. In medial
517 view, the acetabular area is bordered by shallow ridges, between which there is, a triangular
518 and medially prominent interiliac tubercle (Fig. 6B, 6C).

519 **Comparison and comments.** The lack of the vexillum and a poorly developed tuber superior is
520 characteristic of the genus *Bombina* (Böhme, 1977). The ilium differs from *Bombina orientalis*
521 by a poorly developed tuber superior. The ilium from the Selety 1A locality is distinguishable
522 from *Bombina variegata* and resembles *Bombina bombina* in having: (1) a developed pars
523 descendens; (2) a posteroventral ridge of the pars descendens projecting ventrally rather than
524 posteriorly (Böhme, 1977); and (3) a developed preacetabular fossa (Sanchíz & Młynarski,
525 1979). We, therefore, tentatively assign the bone to *B. bombina* due to the absence of well-
526 preserved material of the fire-bellied toads from the Selety 1A locality. The specific assignment
527 of the ilia from the Malyi Kalkaman 2 locality is impossible due to their fragmentary
528 preservation; therefore we describe them as *Bombina* sp.

529 The specimen from the Cherlak locality (Fig. 6D-6F) is greatly damaged with only a few
530 observable characters remaining that allow for its identification within Bombinatoridae. The
531 identifying characters are: (1) a large pars descendens at its anterior section, but dorsally
532 reduced; (2) a present but larger tuber superior than that of the Maly Kalkaman 2 and Selety 1A
533 specimens (within the family, larger tuber superior are present in the *Barbatula* (Folie et al.,
534 2013)); and (3) although the ventral wall of the acetabulum is not preserved, the remaining part
535 of its base allows for the assumption that it was markedly pronounced. Due to the incomplete
536 preservation, the important characters needed for taxonomic identification, e.g. interiliac
537 tubercle and junctura ilioischiastica, cannot be observed. The ilium from the Cherlak locality
538 can, therefore, be tentatively referred to the family Bombinatoridae.

539

540 Family **Pelobatidae** Bonaparte, 1850

541 Genus *Pelobates* Wagler, 1830

542 *Pelobates* sp.

543

544 (Fig. 6G-6I)

545 **Localities and material examined.** Selety 1A, GIN 1110/2001-AM13, 1 right ilium.

546 **Description.** The corpus ossi and distal portion of the iliac shaft are present. The tips of the pars

547 descendens and pars ascendens are broken. The bone surface is smooth and there is no tuber

548 superior. An oblique posterolaterally-anteromedially directed spiral groove extends on the

549 dorsal surface. Laterally, the high and long pars ascendens possesses a supraacetabular fossa

550 (Fig. 6I). The junction between the iliac shaft and corpus ossi is not constricted. The

551 subacetabular groove is shallow and broad. The acetabulum has a nearly triangular form, with a

552 well-marked rim. In medial view, the corpus ilii possesses an interiliac facet with a rugose

553 surface. The interiliac facet is composed of a large lower and a small upper portions. A well-

554 developed interiliac tubercle is visible between these portions (Fig. 6G). The lower portion is

555 ventroposteriorly oblique, whereas the upper one is flat, less rugose and has a concave surface.

556 The rugose surface of the facet indicates an extensive contact between two ilia (Fig. 6G, 6H).

557 The acetabular dorsal tuber is higher than the ventral one.

558 **Comparison and comments.** The ilium can be assigned to the family Pelobatidae based on the

559 absence of a dorsal crest, the absence of a dorsal tubercle and the presence of an oblique spiral

560 groove on the dorsal surface (Roček et al., 2014). The bone has the same characters of the

561 genus *Pelobates*: (1) a high and long pars ascendens; (2) a well-developed spiral groove (Böhme,

562 2010); (3) the lack of a dorsal crest of the iliac shaft (Folie et al., 2013); and (4) a rugose surface

563 of the interiliac facet (*Rage & Hossini, 2000*). However, further identification of the ilium is
564 impossible, as it does not show relevant differences at the specific level.

565

566 Family **Hylidae** *Rafinesque, 1815*

567 Genus **Hyla** *Laurenti, 1768*

568 **Hyla savignyi** *Audouin, 1827*

569 **Hyla** gr. *H. savignyi*

570

571 (Fig. 6J-6O)

572 **Localities and material examined.** Shet-Irgyz 1, GIN 1106/1001-AM01, 1 left ilium; Novaya
573 Stanitsa 1A, GIN 948/2001-AM20, 1 maxilla, GIN 948/2001-AM13, 1 scapula and GIN 948/2001-
574 AM14, 1 sacral vertebra; Lezhanka 2A, GIN 1130/1001-AM29 – -AM32, 4 ilia, GIN 1130/1001-
575 AM33 – -AM36, 4 scapulae and GIN 1130/1001-AM41, 1 trunk vertebra; Cherlak, GIN
576 1130/1001-AM14 – -AM15, 2 ilia; Olkhovka 1B, GIN 1111/2001-AM02, 1 fragmentary ilium;
577 Pavlodar 2B, GIN 1108/2001-AM01 – -AM03, 3 ilia.

578 **Description.** The ilia from all localities resemble the same morphology, i.e. the tuber superior is
579 dorsally prominent and slightly laterally. The tuber superior is located at the anterior corner of
580 the acetabulum. The preserved iliac shaft is nearly cylindrical, slightly mediolaterally
581 compressed and is devoid of crista dorsalis. The supraacetabular part of the ilium is smaller
582 than the preacetabular. The ventroposterior margin of the iliac shaft is connected with the pars
583 descendens by an expanded preacetabular zone, building a broad and thin lamina. The
584 acetabulum has a nearly triangular form. The acetabular rim is prominent at its high

585 ventroanterior edge. The posterodorsal corner of the acetabulum ascends and builds a small
586 and prominent acetabular tuber (Fig. 6L). In medial view, the bone surface is smooth,
587 sometimes with a shallow depression in the middle part of the corpus ossi. In distal view, the
588 junctura ilioischiastica is slender, the acetabulum is high and the interiliac facet displays a well-
589 pronounced ventromedial expansion. The acetabular dorsal tuber is higher than the ventral one
590 (Fig. 6K).

591 The scapula, a triradiate element of the pectoral girdle, is comparatively long (Fig. 6M-6O). The
592 bone surface is relatively smooth and is pierced by several foramina. The corpus scapulae, the
593 middle part of the bone, is slender and long. The pars suprascapularis is preserved in a
594 fragmentary state and most probably was not high. In dorsal view, the elongate pars acromialis
595 is narrow and almost equal in length (Fig. 6M). The shorter and flattened processus glenoidalis
596 is slightly broad. The processus glenoidalis and pars acromialis are separated by relatively deep
597 sinus interglenoidalis (Fig. 6N). The margo posterior, at the corner of the processus glenoidalis
598 and corpus scapula, possesses an oval to elongated-oval angular fossa (Fig. 6N, 6O). The tear-
599 shaped glenoid fossa reaches the posterior corner of the processus glenoidalis. The crista
600 supraglenoidalis is slightly pronounced.

601 **Comparison and comments.** The Siberian fossil tree frog differs from already described fossils
602 and some recent species of the genus *Hyla*. The following recent material is available for
603 comparison: *Hyla savignyi*, Armenia (four individuals, unnr. GPIT specimen), *Hyla orientalis*,
604 Armenia (two individuals, unnr. GPIT specimen) and *Hyla arborea*, Germany ? (one individual,
605 unnr. GPIT specimen). The Siberian forms can be distinguished from *Hyla* sp. (Rudabánya
606 locality in Hungary, middle Late Miocene (Roček, 2005); Bois Roche Cave in France, early Late

607 Pleistocene (Blain & Villa, 2006)), *Hyla arborae* (TD8 locality in Spain, early Middle Pleistocene
608 (Blain, 2009)), *Hyla* cf. *arborea* (Mátraszőlős 2 locality in Hungary, middle Middle Miocene
609 (Venczel, 2004)), *Hyla* gr. *H. arborea* (Capo Mannu D1 Local Fauna in Italy, Late Pliocene
610 (Delfino, Bailon & Pitruzzella, 2011)), *Hyla* aff. *japonica* (Tologoy 38x, Baikal Lake, Russia, late
611 Late Pleistocene (Ratnikov, 1997)) and Recent *Hyla japonica* (Nokariya, 1983) in having: (a) a
612 fossa supraglenoidalis; (b) a slenderer and lower corpus scapula and pars suprascapularis; and
613 (c) a shorter and broader processus glenoidalis. Apart from these differences, the Siberian fossil
614 tree frogs resemble *Hyla* sp. from the Bois Roche Cave, France (Blain & Villa, 2006), and *H.*
615 *arborea* (one individual, unnr. GPIT specimen) in possessing a low and broad processus
616 glenoidalis. The Recent *H. savignyi* is the only tree frog showing a fossa supraglenoidalis like
617 the one present in the studied remains. The Recent *H. savignyi* also possesses other similarities
618 to the fossil tree frog, namely: (1) a slender junctura ilioischiadic; (2) the same position of the
619 tuber superior; (3) comparable acetabular tubers; and (4) a similar slightly curved pars
620 ascendens. There are, however, also differences between the Recent *H. savignyi* and the fossil
621 tree frog. The fossil tree frog has: (1) a dorsally and slightly laterally prominent tuber superior;
622 (2) a deeper and larger fossa supraglenoidalis; and (3) a ventromedial expansion of the
623 interiliac facet; whereas *H. savignyi* has: (1) a dorsally and laterally significantly prominent
624 tuber superior; (2) a shallow and small angular fossa; and (3) the interiliac facet devoid of
625 ventromedial expansion. Among other fossil tree frogs, the Western Siberian *Hyla* sp. has the
626 lowest and broadest processus glenoidalis. Another fossil tree frog *Hyla* sp. reported from the
627 Russian Platform in the Kuznetsovka locality (0.5-0.65 Ma) (Ratnikov, 2002: fig. 2), displays a
628 similar morphology of the ilium as in the Siberian fossil, i.e. the orientation of the tuber

629 superior and the form of the junctura ilioischiadica. Because of the observed differences in both
630 the Recent and fossil forms, as well as the similarities to *H. savignyi*, we assume that the fossil
631 tree frogs from Western Siberian and the Russian Platform, probably represent a new form
632 related to the group of *Hyla savignyi*.

633

634 Family **Bufonidae** Gray, 1825

635 Genus **Bufo** Laurenti, 1768

636 **Bufo bufo** (Linnaeus, 1758)

637

638 (Fig. 6P-6W)

639 **Localities and material examined.** Novaya Stanitsa 1A, GIN 948/2001-AM15, 1 left and GIN
640 948/2001-AM16 – -17, 2 right ilia, GIN 948/2001-AM18 – -19, 2 trunk vertebrae; Borki 1A, GIN
641 1115/1001-AM02, 1 sacral vertebra, GIN 1115/1001-AM03, 1 left ilium; Olkhovka 1B, GIN
642 1111/2001-AM04, 1 left, GIN 1111/2001-AM03, 2 right ilia and GIN 1111/2001-AM05, 1 trunk
643 vertebra; Olkhovka 1C, GIN 1111/3001-AM01, 1 left scapula, GIN 1111/3001-AM02, 1 trunk
644 vertebra and GIN 1111/3001-AM03, 1 urostyle; Lezhanka 2A, GIN 1130/1001-AM37, 1 left ilia,
645 GIN 1130/1001-AM38, 1 left scapula, GIN 1130/1001-AM39, 1 sacral and GIN 1130/1001-AM40,
646 1 trunk vertebrae; Isakovka 1B, GIN 1131/3001-AM01, 1 left ilium; Isakovka 1A: GIN 1131/1001-
647 AM01, -AM05, 2 right ilia; Peshniovo 3, GIN 1118/3001-AM01, 1 sacral vertebra; Lezhanka 1,
648 GIN 1129/1001-AM04, 1 trunk vertebra; Andreievka 1, GIN 1112/2001-AM01 1 right scapula.

649 **Description and comments.** The ilia are large and have a robust corpus ossi. The spiral groove is
650 broad and very shallow. The tuber superior is broad, low, covered with irregular tubercles, and

651 it is situated above the acetabulum (Fig. 6P). The smooth and concave pars descendens is more
652 developed than the pars ascendens. The ventral edge of the pars descendens is thin and
653 lamelliform. The preacetabular fossa is absent (Fig. 6P). In posterior view, the anterolateral
654 edge of the acetabular is strongly curved. The junctura ilioischiastica shows a higher acetabular
655 ventral tuber than the dorsal tuber, and the ventral half of the corpus ossi projects
656 ventromedially (Fig. 6P).

657 The scapula is a robust bone and is longer than it is high. The material is represented by all
658 ontogenetic series. The angular fossa is absent; a shallow groove on the ventral side of the pars
659 acromialis is present and well pronounced in larger individuals. The pars acromialis and corpus
660 scapula have nearly the same height. The pars suprascapularis laterally increases in height. The
661 pars suprascapularis and corpus scapulae (anterior) have smooth surfaces. The base of the
662 lateral edge of the fossa glenoidalis is elevated but does not project laterally. The crista
663 supraglenoidalis is well developed in larger individuals. The anterior margin is concave. The
664 base of the pars acromialis is high and thin (Fig. 6R). There is a shallow and expanded depression
665 in ventral view. The anteromedial margin of the pars acromialis possesses a low tubercle. The
666 transition from the corpus scapula to the pars acromialis is nearly straight and the wall is thin
667 (Fig 6S, 6T).

668 In several localities, the large-sized frog vertebrae and urostyle (Fig. 6U-6W) are present in
669 association with diagnostic elements (ilia and scapula) (e.g. Olkhovka 1C locality) or are isolated
670 (e.g. Pehsniovo 3 locality). These individuals of the same size can be assigned to the large *Bufo*
671 *bufo*. The morphological traits described above (e.g. lack of angular fossa on the scapula and
672 preacetabular fossa on ilium, general outline, form, and size of the scapula and ilium) as well as

673 the bone dimensions are the same as those found in the common toad *Bufo bufo* (Blain, Gibert
674 & Ferràndez-Cañadell, 2010).

675

676 Genus ***Bufotes*** Rafinesque, 1815

677 ***Bufotes viridis*** Laurenti, 1768

678

679 (Fig. 6X-6Z)

680 **Localities and material examined.** Baikadam, GIN 950/2001-AM02 – -AM04, 3 left and GIN
681 950/2001-AM05 – -AM09, 5 right ilia; Malyi Kalkaman 1, GIN 1107/1001-AM02 and -AM03, 1
682 left and 1 right scapulae; Malyi Kalkaman 2, GIN 1107/2001-AM03, 1 right scapula; Znamenka,
683 GIN 1109/1001-AM01 and -AM02, 1 left and 1 right scapulae, GIN 1109/1001-AM03 – -AM07, 5
684 left and GIN 1109/1001-AM08 – -AM11, 4 right ilia; Pavlodar 1A, GIN 640/5001-AM01 – -AM24,
685 240 left and GIN 640/5001-AM25 – -AM58, 34 right ilia, GIN 640/5001-AM63 – -AM77, 15 left
686 and GIN 640/5001-AM78 – -AM88, 11 right scapulae; Cherlak, GIN 1110/2001-AM16, 1 right
687 ilium; Selety 1A, GIN 951/1001-AM08 – -AM10, 3 left and GIN 951/1001-AM11 – -AM14, 4 right
688 ilia; Isakovka 1A, GIN 1131/1001-AM02 – -AM04, 3 left ilia; Kedey, GIN 951/2001-AM01 and – -
689 AM02, 1 left and 1 right ilia; Lebiazhie 1A, GIN 950/3001-AM01, 1 left scapula, GIN 950/3001-
690 AM01 2 left ilia; Lebiazhie 1B, GIN 950/4001-AM01, -AM02, 2 right ilia.

691 **Description and comments.** The iliac shaft is slightly lateromedially compressed and bears a
692 weakly pronounced depression along the middle section. The spiral groove between the corpus
693 ossi and iliac shaft is weakly developed. The tuber superior is low and possesses a uni- or

694 bilabiate protuberance in its central part. The angular fossa is well pronounced. The
695 anteroventral edge of the acetabular rim is straight. The pars descendens projects sharply
696 ventrally. There is no observable 'calamita' ridge (Fig. 6X). The remains show typical features for
697 *Bufotes viridis*: i.e. the form and shape of the tuber superior and acetabulum (Böhme, 1977;
698 Blain, Gibert & Ferràndez-Cañadell, 2010). Due to the absence of well-preserved material, we
699 prefer to tentatively assign the remains to the *Bufotes viridis* group.

700

701 ***Bufo*** sp.

702 **Localities and material examined.** Cherlak, GIN 1110/2001-AM17, 1 left scapula; Olkhovka 1A,
703 GIN 1111/1001-AM01, -AM02, 2 left ilia; Pavlodar 2B, GIN 1108/2001-AM04 – -AM06, 3 left ilia.

704 **Description and comments.** The greatly damaged ilia exhibits the typical morphology of the
705 genus *Bufo*, i.e. the iliac shaft without the dorsal crest and a spiral groove between the shaft
706 and corpus ilii (Böhme, 1977). There is a preacetabular fossa in the caudoventral corner of the
707 acetabulum. The tuber superior is eroded. In medial view, the pars descendens is
708 ventromedially directed.

709

710 Family **Ranidae** Batsch, A. J. G. K., 1796

711 Genus ***Pelophylax*** Fitzinger, 1843

712 ***Pelophylax*** sp.

713

714 (Fig. 6AA-6AD)

715 **Localities and material examined.** Malyi Kalkaman 1, GIN 1107/1001-AM04, 1 left ilium, GIN
716 1107/1001-AM13, 1 left articular; Malyi Kalkaman 2, GIN 1107/2001-AM04, -AM05, 2 right ilia ,
717 and GIN 1107/2001-AM06, 1 left ilium; Petropavlovsk 1, GIN 952/1001-AM01, 1 left ilium;
718 Olkhovka 1C, GIN 1111/3001-AM04, 1 right ilium; Kamyshovo, GIN 1107/1001-AM01, 1 right
719 scapula; Lezhanka 1, GIN 1129/1001-AM05, 1 left and GIN 1129/1001-AM06, 1 right ilia, GIN
720 1129/1001-AM07, 1 left scapula; Andreevka 1, GIN 1112/2001-AM02, 1 right and GIN
721 1112/2001-AM03, 1 left ilia; Livenka, GIN 1129/2001-AM01, 1 right ilium.

722 **Description and comments.** The ilia have a strong, oval, nearly vertically oriented and ventrally
723 well-defined high tuber superior. The dorsal crest is high; anteriorly it is often broken. The tuber
724 superior is high and slightly more S-shaped than the crest; a well-developed supraacetabular
725 fossa is present. Posterior to the tuber, the dorsal margin of the bone is bent ventrally towards
726 the acetabulum. In posterior view, the tuber superior is curved ventromedially (Fig. 6AA). The
727 junctura ilioschiadica is damaged but based on the preserved structures we speculate that it
728 was tall (Fig. 6AB).

729 The scapula is an elongated and short bone. In ventral view, a weakly-developed crista
730 supraglenoidalis is observable. It runs subparallel to the margo posterior and reaches the
731 middle part of the pars suprascapulars (Fig. 6AC, 6AD).

732 The characters listed above, i.e. like the form and orientation of bones, tuber superior and
733 crista supraglenoidalis, allow for the attribution of the fossils to the genus of the green (water)
734 frogs *Pelophylax* (Böhme, 1977; Sanchíz, Schleich & Esteban, 1993; Bailon, 1999; Blain, Bailon &
735 Agustí, 2007). Any further identification is impossible due to the fragmentary preservation of
736 the material.

737

738 Genus ***Rana*** Linnaeus, 1758 (*Linnaeus*, 1758)

739 ***Rana*** sp. / ***Rana temporaria*** Nilsson, 1842 (*Nilsson*, 1842)

740

741 (Fig. 6AE-6AH)

742 **Localities and material examined.** Ayakoz, unnr. HC specimens, numerous ilia, Baikadam, GIN

743 950/2001-AM10, 1 left, GIN 950/2001-AM11 – -AM13, and 3 right ilia; Malyi Kalkaman 1, GIN

744 1107/1001-AM05 – -AM09, 5 left ilia, GIN 1107/1001-AM10, 1 right ilia, GIN 1107/1001-AM01, -

745 AM11, 2 right scapula; Malyi Kalkaman 2, GIN 1107/2001-AM07, 1 right ilium, GIN 1107/2001-

746 AM08 – -AM13, 6 left ilia; Olkhovka 1C, GIN 1111/3001-AM05, 1 right ilium; Lezhanka 1, GIN

747 1129/1001-AM08, 1 left ilium; Kentyubek, unnr. HC specimens, 2 left ilia.

748 **Description.** The ilia have a reduced, compact, anteriorly directed and low tuber superior. The

749 lateral surface is rough. The dorsal crest is low. The pars descendens is more developed than

750 the pars ascendens (Fig.6AE). In posterior view, the junctura ilioschiadica is low (Fig. 6AF) in

751 comparison to the ilium of *Pelophylax* sp. (Fig. 6AA). The tuber superior projects dorsolaterally.

752 The pars descendens projects medially (Fig. 6AE).

753 The middle portions of both scapulae are preserved without the proximal parts of the pars

754 acromialis and suprascapularis. In dorsal view, a crista supraglenoidalis is observable at the

755 processus glenoidalis, which continues until the pars suprascapularis along the longitudinal axis

756 of the bone. It is very prominent and forms a lamelliform convex ridge. The base of the

757 processus glenoidalis is high and straight (Fig. 6AG, 6AH).

758 **Comments.** The ilia and scapulae morphology strongly resembles that of brown frogs genus,
759 *Rana* (Böhme, 1977). Due to the fragmentary preservation of the bone material, any precise
760 taxonomic identification of the frogs from nearly all localities was impossible. The comparison
761 with Recent species (e.g. *Rana temporaria* (unnr. GPIT specimen), *Rana dalamtina* (unnr. GPIT
762 specimen; Bailon, 1999), *Rana graeca* (unnr. GPIT specimen), *Rana arvalis* (unnr. GPIT
763 specimen), *Rana dybowskii* (MNCN 40459), *Rana amurensis* (unnr. GPIT specimen) etc.)
764 revealed more similarities with the European and Western Asiatic species rather than to
765 Eastern Asiatic brown frogs.

766 Only the Malyi Kalkaman 1 locality provided adequate material for specific identification. The
767 ilia and scapulae from this locality's material resembled the Recent species, *Rana temporaria*,
768 which has the widest distribution among the brown frogs in Eurasia. The fossil bones of brown
769 frogs from other Western Siberian localities are described here as *Rana* sp. Due to the poor
770 preservation of the ilia from the Kentyubek locality, it can be only identified at the family level.

771

772 Class **Reptilia** Laurenti, 1768

773 Order **Squamata** Opperl, 1811

774 Suborder **Gekkota** Cuvier, 1817

775 Family **Gekkonidae** Gray, 1825

776 Genus **Alsophylax** Fitzinger, 1843

777 **Alsophylax** sp.

778

779 (Fig. 7)

780 **Locality and material examined.** Cherlak, GIN 1110/2001-RE01 – -RE10, 10 right dentaries, GIN
781 1110/2001-RE11 – -RE24, 14 left dentaries, GIN 1110/2001-RE26 – -RE38, 13 left maxillae, GIN
782 1110/2001-RE39 – -RE43, 5 right maxillae, GIN 1110/2001-RE44, 1 cervical trunk vertebra;
783 Mynsualmas-MSA 3: 1 right maxilla, unnr. GPIT specimen.

784 **Description.**

785 *Tooth morphology.* The teeth are slender, unicuspid, and not narrowly arranged. All maxillaries
786 and dentary teeth are straight, except the most anterior ones on the dentary, which are
787 anteriorly lightly oblique. The central teeth on dental lamina of both the maxilla and dentary
788 are larger than the anterior and posterior ones (Fig. 7C, 7G). The cusps of maxilla teeth are
789 rarely posteriorly oriented. The most complete dentary bone contains at least 17 (in total,
790 probably 20) teeth, counted by both teeth and their alveoles (Fig. 7B-7D).

791 *Dentaries.* The dentary is a slender and elongated. In the symphyseal region, the bone is slightly
792 medially curved. The pars ventralis is assumed to be enlarged, due to the bone posteriorly
793 increasing in height. The dentary is characterised by a completely closed Meckelian canal,
794 which runs along approximately two-thirds of the bone length (Fig. 7B). The symphyseal
795 articulation surface is reduced. It does not build a pronounced articulation surface. The ventral
796 surface of the symphysis bears a longitudinal, posteriorly deepening symphyseal groove, visible
797 in both the lingual and ventral views (Fig. 7B-7D). The Meckelian canal is open posteriorly at
798 about the 15th-16th tooth position. The splenial facet on the dentary, the anterior margin of
799 Meckelian opening, shows a light concave and elongated surface (Fig. 7B-7E). In lateral view,
800 the bone is smooth, and the only complete dentary possesses five foramina that are arranged
801 in a longitudinal row (Fig. 7A). The size of the foramina increases slightly in the anteroposterior

802 direction, also changing in form from a more rounded outline to an oval appearance. The
803 position of the last mental foramen is arranged lingually in front of the posterior opening of the
804 Meckelian canal. The cavity of the Meckelian canal is divided in two, i.e. the upper and lower
805 subcanals, by a distinct horizontal lamella (Fig. 7E). The horizontal lamella runs parallel to the
806 lamina horizontalis and can be observed posteriorly behind the opening of the Meckelian canal.
807 The upper subcanal opens to the labial surface of the bone near to the mental foramina. The
808 symphyseal groove corresponds to the anterior opening of the lower subcanal. In lingual view,
809 the lamina horizontalis is situated in a low position. Its margin is rounded but not prominent. A
810 shallow and anteriorly extending dental shelf divides the lamina horizontalis from the dental
811 lamina (Fig. 7C). Posteriorly, the bone is nearly L-shaped in the transverse section. In all
812 observed specimens, the pars horizontalis is destroyed in the preserved bone. The caudal
813 portion of the paries verticalis shows bifurcation (Fig. 7E), which corresponds to the coronoid
814 insertion.

815 *Maxilla*. The preserved posterior part of the maxillary possesses a relatively low lacrimal facet
816 of the facial process of the maxilla (pars nasalis sensu *Estes* (1969)), while the latter is always
817 not preserved. The internal wall of the maxilla posteriorly bears a small distinct longitudinal
818 groove, running parallel to the lamina horizontalis (Fig. 7F-7H). The groove begins at the
819 posterior basis of the lacrimal facet and continuous until the preserved posterior tip of the
820 bone. The groove narrows at the middle section of the bone (at the position of the 3rd-4th last
821 tooth), where the lacrimal facet terminates. The lamina horizontalis is clearly visible, expands
822 laterally just under the tip of the lacrimal facet and builds a palatine facet (Fig. 7F-7H). The
823 lamina horizontalis becomes distinctly and posteriorly narrower but does not diminish fully at

824 the posterior end of the bone. The jugal process of the maxilla is bifurcated at its distal end (Fig.
825 7H). The maxillary lappet is damaged, but its base is preserved. The internal wall surface of the
826 maxilla contains few rugosities. Here an anteroposteriorly directed, fairly well-pronounced,
827 median ridge, is observed. In labial view, several foramina occur above the dental row. Some of
828 these foramina are arranged in a longitudinal line that corresponds to the foramina for the
829 mandibular division of the fifth cranial (trigeminal) nerve. This line runs parallel to the lamina
830 horizontalis. The last foramen in the row pierces the maxilla at the base of the lacrimal facet
831 under its tip. The bases of the facial process and maxillary lappet lay a relatively large superior
832 alveolar canal (sac, Fig. 7J, 7L) for the maxillary nerve and its accompanying blood vessel. The
833 remaining foramina at the maxilla are dispersed irregularly on the bone surface. The
834 premaxillary process is present, but it is highly damaged. The anterior basis of the lacrimal facet
835 is pierced by a foramen.

836 *Vertebra.* A single cervical vertebra of a gecko specimen shows an elongate amphicoelous
837 centrum (Fig. 7N-7P). The cotyles are approximately circular. In anterior view, the vertebra has
838 a semi-circular outline. In lateral view, the vertebra is anteroposteriorly compressed. The neural
839 arch is concave on both sides. The transverse processes are high, extremely short, and vertically
840 aligned. The distal end of the process is round. The haemal foramina are present at the lower
841 base of the transverse processes. The prezygapophyses are small and slightly prominent. The
842 neural arch is plane and triangular in outline. It possesses a slender and low neural crest. The
843 postzygapophyses are small, nearly invisible and are situated on the ventrolateral edges of the
844 pterygapophysis.

845 **Comparison and comments.** Numerous characters allow for the identification of the material as
846 a member of the family Gekkonidae. These characters are namely: (1) the amphicoelous
847 condition of the vertebra; (2) the maxillae and dentaries bearing numerous pleurodont,
848 isodont, densely packed, cylindrical, and slender monocuspid teeth; (3) the presence of a
849 medially extended dental shelf of the maxilla; and (4) the lingually closed Meckelian canal
850 (*Hoffstetter & Gasc, 1969; Daza, Alifanov & Bauer, 2012*). The gekkonid remains from the
851 Cherkak locality display a low number of teeth on the dentary (up to 20) and a rounded tooth
852 apex (making the teeth digitiform), which are diagnostic characters for the genus *Alsophylax*
853 (*Nikitina & Ananjeva, 2009*). Within the gekkonids, a low number of teeth (up to 20) is also
854 characteristic of *Mediodactylus russowi*, *Phelsuma laticauda*, and *Ph. serraticauda* (*Nikitina,*
855 *2009*). The Siberian fossil geckos can be distinguished from *Mediodactylus* by peculiarities of
856 the maxilla (i.e. the presence of a lingual longitudinal groove and a reduced row of foramina of
857 the trigeminal nerve), the dentary with a distinct and longer horizontal lamella, plus a reduced
858 symphyseal groove. The Recent genus *Phelsuma* can be excluded from consideration since
859 these geckos are restricted to the islands of the southwest part of the Indian Ocean and belong
860 to another zoogeographic zone. The fossil geckos resemble the Recent species *Alsophylax*
861 *pipiens* (see in *Estes (1969); tab. 2C*) in the presence of the prefrontal process and their short
862 row of foramina of the trigeminal nerve, which terminates below the prefrontal process.
863 Further comparison with the Recent genus *Alsophylax* is, however, impossible due to the lack of
864 available comparative osteological material of the Recent *Alsophylax* species.
865 Fossil geckos were present in the Early Miocene of Kazakhstan, as is evident from the
866 Mynsualmas-MSA 3 locality (unnr. GPIT specimen) (*Böhme & Ilg, 2003*). The re-studying of the

867 material revealed that the posterior fragment of a right maxilla shows morphology similar to
868 *Alsophylax* sp. from the Cherlak locality in the presence of a lingual longitudinal groove, the
869 absence of foramina at the posterior portion of the bone and a round tooth apex. The fossil
870 material, however, differs in its larger size (Fig. 7Q). Taking this difference as well as the
871 similarities into account, we tentatively consider the Mynsualmas record as cf. *Alsophylax* sp.
872 This fossil probably represents a larger *Alsophylax* species than those registered in the Western
873 Siberia.

874

875 Suborder **Lacertilia** Owen, 1842 sensu *Estes, Queiroz & Gauthier, 1988*

876 Family **Lacertidae** *Fitzinger, 1826*

877 Genus **Lacerta** *Linnaeus, 1758*

878

879 **Remarks.** The generic assignment of fossil lacertid remains is extremely difficult. This group is
880 anatomically generalised (*Lacera* sensu lato) and shows very few characteristic features (e.g.
881 bone and teeth morphology) for detailed taxonomic assignments (*Böhme, 2010; Böhme &*
882 *Vasilyan, 2014*).

883

884 **Lacerta** s.l. sp. 1.

885

886 (Fig. 8A)

887 **Material.** Baikadam, GIN 650/2001-RE07 – -RE09, 2 (3?) left dentaries, GIN 650/2001-RE10, 1
888 postsacral vertebra; Pavlodar 1A, GIN 640/5001-RE01 – -RE15, -RE41 – -RE4217 left dentaries,
889 GIN 640/5001-RE16 – -RE25, 10 right dentaries.

890 **Description.** The bones bear pleurodont bicuspid teeth. The most completely preserved
891 dentary possesses at least 20 teeth. The pars dentalis is tall , with its height corresponding to
892 two-thirds of the tooth length. The Meckelian groove is open ventrolingually. It starts from the
893 ventral side of the symphysis and posteriorly increases in height. The lamina horizontalis is
894 slightly curved. The anterior portion of the lamina horizontalis is high and broad, reaching its
895 maximal height in its middle section which corresponds to the tenth tooth position. Behind this
896 point, the lamina horizontalis articulates ventrally with the dorsal margin of the splenial and
897 gradually narrows posteriorly. The articulation surface is lingually exposed. The crista dentalis,
898 sensu *Roček* (1984), is not higher but is longer than the ventral margin of the lamina
899 horizontalis. The ventral margin of the crista dentalis, in its posterior half, bears an articulation
900 surface with the ventral margin of the coronoid. A lingually exposed articulation surface of the
901 splenial is located at the posterior portion of the ventral surface of the lamina horizontalis. Up
902 to eight small foramina are present in labial view (Fig. 8A).

903 **Comments.** See in *Lacerta* s.l. sp. 2.

904

905 ***Lacerta*** s.l. sp. 2.

906

907 (Fig. 8B)

908 **Material.** Pavlodar 1A, GIN 640/5001-RE27 – -RE33, 7 left dentaries, GIN 640/5001-RE34 – -
909 RE39, 6 right dentaries; Cherlak, GIN 1110/2001-RE51, 1 right dentary.

910 **Description.** The dentaries possess 19 bicuspid teeth. The pars dentalis is high with its height
911 corresponding to two-thirds of the teeth length. The lamina horizontalis is curved and
912 maintains almost the same height along its entire length. The lamina horizontalis decreases
913 slightly in height only at the 9th-10th tooth positions, where the splenial articulates with the
914 lamina horizontalis. The articulation facet is lingually exposed only in its most posterior portion.
915 The crista dentalis is short but is longer than the ventral margin of the lamina horizontalis. The
916 Meckelian groove is low and ventrolingually open. Up to seven small foramina are present in
917 labial view (Fig. 8B).

918 **Comments.** *Lacerta* s.l. sp. 2 differs from *Lacerta* s.l. sp. 1 in having: (1) a more curved lamina
919 horizontalis that maintains nearly the same height along its length; (2) a higher and broader
920 anterior portion of the lamina horizontalis; (3) a shorter crista dentalis; and (4) a lower
921 Meckelian groove.

922

923 ***Lacerta* s.l. sp. / Lacertidae indet.**

924 **Material.** Malyi Kalkaman 2, GIN 1107/2001-RE01, 1 vertebra; Olkhovka 1A, GIN 1111/1001-
925 RE01 and -RE02, 1 anterior and 1 posterior trunk vertebrae; Cherlak, GIN 1110/2001-RE06, -
926 RE52 – -RE57, 7 trunk vertebrae, GIN 1110/2001-RE47, -RE48, 2 left maxillae, GIN 1110/2001-
927 RE49, 1 right maxilla, GIN 1110/2001-RE50, 1 left dentary; Pavlodar 1A, GIN 640/5001-RE40, 1
928 premaxilla, GIN 640/5001-RE26, numerous fragments of dentaries and maxillae, GIN 640/5001-
929 RE43, 77 vertebrae; Pavlodar 1B, GIN 640/6001-RE01, -RE02, 2 left dentaries, GIN 640/6001-

930 RE03, -RE04, 2 right dentaries; Olkhovka 1B, GIN 1111/2001-RE01, 1 right dentary; Pavlodar 3A,
931 GIN 1108/3001-RE01, 1 right maxilla; Beteke 2, GIN 945/6001-RE01, 1 left dentary; Beteke 4,
932 GIN 945/8001-RE01, 1 left dentary.

933 **Description and comments.** The preserved maxillaries and dentaries possess pleurodont
934 bicuspid teeth. The Meckelian groove is lingually open. The labial surfaces of the maxillaries
935 show no ornamentation. In labial view, the foramina for mandibular division of the fifth cranial
936 (trigeminal) nerve are observable. They are situated along a longitudinal line, parallel to the
937 ventral margin of the bone. The opening of the superior alveolar canal is large. In lingual view, a
938 shallow but broad groove is present at the anterior portion of the frontal process. The large
939 foramen of the fifth cranial (trigeminal) nerve opens at the ventral surface of the lamina
940 horizontalis. A single premaxilla from Pavlodar 1A, GIN 640/5001-RE40 has a tapering nasal
941 process with a row of seven pleurodont and monocuspid teeth.

942 The bone material is extremely fragmentary, and any comparison between different localities
943 was impossible. The fossil remains (maxillae and premaxilla) from Pavlodar 1A do not show any
944 taxonomical differences, so we were not able to group them neither to *Lacerta* s.l. sp. 1 nor
945 *Lacerta* s.l. sp. 2. Besides the jaw material, vertebrae from the trunk region are available in the
946 Maly Kalkaman 2, Olkhovka 1A and Cherlak localities. It was not possible to identify all of
947 remains below the family level.

948

949 Genus *Eremias* Fitzinger, 1843

950 *Eremias* sp.

951

952 (Fig. 8C-8D)

953 **Material.** Pavlodar 2B, GIN 1108/2001-RE01, -RE02, 1 frontal and 1 trunk vertebra.

954 **Description.** The preserved frontal has a sandglass shape and the most anterior and posterior
955 portions are broken. The bone is slightly curved in lateral view. The posterior portion of the
956 dorsal surface is rough. The crista cranii are round and slightly elevated at the narrowest
957 portion of the bone. Anteriorly, these crista cranii increase in height and build the lateral walls
958 of the cranial vault. The anteroventral surface of the bone has two drop-shaped grooves. The
959 posteroventral surface is plain and slightly lower than the anteroventral surface. The prefrontal
960 facets are developed but do not show any lateral extension. The bone margin that connects
961 both facets is concave. In dorsal view, the nasal facets that are situated at the anterolateral
962 corners, are narrow, deep, and elongated (Fig. 8A, 8D).

963 In lateral view, a single preserved trunk vertebra has a rectangular shape. The neural arch is
964 moderately convex. A narrow and deep groove is present at the transition of the neural arch
965 and prezygapophysis. The neural spine is reduced and posteriorly builds a rounded process,
966 projecting over the posterior margin of the arch. The centrum is compressed anteroposteriorly
967 and possesses two shallow subcentral grooves, with a subcentral foramina in each one. The
968 condyle is small, round and is situated in the middle part of the posterior margin of the
969 centrum.

970 **Comments.** Among the Eurasian lacertids, fused dorsally sculptured frontals are known in
971 *Acanthodactylus*, *Eremias*, *Ophisops* (Evans, 2008). Our own observations of Recent species of
972 these genera (*Eremias strauchi*, *Eremias pleskei*, *Eremias arguta*, *Eremias multicellata*, *Ophisops*
973 *elegans*, *Acanthodactylus erythrurus*) allowed for the assignment of the frontals to the genus

974 *Eremias* and to separate them from: (1) *Ophisops* by a robust frontal, more pronounced
975 grooves at the anteroventral bone surface and a lack of the lateral extension of the prefrontal
976 facet; and (2) *Acanthodactylus* by a flat posteroventral surface of the bone and a less curved
977 outline in lateral view. The preserved single vertebra strongly resembles the morphology that is
978 found in *Eremias* (Rage, 1976).

979

980 Order **Testudines** Linnaeus, 1758

981 Suborder **Cryptodira** Cope, 1868

982 Family **Emydidae** (Rafinesque, 1815)

983 Genus ***Eymdoidea*** Gray, 1870²

984 ***Eymdoidea*** sp.

985

986 (Fig. 8E-8G)

987 **Material.** Novaya Stanitsa 1A, GIN 948/2001-RE01, 1 posteriorly incomplete right hypoplastron,
988 GIN 948/2001-RE02, 1 left femur.

989 **Description and comments.** The caudal part of the left hypoplastron, which has a width of 54.3
990 mm , is preserved (Fig. 8E) and probably belongs to a middle-sized individual with a total length
991 of the carapace, approximately 300 mm. In ventral view, the femoral/abdominal sulcus is nearly
992 straight and curves anteriorly only near the lateral edge of the bone, terminating at the base of
993 the inguinal buttress. The bone is comparatively thin, medially from the bridge (4 mm) to

² We follow taxonomy suggested by Fritz, Schmidt & Ernst, 2011 recognizing *Eymdoidea* as a distinct genus from *Emys*.

994 behind the bridge (7.2 mm). The lateral edge of the bone projects slightly posterolaterally. The
995 outline of the femoral/abdominal sulcus and the profile of the lateral edge are similar to those
996 of the emydid genus *Emydoidea* (both fossil and Recent specimens) ((*Chkhikvadze*, 1983), figs.
997 26 and 27, p. 138; (*Holman*, 1995)).

998 An almost complete left femur is available from the same locality where the hypoplastron
999 fragment was found. The bone is slender and bent (Fig. 8F-8G), and is 50.6 mm in length. This
1000 bone could have belonged to an individual of about 300 mm of the carapace length. The femur
1001 lacks its proximal portion (i.e. femoral head, major and minor trochanters). In ventral view, the
1002 fossa is delimited by the trochanters and is observable below the femoral head. The dimension
1003 of the bone is characteristic of aquatic testudinoids. Taking this latter character into account, as
1004 well as the comparable reconstructed total body-sizes of both elements (hypoplastron and
1005 femur) (ca. 300 mm), we consider the remains to belong to the genus *Emydoidea*.

1006

1007 **Testudines** indet.

1008 **Material.** Malyi Kalkaman², GIN 1107/2001-RE02, shell fragment; Shet-Irgyz 1, GIN 1106/1001-
1009 RE01, 1 neuralia; Petropavlovsk 1, GIN 952/1001-RE01, several fragments of carapax; Borki 1B,
1010 GIN 1115/2001-RE01, 1 fragment of carapax.

1011 **Comments.** The preserved remains were not sufficiently informative for any other taxonomic
1012 interpretation.

1013

1014 **DISCUSSION**

1015 **Neogene evolution of amphibian and reptile assemblages in Western Siberia**

1016 In general, and in contrast with the well-studied European fossil record, very is known about
1017 the Neogene herpetofauna from Asia. This record bias is owing to: (1) the less explored and less
1018 extensively studied Neogene deposits on the Asian continent; and (2) the entirely lack of study
1019 on Recent amphibians and reptiles, in spite of the intense investigations around small mammals
1020 by many scholars. The Western Siberian localities provide an exceptional opportunity to fill
1021 these gaps in information and to explore both the unknown diversity of the Asian
1022 herpetofaunal assemblages and the palaeobiogeographic affinities of the Western Siberian
1023 Neogene herpetofauna with the European faunas. Unfortunately, the yielded fossil material
1024 from this study and from previous investigations has thus far not been rich in amphibian and
1025 reptile remains. On average, only four taxa are available from each studied locality. Our
1026 faunistic, palaeogeographic and palaeoclimatic interpretations are, hence, very tentative and
1027 should be taken within this context. The unbiased comparison and analysis of our data are also
1028 hindered by the scarce record of the Asian Neogene fossil fauna. For the comparison with the
1029 European record, we used already published data on amphibian and reptile groups (families,
1030 genus, species, etc.) which have been summarised in the fosFARbase database (*Böhme & Ilg,*
1031 2003). These data are given in the Table S5. In the 'Europe' record, we consider all known fossil
1032 records from Western, Central, and Eastern Europe as well as from Anatolia (Fig. 9). By
1033 analysing the Neogene amphibian and reptile records from Europe and Asia, we were able to
1034 provide useful data that are applicable for fossil calibration of molecular clocks in the
1035 phylogenetic trees.

1036

1037 **Hynobiidae**

1038 The Asiatic salamanders (*Salamandrella* sp.) have the most abundant and frequent record
1039 among the studied Western Siberian localities. These organisms appeared in these areas in the
1040 middle Late Miocene (in the Selety 1A locality) and are present until the early Early Pleistocene.
1041 Although the herpetofaunal assemblages of the older localities are rich and represented by
1042 numerous taxa, they do not contain any hynobiid remains, demonstrating that there is no
1043 sampling bias in their record and that such specimens are not present in earlier localities.
1044 Recently, the oldest record of the genus, *Salamandrella* sp. has been described from the late (?)
1045 Early Miocene of Eastern Siberia (Lake Baikal) (*Syromyatnikova*, 2014), and a new species of
1046 *Salamandrella* is indicated to be present in the Late Miocene locality Ertemte 2, China (*Vasilyan*
1047 et al., 2013). Furthermore, the fossil Asiatic salamander, *Ranodon* cf. *sibiricus* was recovered
1048 from the Early Pleistocene of Southern Kazakhstan (*Averianov & Tjutkova*, 1995), and a
1049 *Salamandrella* sp. was reported from a few Middle Pleistocene age localities in European Russia
1050 (*Ratnikov*, 2010).

1051 In Central Europe, hynobiids (genus *Parahynobius*) appeared at the earliest Late Miocene and is
1052 present in the record until the Middle Pleistocene (*Venczel*, 1999a, 1999b; *Venczel & Hír*, 2013).
1053 According to our unpublished data, the hynobiids are also present in three Ukrainian localities –
1054 Grytsiv (11.1 Ma) (*Kirscher* et al., 2016), earliest Late Miocene; Cherevichnoe lower level,
1055 middle Late Miocene; and Kotlovina lower level, late Pliocene. The Ukrainian occurrences
1056 coincide with both the Central European and Western Siberian records of hynobiids, which at
1057 that time most probably characterised by favourable conditions for hynobiid distribution.
1058 Considering their oldest records, the origin of Hynobiidae was most probably in Eastern Asia in

1059 the early Miocene. We will present a detailed study on the Cenozoic record of fossil Hynobiidae
1060 including the Western Siberian material in a forthcoming paper.

1061

1062 **Cryptobranchidae**

1063 The cryptobranchid remains are known from two localities in the town of Pavlodar and from
1064 three localities in the Zaisan Basin. The stratigraphic positions of the Pavlodar localities are not
1065 clear. The only record of giant salamander that we were able to study is stored at the
1066 Palaeontological Institute of Moscow, Russia. The collection label provides the following
1067 information: 'collected by Gaiduchenko, in 1970, from the Gusiniy Perelet locality, at the
1068 contact of the Aral clays with overlying sands, about 200-300 m south far from the 'Gusiniy
1069 Perelet' [=Pavlodar 1A] locality'. The only explanation of the stratigraphic allocation of the
1070 giant salamander remains is that they originated from the basal horizon of the Pavlodar Svita,
1071 overlaying the 'Aral clays' (or= limnic clays of the Kalkaman Svita). *Gaiduchenko* (1984) and
1072 *Gaiduchenko & Chkhikvadze* (1985) mention a giant salamander (Cryptobranchidae indet.) from
1073 a locality named Detskaya Zheleznaja Doroga (engl. Children Railway) (Fig. 2, Table S1, Data S3),
1074 a sand pit located 10 km south-east from the 'Gusiniy Perelet' [=Pavlodar 1A] locality. The age
1075 of this fossiliferous horizon may fall near the Miocene-Pliocene boundary, an assumption that is
1076 mostly based on geology, age and accompanying fauna (see Data S2). This record from the
1077 Detskaya Zheleznaja Doroga presents the most northern (52.3° N) occurrence of the giant
1078 salamanders in the Northern Hemisphere known so far. Unfortunately, this material was not
1079 available for our study.

1080 Giant salamander remains have also been reported from three Burdigalian localities – Tri
1081 Bogatyrya, Vympel, Poltinik of the Zaisan Basin (Fig. 1, Table S1) (*Chkhikvadze, 1984; Böhme,*
1082 *Vasilyan & Winklhofer, 2012*). The remains were assigned to the species *Andrias karelcapeki* by
1083 *Chkhikvadze (1984)*. The taxonomic validity of the species still requires revision, which is
1084 necessary for any further interpretations.

1085

1086 **Proteidae**

1087 The oldest record of the genus is described as being from the Late Oligocene and was found in
1088 the Aral Formation in the Akespe locality, on the north coast of the Aral Sea, Kazakhstan (cf.
1089 *Mioproteus,*) (*Malakhov, 2003; Bendukidze, Bruijn & Van den Hoek Ostende, Lars W., 2009*).
1090 Here we add to the record a new, more recent Miocene (Aquitanian) Asian occurrence from the
1091 Ayakoz locality, Kazakhstan (Fig. 3D, Table S1). In the Middle Miocene, representatives of this
1092 genus occur in several localities in southern Russia and northern Kazakhstan (Table S1).
1093 According to our assessment, proteids survived until latest Miocene/earliest Pliocene (locality
1094 Petropavlovsk 1/2). The oldest stratigraphic record of *Mioproteus (Mioproteus caucasicus)* in
1095 Europe is described from the mid Aquitanian (early Early Miocene about 20.5-22 Ma) at two
1096 localities Ulm-Uniklinik and Ulm Westtangente of the North Alpine Foreland Basin (*Heizmann et*
1097 *al., 1989*). The fossil proteids are known in Europe until the Pleistocene Epoch (*Böhme & Ilg,*
1098 *2003*). Due to the lack of complete fossil skeletons and unclear taxonomic assignments of the
1099 fossil records, *Malakhov (2003)* preferred to refer all known specimens of *Mioproteus* to the
1100 ‘*Mioproteus caucasicus* complex’, including *Mioproteus* from Ashut, Kazakhstan,
1101 *Mioproteuscaucasicus* from type locality, as well as from the Late Miocene of Czech Republic,

1102 *Mioproteus wezei* from the Pliocene of Poland and from the Early Pleistocene of Moldavia
1103 (*Malakhov*, 2003). Later, *Roček* (2005) considered *M. wezei* as a junior synonym of *M.*
1104 *caucasicus*, although as already mentioned by *Malakhov* (2003), an adequate amount of
1105 material including cranial and postcranial elements is necessary to solve the taxonomic
1106 problems of the genus. *Malakhov* (2003) also suggested an Asiatic origin for the '*Mioproteus*
1107 *caucasicus* complex' and their later distribution into Europe. In summary, the oldest Late
1108 Oligocene record of *Mioproteus* (*Mioproteus* sp.) from Akespe, Kazakhstan and other localities
1109 of younger ages suggest: (1) a probable Asian origin of the genus; (2) the genus was
1110 continuously present in Central Asia/Western Siberia until the early Pliocene; and (3)
1111 *Mioproteus* migrated into Europe in the Early Miocene.

1112

1113 **Salamandridae**

1114 As has already been established, *Chelotriton* is a basket taxon (*Böhme*, 2008) that needs further
1115 taxonomic study. It is one of the fossil amphibians that has an abundant and wide distribution
1116 in the late Paleogene and Neogene localities of Europe. In Asia, the genus was previously
1117 known only from the late Middle Miocene locality Malý Kalkaman 1 (*Tleuberdina*, 1993). Our
1118 study showed that this genus was present at least since the Aquitanian age (the Aykoz locality
1119 in Kazakhstan, Early Miocene) (Table S1), making their Asiatic record older than previously
1120 assumed.

1121 Two localities (Ayakoz and Baikadam) from Western Siberia revealed aff. *Tylototriton*. The
1122 vertebrae showed significant similarities with the Recent East Asiatic genus *Tylototriton*. In
1123 *Böhme & Ilg* (2003) and *Böhme* (2010), the genus *Tylototriton* (cf. *Tylototriton* sp. and

1124 *Tylototriton* sp. nov.) has been reported from several Early Oligocene localities in southern
1125 Germany. Two Siberian records represent the first fossil occurrence of the genus in Asia, which
1126 appeared more recently in the fossil record than in the European occurrence. These Western
1127 Siberian specimens and the European specimens can be clearly separated from each other by
1128 the morphology of the trunk vertebrae. The Siberian salamanders probably represent new
1129 forms, strongly related to the East Asian terrestrial salamander, *Tylototriton*. The sympatric
1130 occurrence of two fossil terrestrial salamander genera *Chelotriton* and *Tylototriton* was
1131 documented for the first time from the Aquitanian age locality Ayakoz.

1132

1133 **Palaeobatrachidae**

1134 The palaeobatrachids are considered a European family, with probable occurrence in North
1135 America at the terminal Cretaceous (*Wuttke et al., 2012*). Records of the palaeobatrachids are
1136 known from the Paleogene Western and Central Europe. It should be taken into account,
1137 however, that records from the Paleogene of Turkey, as well as from the Paleogene and Early to
1138 Middle Miocene of Eastern Europe, are very scarcely known. In the Miocene, palaeobatrachids
1139 appear to have expanded their distribution to Eastern Europe and also reached Anatolia, where
1140 they existed from the latest Oligocene and remained during the entire Early Miocene. During
1141 the Middle Miocene, palaeobatrachids were present in Europe, from Germany to Ukraine
1142 (*Wuttke et al., 2012*). The palaeobatrachid record in Europe is characterised by a four-million-
1143 year-long (ca. 5.6-9.78 Ma) gap in the Late Miocene (Fig. 9). During this gap, no palaeobatrachid
1144 is known from Western to Eastern Europe even in localities rich in diverse herpetofaunal
1145 assemblages (e.g. Staniantsi, Bulgaria; Morskaya 2, Russia, *Böhme & Ilg, 2003*) and where

1146 characterised by favourable environmental conditions for their distribution. After this gap,
1147 palaeobatrachids occur near the Mio-Pliocene transition in studied localities from Italy (Ciabot
1148 Cagna), (*Cavallo et al.*, 1993) and Hungary (Osztramos 1C) (*Venczel*, 2001)). They seems to have
1149 disappeared from Western (Tegelen locality in Holland, *Villa et al.*, 2016) and Central Europe
1150 (Betfia IX/B locality in Romania, *Venczel*, 2000) after the Early Pleistocene and remained
1151 exclusively in Eastern Europe until the middle Pleistocene (Poland – European Russia) (*Wuttke*
1152 et al., 2012). The palaeobatrachids appear to have never reached the east of the Ural
1153 Mountains. Their most eastern distribution is recorded in the Late Pleistocene locality of
1154 Apastovo, in Russia, which is about 600 km west from the Ural Mountains (*Wuttke et al.*, 2012).
1155 The Western Siberian record does not only represent the first and only out-of-Europe
1156 occurrence of the family, but, surprisingly, falls within the Late Miocene palaeobatrachid gap of
1157 the European record. It is possible that palaeobatrachids occupied Western Eurasia again at the
1158 Mio-Pliocene boundary, from the east.

1159

1160 **Bombinatoridae**

1161 The primitive family of aquatic toads Bombinatoridae includes two recent genera: *Bombina* and
1162 *Barbourula*. The family is known since the Maastrichtian, Late Cretaceous in Romania, genus
1163 *Hatzegobatrachus* (*Venczel et al.*, 2016) and the Early Eocene in India, genus *Eobarbourula*
1164 (*Folie et al.*, 2013). The Recent distribution of *Bombina* is confined to continental Europe and
1165 East Asia, representing the western and eastern genetic clades of the genus respectively. In
1166 Europe, two species *Bombina bombina* and *Bombina variegata* are known. *B. bombina* has the
1167 widest distribution and is found in Central to Eastern Europe, whereas *B. variegata* occurs in

1168 Central Europe and in the south-eastern and western parts of Eastern Europe (*Pabijan et al.*,
1169 2013). The fossil record of the fire-bellied toad *Bombina* is patchy and limited to the Neogene
1170 of continental Europe. According to *Sanchíz & Schleich* (1986), the oldest fossil occurrences of
1171 the genus (*Bombina* sp.) are known from two localities in Germany: Weißenburg 6 (early
1172 Aquitanian) and Stubersheim 3 (early Burdigalian) (*Sanchiz & Schleich*, 1986; *Böhme & Ilg*,
1173 2003). The personal observations of one of the co-authors of this study (MB) did not confirm
1174 the Weißenburg 6 record of *Bombina*. Therefore, in the present study, we consider
1175 Stubersheim 3 to be the earliest occurrence of the genus.

1176 Bombinatorids later appeared in Central Europe in the mid Middle Miocene (*Bombina* sp.,
1177 Opole 2, Poland) (*Młynarski et al.*, 1982). Later, fire-bellied toads are present in three localities,
1178 representing the middle Tortonian age, including also the first fossil occurrences of the Recent
1179 European species – *Bombina* sp. from Rudabánya in Hungary (9.9-10.30 Ma) (*Roček*, 2005), *B.*
1180 cf. *bombina* from Kohfidisch in Austria (8.55-8.95 Ma) (*Tempfer*, 2005), and *B. cf. variegata*
1181 from Suchomasty in Czech Republic (8.8-9.2 Ma) (*Hodrová*, 1987). During the Pliocene,
1182 bombinatorids are represented mainly by the species *B. bombina* in six localities within Central
1183 Europe (*Böhme & Ilg*, 2003). The Pleistocene record is the richest in bombinatorid specimens
1184 with record from over 15 localities ranging from Central to Eastern Europe, and in which both
1185 Recent European species, *B. variegata* and *B. bombina*, are documented (*Böhme & Ilg*, 2003)
1186 (Fig. 9, Table S5).

1187 In Western Siberia, bombinatorids are known from three localities: Malyi Kalkaman 2, Selety
1188 1A, and Cherlak. The oldest known record dated back to the late Serravalian (Middle Miocene).
1189 The oldest Messinian Selety 1A locality provided the fossil form of the Recent *B. bombina* (*B. cf.*

1190 *bombina*) (Fig. 9). The last record of the genus dates back to the early Messinian (Late
1191 Miocene). It is interesting to note that the Western Siberian record of the genus does not
1192 coincide with their European occurrences, i.e. they are present during those periods when
1193 *Bombina* is absent in Europe. According to our analysis, it is clear that the ancestor of the '*B.*
1194 *bombina* – *B. variegata*' clade was present in Europe from, at least, the later part of the Early
1195 Miocene. Later in the Middle Miocene they expanded into Western Asia, reaching the east from
1196 the Ural Mountains. The Western Siberian fossil *Bombina* can be clearly osteologically
1197 separated from *Bombina orientalis*, a member of the East Asian clade of the genus. Taking their
1198 recent distribution as well as the fossil record into account, a split of the European and Asian
1199 *Bombina* clades seems most probable in Asia during the Paleogene.

1200

1201 **Pelobatidae**

1202 The family of European spadefoot toads Pelobatidae includes only one extant genus with four
1203 species distributed in north-western Africa, Europe, in small areas that are east of the Ural
1204 Mountains in Russia and in the north of Kazakhstan (*Kuzmin*, 1995). The family has Laurasian
1205 affinities and records are known from the Late Cretaceous in North America. The presence of
1206 pelobatids in Europe dates back to the Early Eocene, as indicated by the fossil genus
1207 *Eopelobates* (Middle Eocene – late Pliocene), as well as by the fossil forms of the Recent genus
1208 *Pelobates* (Middle Oligocene – Recent) (*Roček et al.*, 2014). The Asian record of Pelobatidae is
1209 very limited and includes forms from the Eo-Oligocene of Kazakhstan (*Chkhikvadze*, 1985) and
1210 Eocene of India (*Folie et al.*, 2013). Recently, *Roček et al.* (2014) excluded the genus *Uldzinia*
1211 (Oligocene, Mongolia) (*Gubin*, 1995) from the family Pelobatidae. The Kazakhstan fossil record

1212 of the family (*Chkhikvadze*, 1985, 1998) includes numerous remains of Pelobatidae indet. from:
1213 (1) the localities of the Zaisan Basin from the Upper Aksyr Svita³, early Priabonian; rare finds in
1214 the Kusto Svita and basal horizon of Buran Svita⁴, late Priabonian and earliest Rupelian;
1215 abundant occurrences in the Buran Svita⁵, early Rupelian and (2) large-sized spadefoot toads
1216 from the Kyzyl-Kak locality of the Turgay Basin, Central Kazakhstan, Late Oligocene
1217 (*Chkhikvadze*, 1998). Revision of this rich pelobatid record from the Zaisan Basin was not
1218 possible due to the lack of descriptions and illustrations of the material as well as the difficulty
1219 in accessing the specimens. Nevertheless, taking the Paleogene fossil records into account, we
1220 inferred that the spadefoot toads may have dispersed from Europe to Western Asia during the
1221 Late Eocene to Early Oligocene. It cannot be ascertained if the *Pelobates* sp. from the Selety 1A
1222 (early Messinian, Miocene) is a European or Asian migrant.

1223

1224 **Hylidae**

1225 The family of tree frogs, Hylidae, has a wide distribution in Eurasia and is represented by the
1226 monophyletic genus *Hyla*. The most recent phylogenetic study of the genus *Hyla* by *Li et al.*
1227 (2015) recognised two closely related clades in Eurasia, namely the West Palaeartic *arborea*-
1228 group and East Palaeartic *chinensis*-group, as well as a small East Palaeartic *japonica*-group
1229 that is related to the North American clade of *Hyla*. The revision of the Western Eurasian *Hyla*

³ localities: Zertsalo [Sunduk Section], lager Biryukova [Kiin-Kerish Section], lower faunistic level of Plesh [Kusto-Kyzylkain Section], probably also Tabtym [Sarykamysh Section]

⁴ localities: main level of Plesh, Tuzkabak, Cherepakhovoe Pole [Tayzhuzgen Section], Raskop [Aksyr Section], Tyubiteika, sopki 'Rybnaya' and Kontrolnaya [Juvan-Kara Section]

⁵ localities: Maylibay, Tologay [Tayzhuzgen Section], Podorozhnik [Jaman-Kara Section]

1230 phylogeny, based on molecular genetic studies, revealed a high diversity in the area containing
1231 about eight (?nine) (*Li et al.*, 2015) or ten (*Gvoždík et al.*, 2010) species. Among them are two
1232 clades: (1) *H. savignyi* in the east (Levant and the area of Turkey, Iran, Armenia, Georgia) and (2)
1233 *H. arborea* (Western, Central Europe and Balkan) + *H. orientalis* (South-eastern Europe,
1234 Georgia, Armenia, Iran), which have wide distributions in the east and west respectively (*Stöck*
1235 *et al.*, 2008a; *Gvoždík et al.*, 2010).

1236 The oldest European record of the genus is known from the Oberdorf O4 locality, late Early
1237 Miocene, Austria (*Sanchíz*, 1998b). After an interruption/gap of approximately three million
1238 years, records of the genus continued in the late Langhian with the first fossil appearance of the
1239 Recent species *H. arborea* (*H. cf. arborea*, Mátraszőlős 2, Hungary) (*Venczel*, 2004). The record
1240 is almost consistent in the entire Neogene and Quaternary periods of Europe (Fig. 9). There is
1241 quite an abundant record of the genus with the oldest and first occurrences of *H. savignyi* (*H.*
1242 *cf. savignyi*) derived from five localities in Western Siberia, dating back to the late Late Miocene
1243 and early Early Pliocene. Apart from the distribution in Siberia, *H. savignyi* also may occur in
1244 southern Russia, in the Middle Pleistocene (*Ratnikov*, 2002)(see 'Comparison and Discussion' in
1245 *Hyla gr. H. savignyi*), representing the youngest fossil record of the species.

1246 Based on the fossil record of the tree frogs, we concluded that two large Western Eurasian
1247 clades split in Europe during the Middle Miocene. Our data indicated older ages for the first
1248 fossil occurrences of these clades than has been previously estimated from molecular data in
1249 two recent studies (*Gvoždík et al.*, 2010; *Li et al.*, 2015). *Gvoždík et al.* (2010)⁶ suggested that

⁶ The divergence dates of split events were estimated by a relaxed molecular clock approach, based on the mitochondrial data set, where the calibration with fossil record is missing.

1250 the split of *H. orientalis/arborea* and *H. savignyi* occurred 11.1 Ma (early Late Miocene, early
1251 Tortonian), which is approximately three million years younger than the first fossil occurrence
1252 of *H. cf. arborea* (Table S5). Whereas, without calibrating the molecular clock using the oldest
1253 European fossil *Hyla* (*Hyla* sp., Oberdorf O4 locality in Austria), *Li et al. (2015)* estimated this
1254 split to have occurred at 12-20 Ma, during a time interval in which the oldest fossil tree frogs
1255 related to the Recent *H. arborea* occurred. In both of the cases, the interpretation of the
1256 molecular phylogeny of the group can be improved by calibrating the phylogenetic tree with
1257 the fossil record introduced in this study.

1258 Considering our data and the results presented by *Li et al. (2015)*, we suggest the following
1259 distribution pattern for the West Palaearctic *H. arborea*-group: (1) the group entered Eurasia
1260 from the east via Beringia from North America, during the Paleogene; and (2) the ancestors of
1261 the group reached Europe during the early Miocene via the Tugai Strait between Europe and
1262 Asia (the Turgai Strait) and diversified, apparently, in Western Siberian. The Late Miocene and
1263 Early Pliocene records represent the most eastern expansion of the European genera, when the
1264 climatic conditions were still favourable for their distribution; it is conceivable for us that the *H.*
1265 *savignyi* may have potentially so far not found fossil occurrences in the Miocene of Eastern
1266 Europe and/or from the Caucasus in the south.

1267

1268 **Bufonidae**

1269 Two groups of toads were found in the studied localities in Western Siberian; namely the
1270 common (*Bufo bufo*) and the green toads (*Bufo cf. viridis*) (Fig. 7F-7K, Table S1). The toads of

1271 both groups are, with their occurrences are the most abundant among frogs remains found at
1272 the fossil localities.

1273 **Common toads.** *Bufo bufo* is the Recent species with the widest distribution (i.e. Central,
1274 Southern, Eastern Europe and Western and Eastern Asia) of all members of the common toads
1275 *Bufo bufo* species group. This group includes three other species with limited distribution,
1276 namely: *Bufo spinosus* (northern Africa, Western Europe), *Bufo eichwaldi* (south coast of the
1277 Caspian Sea), and *Bufo verrucosissimus* (east of the Black Sea) (Arntzen et al., 2013). These
1278 species are known also as the western group of the genus. Their nearby Eastern Asian relatives
1279 - the eastern group, include the *Bufo gargarizans* species group. The Western Siberian fossil
1280 record of the *Bufo bufo* species group is restricted to the late Late Miocene to the early Early
1281 Pliocene, which in comparison to the European record, is very poorly represented. The oldest
1282 toad remains that are assigned to the *Bufo bufo* species group are from the Middle Miocene of
1283 Slovakia: *B. bufo* from the Devinská Nová Ves – Zapfe’s fissure locality, 13.7-14 Ma (Hodrová,
1284 1980; Böhme, 2003) and *B. cf. bufo* from the Devinská Nová Ves – Bonanza locality, 13.5-13.7
1285 Ma (Hodrová, 1988). Then, since 9.2 Ma during the Late Miocene (Suchomasty locality in the
1286 Czech Republic) (Hodrová, 1987), *B. bufo* representatives are present in Central Europe and
1287 extend their distribution across Europe. At ca. 4.7 Ma, remains of the common toad, exhibiting
1288 characters of the Recent *B. spinosus*, appeared in Spain, in the Celadas 6 locality (Böhme & Ilg,
1289 2003). The oldest fossil remains referred to *B. verrucosissimus* were recovered from the Late
1290 Pliocene (3.0-3.8 Ma) in the Apastovo locality in Russia (Ratnikov, 2001). The Western Siberian
1291 record suggests at least a late Miocene dispersal of *B. bufo* to the east, reaching the present
1292 distribution area of the species. Considering the genomic data of Recuero et al. (2012), these

1293 'migrants' should represent the common ancestor of the *B. bufo* + *B. verrucosissimus* clade,
1294 expanding to the east into Asia and to the south into Eastern Europe. This bufonids most
1295 probably remained, permanently, in these areas, until present times. The lack of their
1296 representation in the fossil record in the Late Pliocene and Quaternary sites can be explained
1297 by sampling bias. Although *B. bufo* and *B. verrucosissimus* do not occur sympatrically nowadays,
1298 specimens of both these species have been found together in two Middle Pleistocene localities
1299 (Koziy Ovrage and Yablonovets from Russia; see more in Table S5).

1300 Two recent molecular studies (*Garcia-Porta et al., 2012; Recuero et al., 2012; pp. 71-86*)
1301 suggested models of palaeobiogeographic history and timing of major cladogenetic events in
1302 the *B. bufo* species group; e.g. the origin in South-western Asia and subsequent migration into
1303 Europe via Anatolia. These studies, however, did not consider the entire fossil record, including
1304 the oldest record of the groups from the Middle Miocene of Slovakia (*Hodrova, 1980*) nor those
1305 of the species group in both their calibration of the molecular clock and palaeogeographic
1306 considerations. The updating and improvement of the distribution models are, therefore,
1307 necessary. Moreover, further finds of the fossil forms of south-eastern species *B. eichwaldi* will
1308 help to reveal the place of origin and distribution routes of the ancestors of the group. Although
1309 only the molecular clock, and not the entire fossil record of the group has been used for the
1310 calibration, results from mtDNA sequencing seem to provide reliable data on diversification
1311 rates within the *B. bufo* species group, which can be confirmed by first appearances of the
1312 fossils related to each Recent species.

1313 **Green toads.** The range of the widely distributed *Bufotes viridis* species group (or *Bufotes viridis*
1314 *sensu lato*) extends across Central Europe to Central Asia, as well as the entire northern Africa

1315 and Mediterranean area, including numerous islands. The species complex is highly diverse and
1316 includes over ten recognised species, e.g. *Bufo balearicus* (southern Mediterranean and
1317 Apennine Peninsula, Corsica, Sardinia, Balearic Islands), *Bufo boulengeri* (northern Africa),
1318 *Bufo siculus* (Sicily), *B. viridis* (Central and Eastern Europe), *Bufo variabilis* (Balkans,
1319 Anatolia, Caucasus) etc., found in a diverse range of environments (Stöck et al., 2006; Stöck et
1320 al., 2008b). Among them, however, no valuable osteological characters has been established for
1321 taxonomic identification (Blain, Gibert & Ferràndez-Cañadell, 2010). Hence, no precise specific
1322 assignment of any fossil material is possible. Blain, Gibert & Ferràndez-Cañadell (2010) recently
1323 showed that the green toads were also present in the Iberian Peninsula in the Early Pleistocene,
1324 1.1-1.3 Ma, and suggested that they became extinct due to climatic changes and/or
1325 competition.

1326 In the studied Western Siberian localities, fossil remains that are related to *B. viridis* are the
1327 most frequently occurring element in the Western Siberian herpetofauna. This species is almost
1328 permanently present from the Middle Miocene to Early Pleistocene. Specimens are found in
1329 the late Middle Miocene localities, and although there are gaps in the record, remains are
1330 present in the late Late Miocene to Early Pleistocene localities (Table S1). In the youngest
1331 localities (Olkhovka 1A, Lebiazhie 1A, Lebiazhie 1B), they are found as a sole taxon. Further
1332 fossils assigned to the family Bufonidae (Bufonidae indet.) were already reported from the
1333 Kentyubek locality in the Turgay Basin, from the Middle Miocene (Bendukidze & Chkhikvadze,
1334 1976), and from two localities in the Zaisan Basin: the Zmei Gorynych locality in Akzhar Svita,
1335 from the Early Miocene (Chkhikvadze, 1985) and from the early Rupelian age fossil sites (see
1336 section 'Pelobatidae') of the Buran Svita (Chkhikvadze, 1998). Malakhov (2005) described the

1337 stratigraphically oldest green toad fossil, *B. aff. viridis*, from the early Early Miocene (20.4-22.5
1338 Ma, Aquitanian) locality of Ayakoz in North-eastern Kazakhstan (Fig. 1, Table S1). *B. aff. viridis*
1339 from the Ayakoz locality is older than the *B. aff. viridis* from the Early Miocene Keseköy locality
1340 (18-20 Ma) in north-western Turkey (Claessens, Leon P. A. M., 1997). All the occurrences of the
1341 oldest European fossils of green toads are from the Early Miocene: Vieux-Collonges locality in
1342 France (14-17 Ma), (Bailon & Hossini, 1990); Petersbuch 2 and 7 (17.5-18 Ma) localities in
1343 Germany (Böhme & Ilg, 2003); and probably the Córcoles locality (17-18 Ma) in Spain (Sanchíz,
1344 1998a). Once the green toads entered Europe, they became a regular element of the European
1345 Neogene and Quaternary herpetofaunal assemblages (Fig. 9). Besides *B. aff. viridis*, the
1346 European record of green toads includes another species, *Bufoetes priscus*, from four localities of
1347 the latest Early Miocene to the earliest Late Miocene age (see Table S5). Taking into account
1348 the *B. viridis* Neogene records and the bufonid records from the Eurasian Paleogene, we
1349 suggest that the group arrived in the Old World in the Paleocene (Rage, 2003), entered Central
1350 Asia in the Early Oligocene and diversified there. Although we were not able to study the
1351 Paleogene bufonid record from Kazakhstan, taking into consideration the palaeogeography of
1352 common and green frogs, the assignment of the Early Oligocene Kazakhstan record to the green
1353 toad seems most probable. Apparently, the Early Oligocene forms were ancestral to the *B.*
1354 *viridis* lineage, which evolved in Central Asia in the Early Miocene. This assumption is also
1355 supported by molecular data suggesting that: (1) the green toad clade underwent
1356 diversification in Asia during the Oligocene/early Miocene; and (2) a high genomic and specific
1357 diversity is found within the Central Asian green toads (Stöck et al., 2006). Present in the
1358 Central Asian fossil record from the Early Miocene; they consequently dispersed via Anatolia in

1359 the early Burdigalian into Europe during the middle Burdigalian. Apparently, the European
1360 Neogene record should not necessarily represent one 'lineage' or one dispersal event of the *B.*
1361 *viridis* group from Asia. Several migration events most probably took place during the Miocene.
1362 The descendants of these events were replaced later by the ancestors of the Recent species *B.*
1363 *viridis*, *B. variabilis*, etc. as indicated by the genetic data at the Mio-Pliocene transition (Stöck et
1364 al., 2006). Prospective further studies could include: (1) the verification of dispersal events in
1365 the European fossil record, with help of an abundant and species-rich fossil material from
1366 stratigraphically well-dated localities; (2) the exploration the Miocene record of Anatolian and
1367 South-eastern Europe, as well as the Paleogene record of Asia; and (3) a challenging project of
1368 establishing the osteological characters that are important for the systematic identification of
1369 the members of the *B. viridis* species group.

1370

1371 **Ranidae**

1372 The family of true frogs, Ranidae, are present in the Western Siberian record by both green
1373 (*Pelophylax* sp.) and brown (*Rana* sp.) frogs. The green frogs appear more frequently in the
1374 record than the brown frogs. Both frog genera are common amphibians in the Recent
1375 herpetofauna of the area. Besides this record, further true frog finds (e.g. Ranidae indet.) are
1376 reported from the early Rupelian age fossil sites (see the list of the locality section
1377 'Pelobatidae') of the Buran Svita, Zaisan Basin. We were not able to revise their taxonomic
1378 validity due to lack of figured fossils and the inaccessibility of the material.

1379 **Green frogs.** The genus *Rana* includes 21 Recent species of aquatic frogs having a wide
1380 distribution ranging from northern Africa, Europe to Eastern Asia. Two genetically distinct

1381 clades, i.e. Western Palaearctic and the Far East, are recognised within the green frogs genus
1382 *Pelophylax* (LyMBERAKIS et al., 2007). The oldest green frog record from the Western Siberian
1383 (*Pelophylax* sp.) is dated back to the late Middle Miocene, coinciding stratigraphically with the
1384 Eastern Siberian record of the group (Middle Miocene, ca. 13 Ma, Tagay Section, Baikal Lake,
1385 Russia) (Daxner-Höck et al., 2013). Records of this group are present in the studied localities
1386 until the late Early Pliocene with long (during the Late Miocene) and short (during the Early
1387 Pliocene) gaps in the fossil record. Due to the fragmentary preservation of the studied bones as
1388 well as the lack of other informative elements of the skeleton (e.g. frontoparietals), any
1389 assignment to the Recent green frog species was impossible. Considering the present
1390 distribution of the two green frog clades, an affiliation of the Western Siberian fossil record to
1391 the Western Palaearctic clade is most probable.

1392 Despite the being only a few green frog records described in this study, these records still
1393 significantly expand the previously scarce and poorly known fossil history of the genus.
1394 Moreover, both of the Middle Miocene records from Western and Eastern Siberia represent the
1395 oldest records of the green frogs in the Asian continent. Although an Asiatic origin of the green
1396 frogs has been already assumed by several authors e.g. Sanchíz, Schleich & Esteban (1993),
1397 LyMBERAKIS et al. (2007), the earliest frog remains have been assigned to the *Pelophylax*
1398 *ridibundus* species group, which occurred in Europe in the early Oligocene (Möhren 13 locality,
1399 Germany) (Sanchíz, Schleich & Esteban, 1993). Its affiliation to a living species is impossible. In
1400 Europe, the fossil record of *Pelophylax* is continuous and is maintained through the Oligocene
1401 and entire Neogene (Table S5). Nevertheless, a well-documented Paleogene record of the
1402 group is not available from Asia and, therefore, any interpretations would not be confident. The

1403 only possible scenario, taking into account both the fossil record and genomic data, is that the
1404 Western Palaeartic green frogs split from their Far East sister clade during the Eocene; they
1405 diversified in the territory of Europe and/or Western Asia during the Oligocene; they dispersed
1406 back to the East in the middle Miocene; and eventually reached the territory of the Western
1407 Siberia.

1408 **Brown frogs.** The genus *Rana* (subgenus *Rana* sensu *Veith, Kosuch & Vences, (2003)* is
1409 comprised of more than 15 species that are distributed throughout Eurasia. Similar to green
1410 frogs, there are two known lineages from the brown frog species, namely: the Western and the
1411 Eastern Palaeartic lineages (*Veith, Kosuch & Vences, 2003*). Based on the osteological
1412 characters, the studied Western Siberian brown frog remains show a relation to the Western
1413 Asiatic lineage of the genus *Rana*, more precisely to the *Rana temporaria* species group (sensu
1414 *Veith, Kosuch & Vences, 2003*). Among the late Paleogene and Early Miocene fossil frogs
1415 (*Böhme & Ilg, 2003*), in which the generic identification is unclear (*Rana* vel *Pelophylax*), only
1416 the frog remains from the Early Miocene in Dietrichsberg, Germany (*Böhme, 2001*) have
1417 definitely been assigned to the brown frog *R. cf. temporaria*, representing the oldest known
1418 record of the group so far. As already suggested by *Böhme (2001)*, brown frogs migrated from
1419 their possible centre of origin in Western or Central Asia to Europe during the second half of
1420 the Early Miocene. This hypothesis is confirmed by the brown frog fossils from the Ayakoz
1421 locality in Kazakhstan, which dates back to the Aquitanian age and are stratigraphically older
1422 than the Dietrichsberg fossil frogs. The present-day biogeography and diversity of brown frogs,
1423 the presence of a distinct Eastern Palaeartic lineage in Eastern Asia as well as the Asian
1424 distribution of many European species provide further support for an Asiatic origin. Most likely,

1425 the dispersal route of the brown frogs was similar to that of the green toad (*Bufo cf. viridis*)
1426 whereby dispersal into Europe occurred via Anatolia, during the Early Miocene.

1427 It is interesting to note that the earliest brown frog from the studied Western Siberian localities
1428 (Malyi Kalkaman 2) shows osteological similarities with the Recent species *R. temporaria*,
1429 representing herewith the oldest fossil record of the species in the east.

1430 Previous molecular studies (Veith, Kosuch & Vences, 2003; Lymberakis et al., 2007), on both
1431 green and brown frogs, aimed to reconstruct their phylogenetic relationships, suggest models
1432 of biogeographic history as well as suggest when the splits between different genera, clades,
1433 species, etc. occurred. Such studies have provided contradictory results also for this group, e.g.
1434 the split of *Rana* and *Pelophylax* was at 9.32 Ma (Veith, Kosuch & Vences, 2003), whereas
1435 Lymberakis et al. (2007) estimated the split of the Western Palearctic and Far East lineages of
1436 *Pelophylax* to have occurred significantly earlier, i.e. 15 Ma before. Here neither geologic
1437 events nor the fossil records have been used consistently for the calibration of the molecular
1438 clock. Thus, the recalibrating of the timing for the splits with the new fossil finds provides a
1439 more reliable basis for phylogenetic reconstructions.

1440 For the better understanding of relationships between these groups, as well as to reveal more
1441 around the origin and palaeobiogeographic history of them, it would be interesting to review
1442 the specimens of the Paleocene frogs (Ranidae indet.) from the early Rupelian fossil sites (see
1443 section 'Pelobatidae') of the Buran Svita in the Zaisan Basin (Chkhikvadze, 1998). The
1444 incorporation of such a review, however, was not possible in the present study, due to the lack
1445 of figures of the fossils and the inaccessibility of the material.

1446

1447 **Gekkonidae**

1448 The family Gekkonidae is represented in the Western Siberian fossil record by the straight-
1449 fingered or even-fingered geckos, genus *Alsophylax*. They occur only in the Cherlak locality,
1450 dated back to the terminal Miocene, ca. 5.9 Ma. *Alsophylax* sp. is the most abundant element in
1451 the herpetofaunal assemblage of the Cherlak locality, with approximately 70% of the
1452 identifiable bone material belonging to this taxon. The genus *Alsophylax* is mainly distributed in
1453 Central Asia, partly occurring also in Mongolia and China. These geckos prefer habitats in arid
1454 and warm landscapes (Ananjeva et al., 2006). The appearance of these dry and warm adapted
1455 geckos in Western Siberia, which is four degrees north of their present occurrence, suggests a
1456 shift of the arid environment from the south to the north at the end of the Late Miocene (see
1457 below). It is interesting to note that out of the seven gecko genera, e.g. *Eublephareus*,
1458 *Mediadactylus*, *Terratoscincus* (Ananjeva et al., 2006) inhabiting Central Asia, only *Alsophylax*,
1459 which has the most northern distribution, occurs in the fossil record. Apparently, this genus is
1460 ecologically more adaptable in comparison to other genera, not only in the present, but
1461 probably also in the past.

1462

1463 **Lacertidae**

1464 Lacertid remains are the most frequent fossil bones among those of lizards occurring in
1465 Western Siberian localities. They are very rare in the Middle Miocene faunas, but occur more
1466 frequently in the Late Miocene, Pliocene, and Pleistocene localities. In the middle Late Miocene
1467 locality Pavlodar 1A (ca. 7.25 Ma), two taxa (*Lacerta* s.l. sp. 1 and sp. 2) occur sympatrically.
1468 *Eremias* sp. appears in the Western Siberian record in the Pliocene. This genus is widely

1469 distributed in the Central Asian steppes, inhabiting dry and warm habitats (*Ananjeva et al.*,
1470 2006).

1471

1472 **Emydidae**

1473 *Emydoidea* sp. is the only turtle identified from the studied fossil sites. The present-day
1474 distribution of the monotypic genus *Emydoidea* is restricted to the water bodies of the north-
1475 eastern territory of the USA. In Eurasia, fossil forms of this aquatic genus appear in the fossil
1476 record in Central Kazakhstan since the Middle Miocene (*Emydoidea tasbaka*, the Kentyubek
1477 locality in the Turgay Basin) (*Chkhikvadze*, 1989). Fossil forms have also been reported in
1478 Eastern Europe from the Late Miocene (*Emydoidea tarashchuki*, Krivoy Rog locality in Ukraine
1479 and Pantishara (8.7-9.2 Ma) in Georgia) (*Chkhikvadze*, 1980);(*Chkhikvadze*, 2003). The Siberian
1480 record indicates their occurrence in Asia also during the Late Miocene, which, interestingly, is
1481 located much further north than their Middle Miocene record from Kazakhstan. According to
1482 *Chkhikvadze* (2003), representatives may have also been present in Eastern Europe during the
1483 Pliocene. We avoid interpreting palaeobiogeography, stratigraphic distribution, etc. of this
1484 genus, since the available published material (e.g. *Chkhikvadze*, 1983, 1989), together with
1485 other extinct testudinoid taxa from Kazakhstan and Eastern Europe, is insufficiently described
1486 and poorly illustrated, requiring thorough revision. Nevertheless, we used the available
1487 published data on both freshwater turtles and terrestrial tortoises to attempt to interpret the
1488 record at the family level (Table 2). The turtle records from three well-explored regions in the
1489 studying area, i.e. Zaisan Basin, Turgay Basins and Western Siberia, are summarised in the Table
1490 2 . Throughout the entire Early Miocene in the Zaisan Basin, the turtle fauna is dominated by

1491 aquatic forms, i.e. out of eight taxa only two are tortoises (*Protestudo* spp.). The aquatic forms
1492 remained dominant in the Zaisan Basin during the Middle Miocene, the terrestrial family
1493 Testudinidae completely replaced the aquatic turtles (Emydidae, Trionychidae) in the end of the
1494 Middle Miocene and became the only family present in the younger deposits of the Late
1495 Miocene. Similar to the Zaisan Basin, the aquatic forms represent the Middle Miocene turtle
1496 fauna in two adjacent regions, the Turgay Basin in the west and Western Siberia in the north.
1497 Subsequently, in the beginning of the early Late Miocene, a testudinid appears in Western
1498 Siberia and is replaced by an emydid towards the end of the late Late Miocene and a chelydrid
1499 at the Mio-Pliocene transition. The absence of tortoises since the end of the Late Miocene in
1500 Western Siberia and the Plio-Pleistocene in the Zaisan Basin can be explained by a less
1501 favourable, probably colder (MAT <15°C, cold month temperature CMT <8°C) climate. Since the
1502 late Late Miocene, the emydid and chelydrid aquatic turtles are the only chelonids in Western
1503 Siberia. The presence of these chelonids not only indicate a humid environment with standing
1504 water-bodies but most probably also a cooler climate (for emydids: MAT>8°C, CM>-1.4 °C),
1505 since, in general, aquatic turtles can tolerate much colder conditions than tortoises, in that an
1506 aquatic environment acts as thermal buffer, consequently enabling aquatic turtles to populate
1507 higher poleward latitudes.

1508

1509 **Palaeobiogeographic considerations**

1510 By comparing the spatial and temporal patterns between European and Asian fossil records,
1511 including the first and last fossil occurrences, combined with an analysis of the available

1512 genomic data of the recent relatives of the fossil groups present in the studied material, certain
1513 palaeogeographic distribution patterns can be revealed along with new interpretations.
1514 Our analysis suggests a Western Asiatic origin for Hynobiidae, Proteidae, aff. *Tylototriton*,
1515 *Bufo viridis* species group and brown frogs, *Rana*. The green toads and brown frogs
1516 dispersed coincidentally in the earliest Miocene wherein, and at least for the *Bufo viridis*
1517 group, Anatolia was involved. Anatolia also played an important role in the distribution of the
1518 *Bufo bufo* species group; however, any age estimation of the event is not available. A
1519 salamander, showing affinities to the clade of the Recent East Asian genera *Tylototriton* +
1520 *Echinotriton*, is present in Western Siberia, most probably representing the forms similar to that
1521 of the early Oligocene (aff. *Tylototriton*) in Europe, a sister group of the recent clade. In order to
1522 resolve the affiliations of these fossils, further Paleogene material from both the Asia and
1523 European continents are necessary.
1524 An eastward dispersal from Europe into Western Asia can be observed over a period ranging
1525 from the Middle to Late Miocene, based on the current data available from both European and
1526 Asiatic records, for at least seven amphibian groups (family Palaeobatrachidae, genera
1527 *Chelotriton*, *Pelobates*, *Bombina* (i.e. *Bombina* (cf.) *bombina*), *Hyla* (i.e. *Hyla* cf. *savignyi*),
1528 *Pelophylax* ?, *Bufo bufo* species group). Besides the amphibians, some Western Siberian
1529 reptiles, such as the glass lizards and snakes from the Middle Miocene, show European
1530 affinities, resembling the Central European faunas (Vasilyan, Böhme & Klembara, 2016).
1531 The amphibian genera *Bombina*, *Hyla*, *Bufo*, *Rana*, and *Pelophylax* resemble a comparable
1532 palaeobiogeographic pattern: the molecular genetic data showed the presence of two clearly
1533 separable western and eastern clades (species groups) in each of these genera. In all cases, it

1534 was possible to morphologically attribute the Western Siberian fossil amphibians to the
1535 western clades or species of the clades. It is interesting to note that even though the first fossil
1536 occurrences of these genera have different stratigraphic ages, they are found exclusively in
1537 Europe (see Fig. 9, Table S5). To explain this common pattern, we hypothesise that the western
1538 and eastern clades had already split in the Paleogene, most probably in the western or central
1539 parts of Asia, and subsequently dispersed into Europe.

1540 The Western Siberian fossil *Mioproteus*, *Chelotriton*, *Bombina*, *Paleobatidae*, *Hyla*, *Bufo bufo*
1541 and *Rana temporaria* represent the most eastern records of those groups found in the Eurasian
1542 fossil record. In comparison to their present-day geography, the Western Eurasian species of
1543 the genera *Bombina* and *Hyla*, respectively, show wider distribution ranges during the Middle
1544 to Late Miocene, and Late Miocene to Early Pliocene. The palaeogeographic affinity of the
1545 earliest Messinian pelobatid (locality Selety 1A) is still unclear. Considering the geographic
1546 location of the fossil site, its relation to the Recent genus *Pelobates* seems most possible.

1547 In *Chkhikvadze* (1985), two lizards *Varanus* sp. and Agamidae indet. have been reported from
1548 three Miocene localities of the Zaisan Basin. Although the taxonomic assignment of the remains
1549 could not be verified in this study, we adopt the identifications for biogeographic and
1550 palaeoenvironmental interpretations. These lizards are currently widely distributed in Central
1551 Asia. *Varanus*, being a thermophilous reptile species, is restricted to the southern part of the
1552 region. Its presence in the early Late Miocene of the Zaisan Basin aids in characterisation of the
1553 climate of the Sarybulak Svita, in the beginning of the late Miocene, i.e. a probable mean
1554 annual temperature of not less than 14.8 °C (*Böhme*, 2003).

1555 In summary, Western Siberia (Central Asia) can be hypothesised as a centre of evolution and
1556 dispersal for several temperate Neogene herpetofaunal taxa, e.g. the genera *Salamandrella* and
1557 *Mioproteus*, the green toad *Bufo viridis* species group and brown frog *Rana*. The Neogene
1558 herpetofauna of Western Siberia and the adjacent areas has significant similarities with the
1559 European amphibian and reptile assemblages. The Western Palaearctic herpetofauna gradually
1560 entered the Siberian territory from Europe, between the Middle Miocene to Early Pliocene,
1561 strongly shaping the herpetofauna of Western Siberia and partially retaining the faunal
1562 elements of an Asiatic origin (e.g. Hynobiidae, Proteidae, and *Alsophylax*). The faunal diversity
1563 of the fossil record collapses significantly after the Early Pliocene. Only a few amphibians and
1564 reptiles, e.g. *Salamandrella*, *Bufo*, *Lacerta*, and *Vipera* are present in the Pliocene fossil
1565 record, being able to survive in the increasingly less favourable environments to form the main
1566 part of the present-day Western Siberian herpetofauna.

1567 The palaeobiogeographic analysis of the recent amphibian faunas of Western Asia (*Savage*;
1568 *Garcia-Porta et al., 2012*) hypothesised a progressive aridification of Central Asia linked with
1569 the global cooling trends during the Miocene, forcing amphibians to shift their distribution to
1570 the south.

1571

1572 **Palaeoclimatic implications**

1573 The Neogene climate evolution of Western Siberia has been previously reconstructed based on
1574 palynofloras, showing a progressive change in environmental conditions, i.e. in the climate and
1575 vegetation, during the Miocene (*Arkhipov et al., 2005*). Between the Early to Late Miocene, a
1576 warm and humid climate was replaced by a warm temperate climate in the Middle Miocene

1577 and a boreal-warm temperate climate in the Late Miocene. Towards the end of the Miocene, a
1578 drastic climatic shift took place resulting in semiarid and arid conditions. The Pliocene climate is
1579 predominated by frequent changes between semiarid forest-steppe/steppe and arid desert
1580 environments, however, from the Late Pliocene the environment changes into subarctic
1581 (*Arkhipov et al., 2005; fig. 46, p. 76*).

1582 At a lower temporal resolution, the testudinoid fossil records from the Zaisan Basin, the Turgay
1583 Basin, and Western Siberia confirm a general trend towards aridity in the Neogene (Data S4).
1584 Based on the environmental requirement (aquatic or terrestrial) of the testudinoids from the
1585 Zaisan Basin, we infer that the climate changed from humid to dry. We further infer that the
1586 Early and Middle Miocene was mostly humid (dominance of aquatic families), whereas the
1587 presence of exclusively terrestrial forms (tortoises) from the latest Middle Miocene to Late
1588 Miocene indicates dry and open habitats in the Zaisan Basin. Unfortunately, it is impossible to
1589 make any quantification of the palaeoprecipitation values based on these limited taxa and well-
1590 documented herpetofaunal assemblages are necessary from these deposits for further
1591 environmental reconstructions.

1592 To establish a better palaeoclimatic understanding, we estimated palaeoprecipitation values for
1593 12 data points (Table S4). These localities provided six and more amphibian and reptile taxa,
1594 applicable for the bioclimatic analysis (*Böhme et al., 2006*). Even so, our data do not be enable
1595 accurate reconstruction of the climate development over the Middle Miocene to earliest
1596 Pleistocene in Western Siberia. The climate development can, therefore, only be reconstructed
1597 and discussed for several short intervals. Nevertheless, our estimations rather show a dynamic
1598 climate development in the Neogene of Western Siberia, with larger precipitation amplitudes,

1599 ranging from 158 mm to over 1500 mm per year (Table S1, Fig. 10), than previously estimated
1600 using palynological data (*Arkhipov et al., 2005*). Apart from the fluctuating humidity factor, in
1601 general, the MAP was significantly above the present day values (reaching 550% of the present-
1602 day values) (Fig. 10). Only two localities are characterised by drier climates, the late Serravallian
1603 (ca. 12.1 Ma) and the late Messinian (5.9 Ma), exhibiting either present-day or below present-
1604 day levels.

1605

1606 **Reliability of precipitation estimates**

1607 The accuracy of precipitation estimates, based on bioclimatic analysis of herpetofauna,
1608 depends primarily on the taxon counts and the assumption of low (stochastic) taphonomic bias
1609 (*Böhme et al., 2006*). In Western Siberia, some of the documented localities were rich in aquatic
1610 herpetofauna, e.g. composed by freshwater turtles, giant salamanders, proteids, etc. , but small
1611 terrestrial forms (e.g. lizards and anguids) were absent, indicating a possible non-stochastic
1612 taphonomic bias (i.e. exclusion of elements of certain habitats). These localities will result in a
1613 bias in humidity estimates toward the wet end. Examples of such localities include Kentyubek
1614 and Novaya Stanitsa 1A, where the numeric results well exceed the MAP of 1600 mm, the
1615 upper limit to which the eco-physiologic index – humidity relation is calibrated (see details in
1616 *Böhme et al., 2006*). In these cases, we restrict our estimates to a limit of 1500 mm.

1617 **Aquitanian**

1618 For the Aquitanian age Ayakoz locality, we estimated a MAP value of 945 mm, representing
1619 more than three times higher rainfall in comparison to the recent times. Using the palynologic
1620 data, *Arkhipov et al. (2005)* estimated a humid climate with MAP 800 mm for the Abrosimov

1621 Svita (Aquitanian age) in Western Siberia. Besides this study and based on the data of fossil
1622 macroflora, *Bruch & Zhilin* (2007) estimated similar values of precipitation (935 to 1232 mm)
1623 for about 30 Aquitanian age localities, distributed from Western to Eastern Kazakhstan. Our
1624 reconstruction, therefore, appears to fit well within the historical precipitation estimates of the
1625 region.

1626 **Akzhar Svita**

1627 Towards the end of the late Early Miocene (Burdigalian), an elevated humidity in Western
1628 Siberia can be suggested based on the presence of the giant salamander in three localities of
1629 the Zaisan Basin (Tri Bogatyrya, Vympel, and Poltinik). As already suggested, their occurrence
1630 indicates a high rainfall for those time periods (MAP > 900 mm), as well as an increased basinal
1631 relief enabling the distribution and reproduction of this group in the lowland settings (*Böhme,*
1632 *Vasilyan & Winklhofer*, 2012). This period of the Akzhar Svita also corresponds to the folding
1633 and uplift of the Altai Mountains (*Zykin*, 2012; p. 394), from which the establishment of the
1634 higher basinal relief was possible.

1635 **Late Serravallian**

1636 In contrast to the already known climate development suggested by *Arkhipov et al.* (2005), our
1637 data suggest that there were strong humidity fluctuations during the late Middle Miocene (late
1638 Serravallian), with MAP values ranging between 282, 884 and 1108 mm (Fig. 10). The only
1639 botanical data of this time (*Bescheul* macroflora) point to a warm-temperate and humid (MAP
1640 ~700 mm) climate (*Arkhipov et al.*, 2005), which best compares to our Malyi Kalkaman 2 results
1641 (MAP 884 mm).

1642 **Novostanitsa Svita**

1643 Although the herpetofaunal assemblage for the early Messinian locality Novaya Stanitsa 1A is
1644 incomplete, a very high MAP value of at least 1500 mm can be estimated. The value indicates a
1645 significantly higher humidity than of Tortonian-Messinian boundary and late Messinian (see
1646 below). Our data are contrary to the palynologic results, which gave lower estimates (400-450
1647 mm; Arkhipov *et al.* 2005).

1648 **Rytov Svita**

1649 The Cherlak locality (5.9 Ma, Rytov Suite) is characterised by a rather dry climate (MAP 255
1650 mm), with a similar humidity level to that of the present-day (Fig. 10). Our data for a warm and
1651 dry climate are confirmed by the presence of: (1) gekkonid *Alsophylax*; (2) mollusc fauna
1652 containing thermophilous species; (3) the small mammal fauna, represented mainly by pikas,
1653 hamsters and jerboas, characteristic for open and dry habitats (Zykin, 2012); and (4) ostriches
1654 (*Struthiolithus* sp.) and camels (*Paracamelus* sp.) in this svita (Shpanskiy, 2008). Arkhipov *et al.*
1655 (2005) summarised the available palynological and vegetation data of the svita and reported
1656 the presence of a poor (due to the oxidation) spectra containing xerophyte plants (Asteraceae,
1657 Chenopodiaceae), characterising desert and steppe environments. Interestingly, his results
1658 proposed a northward shift of dry steppe and desert environments by 4° (to the latitude of
1659 56°), which concurs with our data, as is indicated by the presence of the steppe-dwelling
1660 gekkonid *Alsophylax* sp. (see the section 'Gekkonidae').

1661 **Miocene-Pliocene transition (Detskaya Zhelznaja Daroga)**

1662 Even though the precise taxonomic identification of the Western Siberian and Zaisan
1663 cryptobranchids, is unclear at the generic or species level, their occurrence indicates a high
1664 rainfall > 900 mm MAP (Böhme, Vasilyan & Winklhofer, 2012) during the Burdigalian age in the

1665 Zaisan Basin and the Miocene-Pliocene transition in Western Siberia. Besides the presence of
1666 Cryptobranchidae indet. from the locality Detskaya Zheleznaja Doroga, the co-occurrence of the
1667 aquatic chelonids *Chelydropsis kuznetsovi* and probable *Sakya* sp. (Gaiduchenko, 1984;
1668 Gaiduchenko & Chkhikvadze, 1985) confirms the presence of a high degree of precipitation at
1669 the Miocene-Pliocene boundary in Western Siberia.

1670 **Earliest Pliocene (Olkhovka 1A-1C)**

1671 Our earliest Pliocene humidity data are estimated based on the fauna from the localities
1672 Olkhovka 1A, 1B and 1C, for which no correlation data is available for regional svitas (see
1673 'Geology and Stratigraphy section'). Nevertheless, the results still indicate significant
1674 precipitation (MAP 575 mm), well above the present-day values for this region. These findings
1675 correspond well with the similar aged Speranovskaya palynoflora (Volkova, 1984), which
1676 indicates the presence of warm forests and forest-steppes with MAP estimates between 500-
1677 550 mm (Arkhipov et al., 2005).

1678

1679 **Conclusions**

1680 In summary, over 50 salamander, frog, lizard, snake and turtle taxa have been assigned to
1681 specimens from more than 40 Western Siberian localities that range in age from the Middle
1682 Miocene to the Pleistocene (Table S1). The late Middle Miocene localities have the most diverse
1683 faunas including all the main groups of the herpetofauna. According to our analysis, the fossil
1684 fauna contains taxa showing an Asian (Eastern Palaearctic) origin, such as Hynobiidae,
1685 Proteidae, *Bufotes viridis* species group and *Rana*, *Varanus*, and Agamidae. The main part of the
1686 herpetofaunal assemblage, including Palaeobatrachidae, Paleobatidae, the genera *Chelotriton*,

1687 *Bombina* (i.e. *Bombina* (cf.) *bombina*), *Hyla* (i.e. *Hyla* (cf.) *savignyi*), *Pelophylax* ?, *Bufo bufo*,
1688 *Ophisaurus* sp. (Vasilyan, Böhme & Klembara, 2016), has European (Western Palaearctic)
1689 affinities. The Western Siberian records of *Mioproteus*, *Chelotriton*, *Bombina*, Paleobatidae,
1690 *Hyla*, *Bufo bufo*, and *Rana temporaria* represent the most eastern occurrences of these groups
1691 in Eurasia. The earliest Miocene dispersal of the green toad, *Bufo viridis* species group into
1692 Europe from Asia via Anatolia, can be inferred. We suggest the same distribution pattern for
1693 brown frogs, *Rana*, too. In this scope, it will be important to perform future detailed studies on
1694 the Neogene record of the amphibian and reptile faunas in Anatolia and analyse them in a
1695 palaeobiogeographic context.

1696 According to our study, the precipitation development in Western Siberia shows high-
1697 amplitude changes during the studied intervals. Aside from the certain time periods, i.e. late
1698 Seravalian and late Messinian, the palaeorainfall in Western Siberia was estimated to be
1699 significantly higher than the present-day values. The best results on precipitation estimates that
1700 we were able to reconstruct, with reliable age constrain, were for the period from 6.6 to ~4.5
1701 Ma. These results indicate a humid climate during the early Messinian; a dry climate during the
1702 late Messinian; a very humid climate during the Miocene-Pliocene transition and a humid
1703 climate during the earliest Pliocene (Data S4, Fig. 10). The decreasing tendency of the
1704 herpetofaunal diversity towards the end of the Neogene and Quaternary could be attributed to
1705 the progressive global cooling and forced ice-sheet development in the Northern Hemisphere.

1706

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1718

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Figure 1(on next page)

Map of Eurasia (A) showing location of the Western Siberian studied fossil sites (B) (1-38, 58; black-thin outlined circles) as well as localities known from the literature (39-57; white - thick outlined circles).

1, Baikadam; 2, Malyi Kalkaman 2; 3, Malyi Kalkaman 1; 4, Shet-Irgyz 1; 5, Petropavlovsk 1; 6, Znamenka; 7, Pavlodar 1A; 8, Selety 1A; 9, Kedey; 10, Novaya Stanitsa 1A; 11, Borki 1A; 12, Lezhanka 2A; 13, Cherlak; 14, Pavlodar 1B; 15, Lezhanka 2B; 16, Olkhovka 1A; 17, Olkhovka 1B; 18, Olkhovka 1C; 19, Iskakovka 2A; 20, Isakovka 1A; 21, Peshniovo 3; 22, Isakovka 1B; 23, Kamyshlovo; 24, Beteke 1B; 25, Pavlodar 2B; 26, Pavlodar 3A; 27, Lezhanka 1; 28, Andreievka-Speransko; 29, Andreievka 1; 30, Livenka; 31, Beteke 1C; 32, Lebiazhie 1A; 33, Lebiazhie 1B; 34, Podpusk 1; 35, Beteke 2; 36, Beteke 4; 37, Kamen-na-Obi; 38, Razdole; 39, Akespe; 40, Ayakoz; 41, Golubye Peski; 42, Zmei Gorynych; 43, Vympel; 44, Poltinik; 45, Zaezd; 46, Tri Bogatyrja; 47, Kaymanovaja cherepakha; 48, Ryzhaya II; 49, Kentyubek; 50, Ashut; 51, Point "Y"; 52, Sarybulak Svita; 53, Kalmakpai Svita; 54, Karabastuz; 55, Kalmakpai; 56, Petropavlovsk 1/2; 57, Detskaya zheleznaja doroga; 58, Shet-Irgyz 2. Map data © 2016 Google and Map data © OpenStreetMap contributors, CC BY-SA.



Figure 2 (on next page)

Compiled stratigraphy of the Middle Miocene – Early Pleistocene studied localities in Western Siberia grouped in the svitas and relative to their geographic positions.

The localities without assignment into a certain svita are given in coloured frames according to age. In the right column the small mammalian biochronologic data (species or lineages) used for age estimations are given, the numbers accompanying the species refer the locality numbers. The arrows left from the small mammal taxa indicates its/their first appearance.

Abbreviations: klm – Kalkaman, ish – Ishim, pv – Pavlodar, kd – Kedey, nst – Novaya Stanitsa, rt – Rytov, is – Isakov, psh – Peshnev, krt – Krutogor, bt – Betekey, liv – Levetin, irt – Irtysh, kar – Karagash.

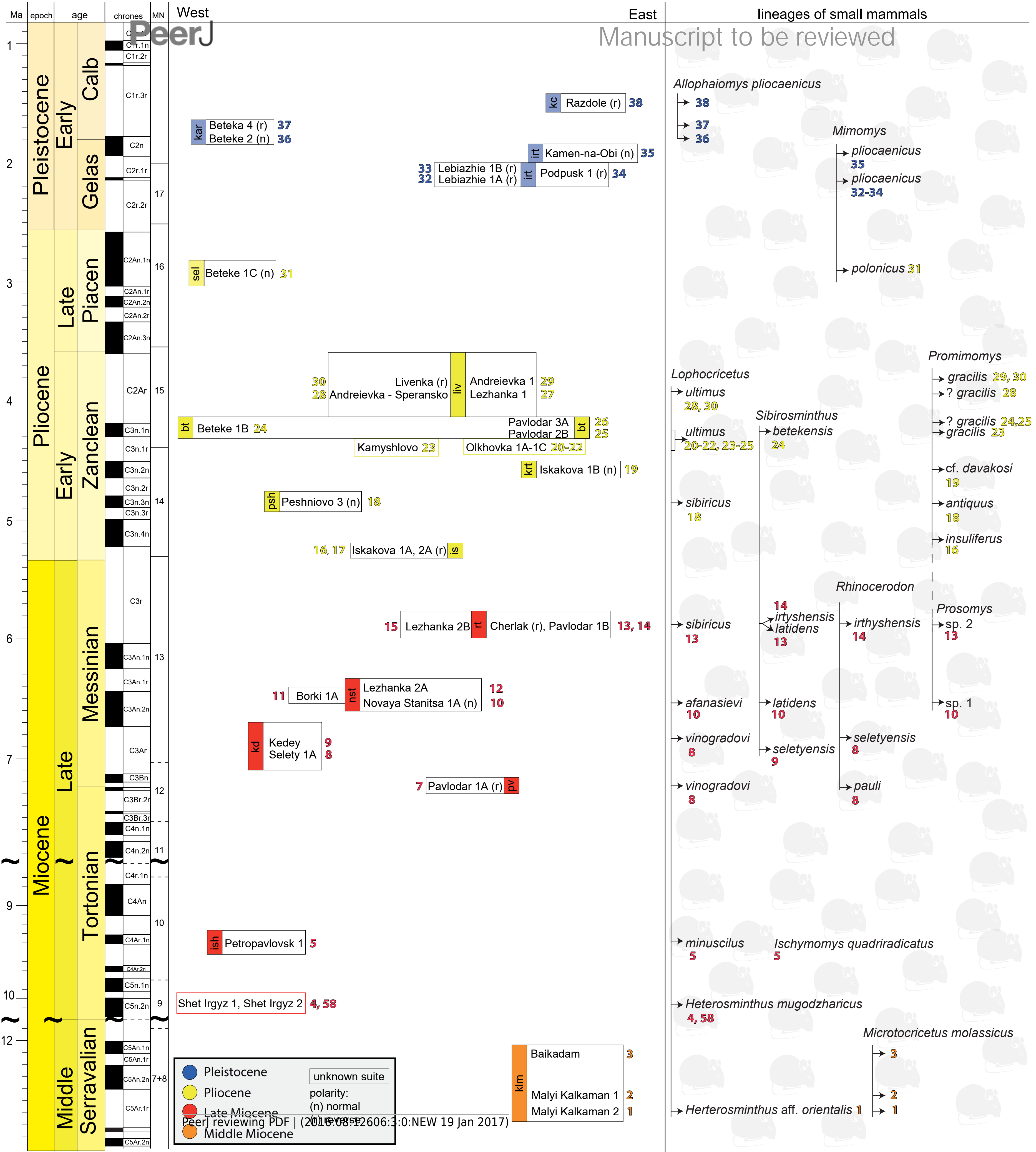


Figure 3

Salamander remains from Western Siberian localities.

A-C, Cryptobranchidae indet. from the loc. ? Gusiny Perelet, unnr. PIN specimens; **A**, fragmentary right dentary, natural cross-section; **B**, the same dentary, in lingual view; **C**, a jaw fragment, lingual view; **D-G**, *Salamandrella* sp., Lezhanka 2A, GIN 1130/1001-AM01, trunk vertebra; **H-S**, *Mioproteus* sp.; **H-L**, loc. Ayakoz, trunk vertebra, GNM unnr. specimen; **M-O**, trunk vertebra, Borki 1A, GIN 1115/1001-AM01; **P-Q**, right premaxilla, Malyi Kalkaman 2, GIN 1107/2001-AM01; **R-S**, left premaxilla loc. Grytsiv (Ukraine), unnr. MNMHK specimen; **T-X**, trunk vertebrae of aff. *Chelotriton* sp., loc. Ayakoz, GNM unnr. specimen; **Y**, *Chelotriton* sp. from Malyi Kalkaman 2, GNM unnr. specimen; **D, H, M, P, R, T, Y**, dorsal view; **E, I, N, Q, S, U**, ventral view; **F, J, O, V**, lateral view; **G, K, W**, anterior view; **L, X**, posterior view. Scale bars: A-C = 5 mm; D-G = 0.5 mm; H-Y = 1 mm.

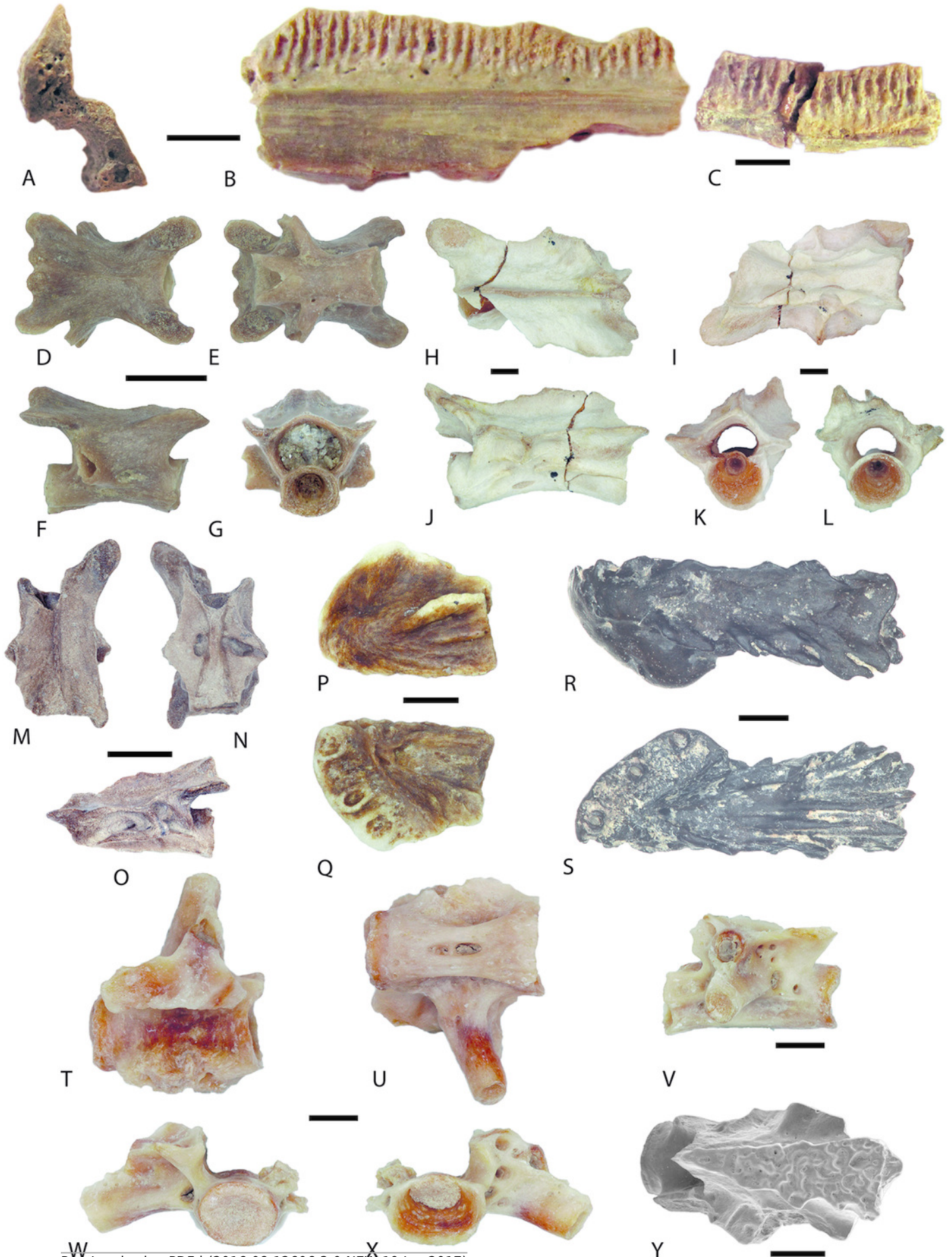


Figure 4

Trunk vertebrae of fossil aff. *Tylototriton* (A-K), and recent *Tylototriton*, *Echinotriton* and *Cynops* (L-AE).

A-E, aff. *Tylototriton* sp., locality Ayakoz, GNM unnr. specimen; **F-K**, GIN 950/2001-AM14 and GIN 950/2001-AM01, loc. Baikadam; **L-P**, *Tylototriton verrucosus*, GPIT unnr. specimen; **Q-U**, *Tylototriton shanjing*, GPIT unnr. specimens; **V-Z**, *Echinotriton andersoni*, GPIT unnr. specimen; **AA-AE**, *Cynops pyrrhogaster*, GPIT unnr. specimen; **A, F, G, L, Q, V, AA**, lateral view; **B, H, M, R, W, AB**, dorsal view; **C, I, N, S, X, AC**, ventral view; **D, J, O, T, Y, AD**, anterior view; **E, K, P, U, Z, AE**, posterior view. Scale bars = 2 mm.

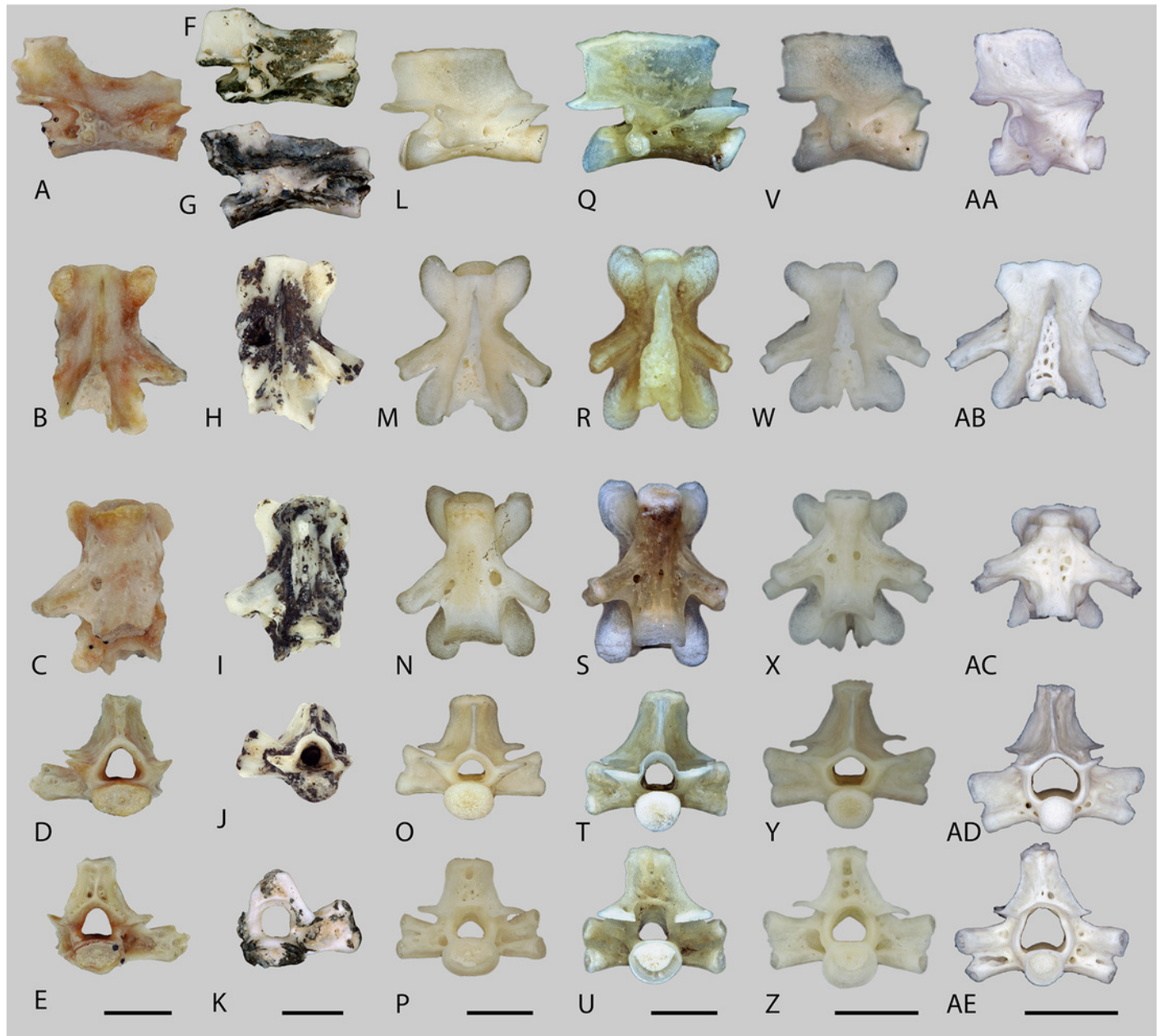


Figure 5

Palaeobatrachid sphenethmoids.

A-D, Palaeobatrachidae indet., Novaya Stanitsa 1A, GIN 948/2001-AM12; **E-H**, *Palaeobatrachus* sp. from Grytsiv (Ukraine), unnr. NMNHK specimen; **A, E**, ventral view; **B, F**, dorsal view; **C, G**, anterior view; **D, H**, lateral view. Abbreviations: ao, antrum olfactorium; alo, antrum pro lobo olfactorio; is, incisura semielliptical; ff, frontoparietal facet; lp, lateral processes; ls, lamina supraorbitalis; nf, nasal facet; onf, orbitonasal foramina; olf, olfactory foramina; pf, parasphenoid facet. Scale equals 1 mm

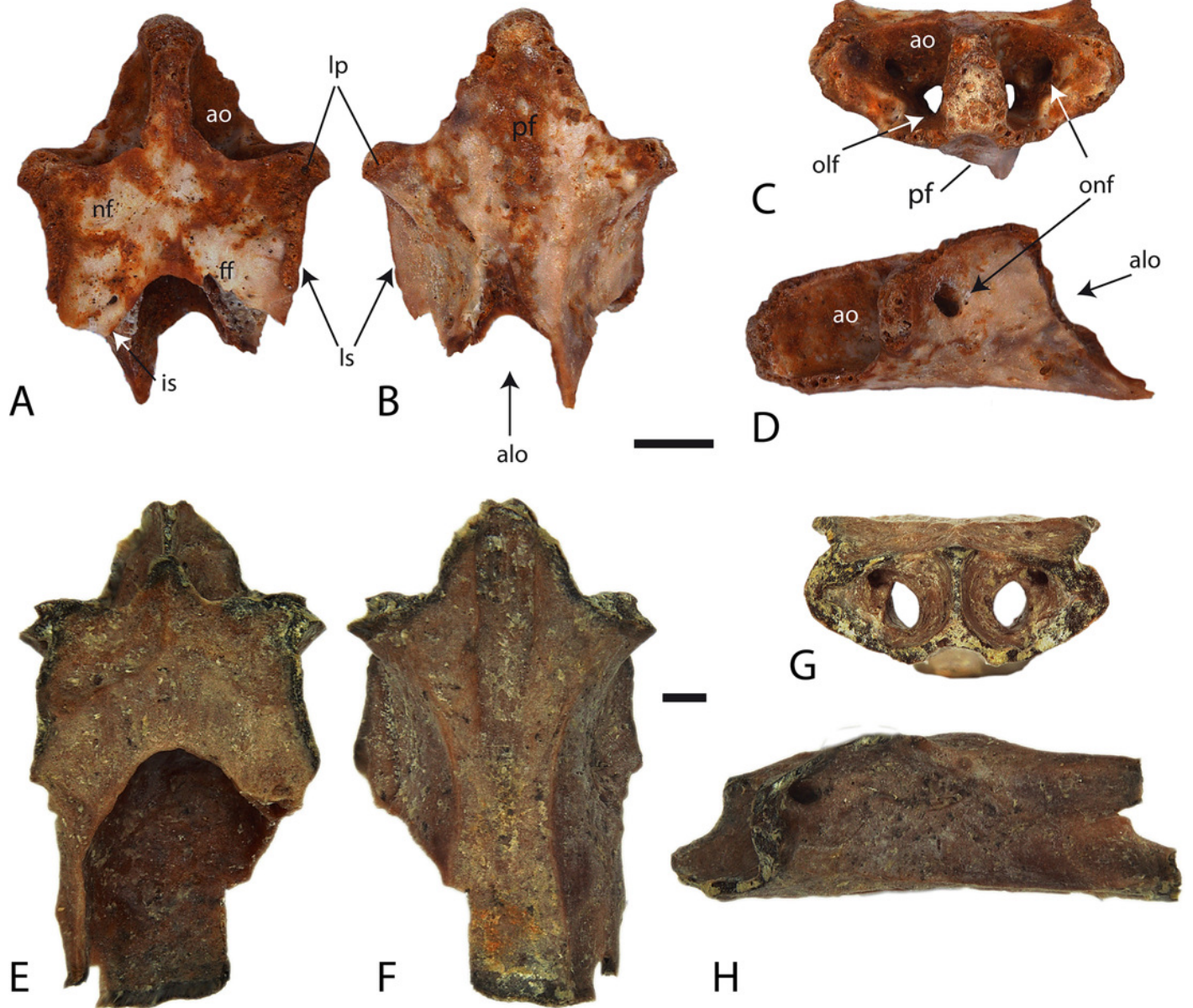


Figure 6

Fossil frogs from Western Siberia.

A-L, P-Q, X, AA, AB, A, AF, Iliia; **A-C** *Bombina* cf. *bombina*, Selety 1A, GIN 951/1001-AM06; **D-E**, *Bombina* sp., Cherlak, GIN 1110/2001-AM13; **G-I**, *Pelobates* sp., Selety 1A, GIN 951/1001-AM07; **J-L**, *Hyla* gr. *H. savignyi*, Lezhanka 2A, GIN 1130/1001-AM29; **P, Q**, *Bufo bufo*, Olkhovka 1B, GIN 11 11/2001-AM03; **X**, *Bufo* cf. *viridis*, Pavlodar 1A, GIN 640/5001-AM01; **AA, AB**, *Pelophylax* sp., Lezhanka 1, GIN 1129/1001-AM05; **AE-AF**, *Rana arvalis*, Malyi Kalkaman 1, GIN 1107/1001-AM10; **A, D, G, J, P, U, AA, AE**, in lateral view; **B, E, H, K, Q, AB**, in proximal view; **C, F, I, L**, in medial view; **M-O, R-T, Y, Z, AC, AD, AG, AH**, Scapulae of frogs; **M-O**, *Hyla* gr. *H. savignyi* from Lezhanka 2A, GIN 1130/1001-AM33; **R-T**, *Bufo bufo*, Olkhovka 1C, GIN 1111/3001-AM01; **Y-Z**, *Bufo* cf. *viridis*, Pavlodar 1A, GIN 640/5001-AM63; **AC-AD**, *Pelophylax* sp., Lezhanka 1, GIN 1129/1001-AM07; **AG-AH**, *Rana temporaria*, Malyi Kalkaman 1, GIN 1107/1001-AM01; **M, R, Y, AC, AG**, dorsal view; **N, S, Z, AD, AH**, ventral view; **O, T**, posterior view; **U, V**, trunk vertebra of *Bufo bufo*, Olkhovka 1C, GIN 1111/3001-AM02; **U**, anterior view; **V**, lateral view; **W**, urostyle of *Bufo bufo*, Olkhovka 1C, GIN 1111/3001-AM03, dorsal view. The arrows show the position of the angular fossa. Scale bars: A-Q, AA-AD, AG, AH = 1 mm, R-Z, AE, AF = 2 mm.

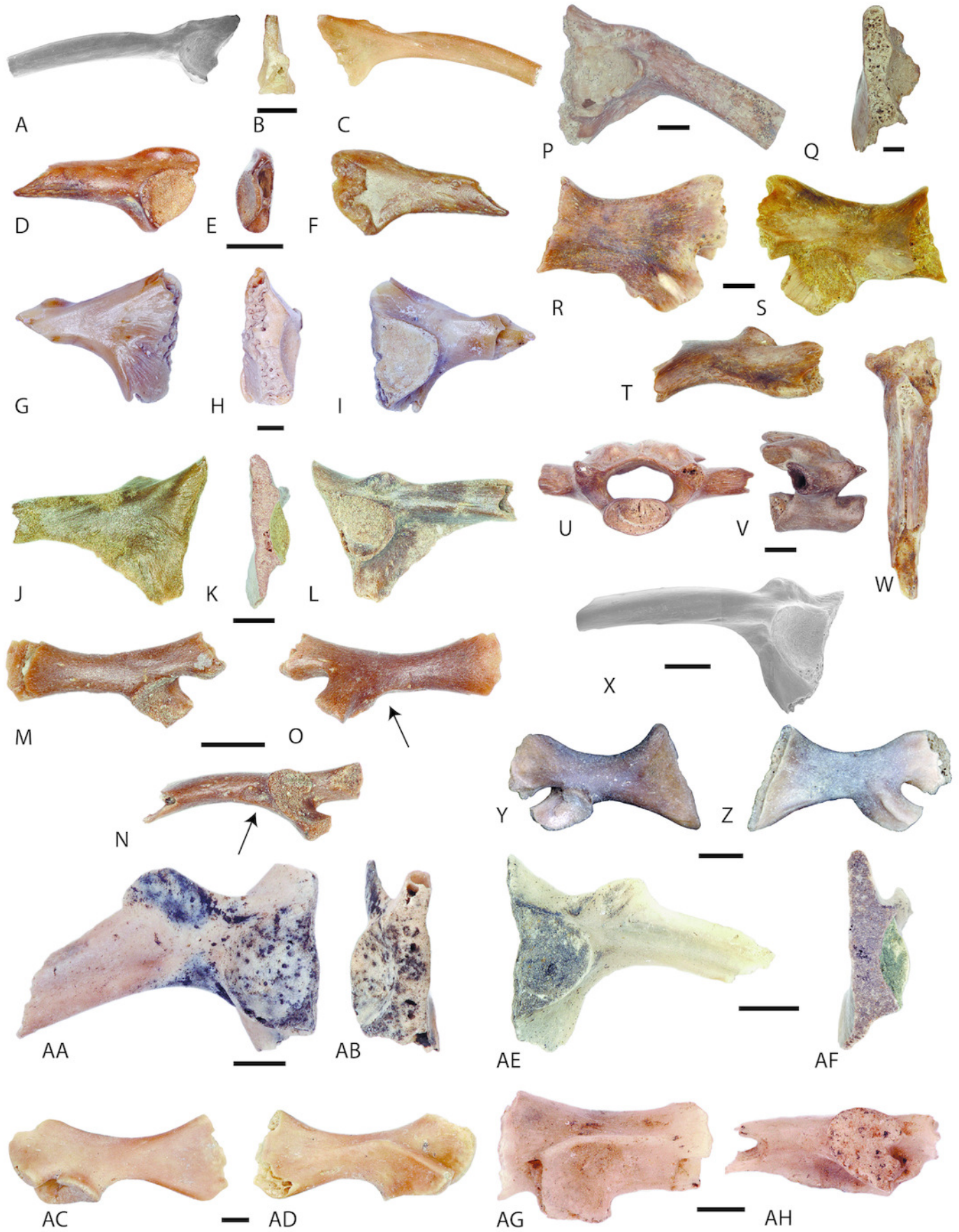


Figure 7

Alsophylax sp. from the localities Cherlak (A-P) and Mynsualmas-MSA 3 (Q).

A-E, two left dentaries; **A-D**, left dentary, GIN 1110/2001-RE11; **A**, mirrored labial view; **B-D**, mirrored lingual view; **C**, symphyseal region in lingual view; **D**, the same region in ventral view, both display the symphyseal groove; **E**, posterior fragment of left dentary, GIN 1110/2001-RE12, lingual view; **F-M**, five maxillae; **F-G**, left maxilla, GIN 1110/2001-RE26, lingual view; **H**, right maxilla, GIN 1110/2001-RE39, lingual view; **I-J**, right, GIN 1110/2001-RE40 and **K-L**, left maxillae GIN 1110/2001-RE27; **I, K**, lingual view; **J, L**, labial view; **M**, left maxilla, GIN 1110/2001-RE28, labial view; **N-P**, cervical vertebra, GIN 1110/2001-RE44; **N**, anterior view; **O**, left lateral view; **P**, posterior view; **Q**, right dentary, unnr. GPIT specimen, lingual view. Abbreviations: dl, dental lamina; ds, dental shelf; fcpr, facial process of maxilla; fMx5, foramina for mandibular division of the fifth cranial (trigeminal) nerve; hfr, haemal foramen; hl, horizontal lamella; lf, lacrimal facet; lg, longitudinal groove; lh, lamina horizontalis; mc, Meckelian canal; na, neural arch; nc, neural canal; nf, nasal facet; pfc, palatine facet; ph, paries horizontalis; prz, prezygapophysis; psz, postzygapophysis; pv, paries verticalis; pxp, premaxillary process; pyp, pterygapophysis; sac, opening of superior alveolar canal; sg, symphyseal groove; sf, **splenea** facet; tpr, transverse process.



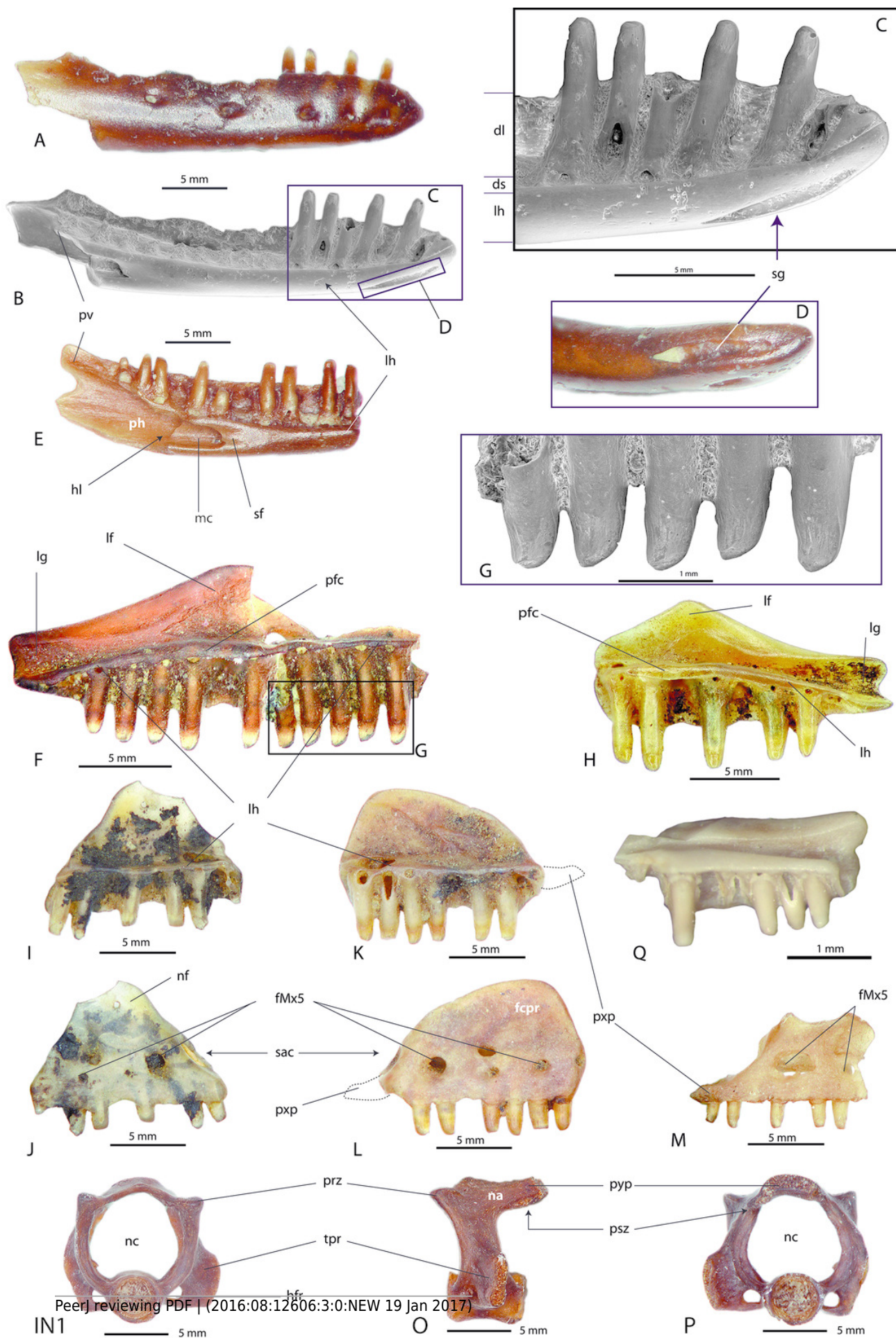


Figure 8

Lizard and turtle remains from the Western Siberian localities.

A, *Lacerta* s.l. sp. 1, left dentary, Pavlodar 1A, GIN 640/5001-RE01, lingual view; **B**, *Lacerta* s.l. sp. 2, right dentary, Pavlodar 1A, GIN 640/5001-RE34, lingual view; **C-D**, *Eremias* sp., frontal, Pavlodar 2B, GIN 1108/2001-RE01; **C**, dorsal view; **D**, ventral views; **E**, *Emydoidea* sp., fragment of right hypoplastron, GIN 948/2001-RE01, ventral view; **F-G**, *Emydoidea* sp., left femur, GIN 948/2001-RE02; **F** cranial view; **G**, ventral view. Scale bars: A, C, D = 2 mm; B = 1 mm; E-G = 1 cm.

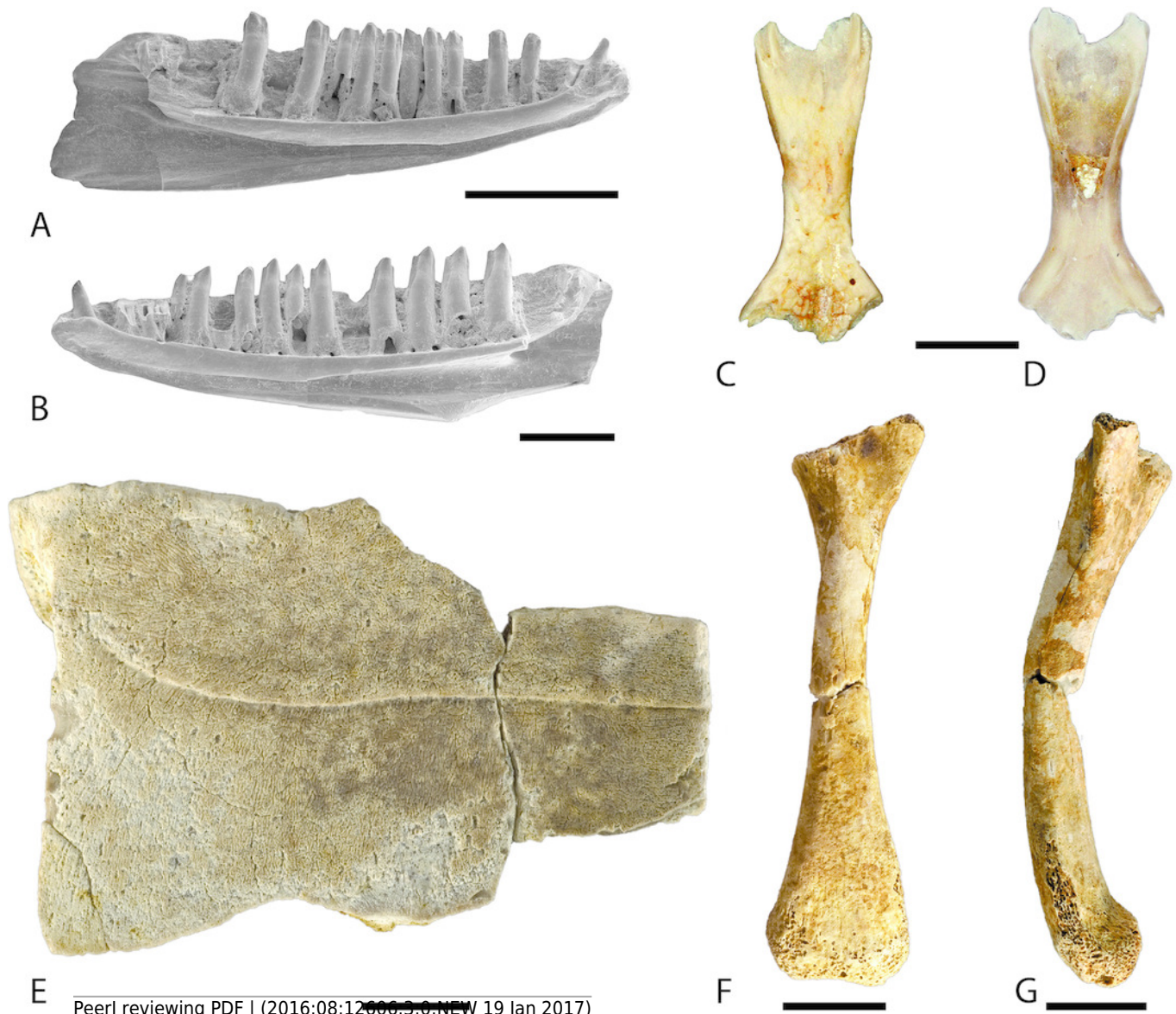


Figure 9 (on next page)

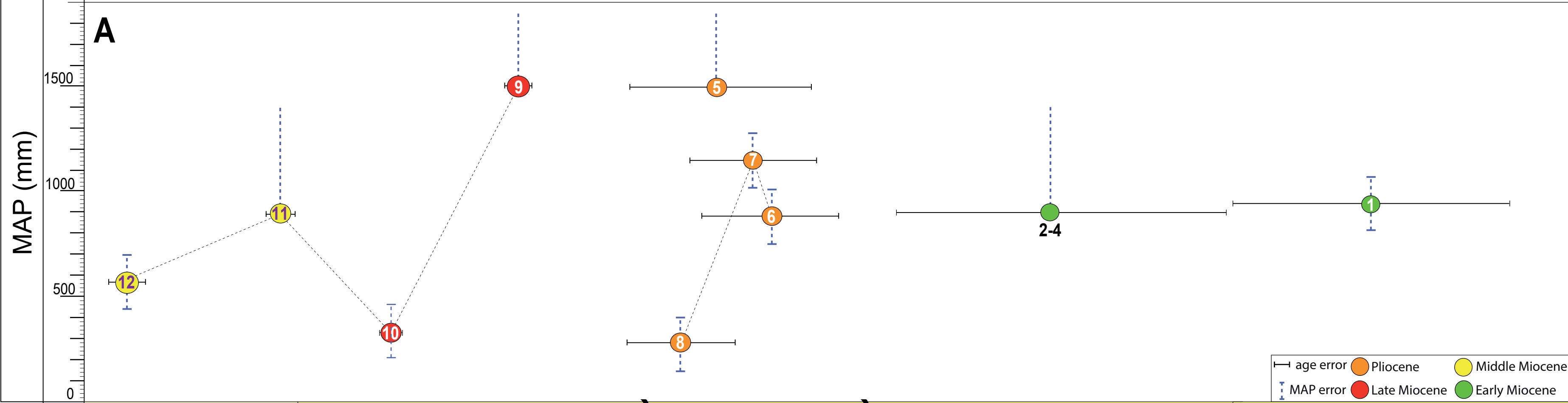
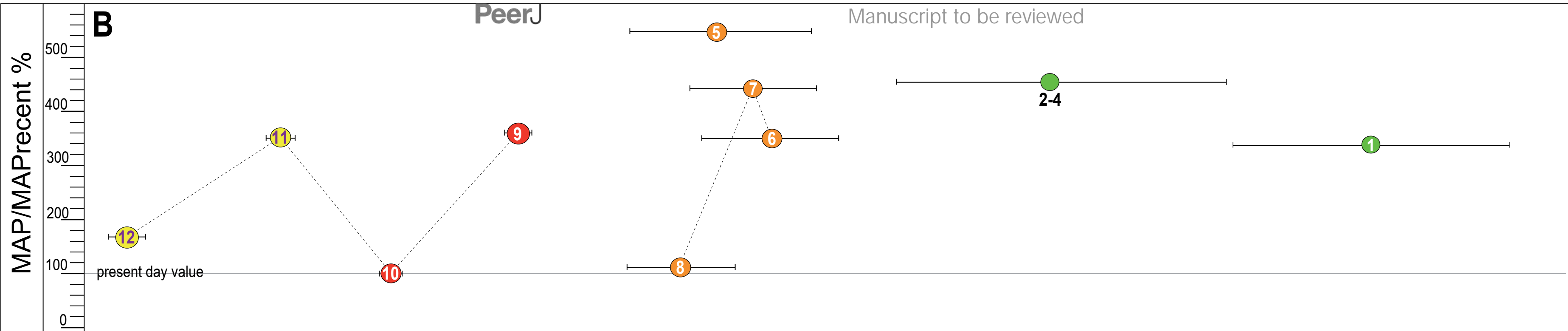
The European (Suppl. 3) and Western Siberian (present study) Neogene fossil record of the studied amphibian groups.

Detailed list of the localities see the Additional file 3 and for the family Cryptobranchidae – Böhme *et al.* (2012; table 1). The occurrences of each group in Europe and Western Siberia are given in the same colour. The Paleogene records of the groups are indicated by arrows. Abbreviations: Hyn, Hynobiidae; Cry, Cryptobranchidae; Prot, Proteidae; Chel, *Chelotriton*; Tylt, *Tylototriton*; Bomb, *Bombina*; red balk, *Bombina* (cf.) *variegata*; black balk, *Bombina* (cf.) *bombina*; Palbr, Palaeobatrachidae; Pelb, Pelobatidae; Hyla, *Hyla*; white balk, *Hyla* (cf.) *arborea*; Bbuf, *Bufo bufo* (group); Bvir, *Bufotes* (cf.) *viridis* / group of *Bufotes viridis*; Rana, *Rana* (cf.) *temporaria*; Pelx, *Pelophylax*.

Figure 10(on next page)

Palaeoprecipitation development of Western Siberia including the Zaisan Basin.

A, curve displaying the development of the absolute values of mean annual precipitation (MAP); B, the ratio of MAP to recent precipitation value ($MAP / MAP_{recent} 100\%$), dashed black line (100 %) indicates the recent precipitation values. Localities: 1, Ayakoz; 2, Vympel; 3, Poltinik; 4, Tri Bogatyrja; 5, Kentyubek; 6, Malyi Kalkaman 2; 7, Malyi Kalkaman 1; 8, Baikadam; 9, Novaya Stanitsa 1A; 10, Cherlak; 11, Detskaya zheleznaya doroga; 12, Olkhovka 1B.



← age error ● Pliocene ● Middle Miocene
 ↓ MAP error ● Late Miocene ● Early Miocene

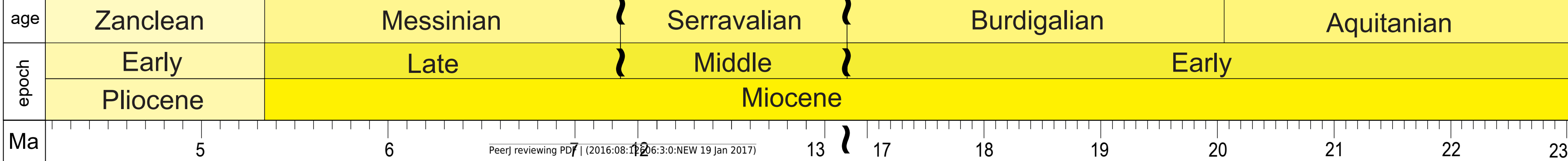


Table 1 (on next page)

Recent herpetofauna of south-western part of Siberia (Ob and Irtysh River drainages) according to different authors.

Recent herpetofauna of south-western part of Siberia (Ob and Irtysh River drainages) according to different authors. References: 1 - *Kuzmin* (1995) ; 2 - *Borkin* (1999) ; 3 - *Ravkin, Bogomolova & Chesnokova* (2010) ; 4 - *Amphibiaweb* (2016) ; 5 - *Ananjeva et al.* (2006).

Abbreviation: is - insular occurrence.

1

Taxa		References				
		1	2	3	4	5
Caudata	<i>Salamandrella keyserlingii</i>	+	+	+	+	
	<i>Lissotriton vulgaris</i>	is	+	-	+	
Anura	<i>Rana arvalis</i>	+	+	+	+	
	<i>Rana amurensis</i>	+	+	+	+	
	<i>Rana temporaria</i>	+	-	+	+	
	<i>Pelophylax ridibundus</i>	is	-	+	is	
	<i>Bufo viridis</i>	is	+	+	is	
	<i>Bufo variabilis</i>	-	-	-	+	
	<i>Bufo bufo</i>	+	+	+	+	
	<i>Bufo gargarizans</i>	-	?	-	is	
Lacertoidea	<i>Lacerta agilis</i>					+
	<i>Zootoca vivipara</i>					+
Serpentes	<i>Elaphe dione</i>					+
	<i>Natrix natrix</i>					+
	<i>Vipera berus</i>					+
	<i>Vipera renardi</i>					+
	<i>Gloydius halys</i>					+

2

3

Table 2(on next page)

Neogene testudinoid fauna of Western Siberia and the Zaisan and Turgay Basins.

According to *Chkhikvadze* (1989), as well as the superscriptions before the taxa indicate the references: * *Kordikova* (1994) ; ** *Danilov, Cherepanov & Vitek* (2013) , *** *Kentyubek fauna* (Suppl. 3), * *Chkhikvadze* (1989) and ** *Tleuberdina et al.* (1993) , ^k *Kuznetsov* (1982) , ^o our results. The aquatic families are indicated with blue and terrestrial families with dark yellow colour. Abbreviations: Ty - Trionychidae, Cy - Chelydridae, Pl - Platysternidae, Ts - Testudinidae, Ey - Emydidae, Gey - Geoemydidae.

1
2

Stage		Zaisan Basin			Turgay Basin	Western Siberia	
		Suite	Turtle "Stage"*-	taxa	taxa	taxa	
Pliocene						<i>Chelydropsis kuznetsovi</i> (Cy), ?Sakya sp. (Ey)	
Miocene	Late	Karabulak		* <i>Protestudo illiberalis</i> (Ts)		^o <i>Emydoidea</i> sp. (Ey)	
		Kalmakpai		<i>Protestudo kegenica</i> (Ts)		^k <i>Protestudo karabastusica</i> (Ts)	
	Middle	Sarybulak	up.	* <i>Protestudo darewskii</i> (Ts)	*** <i>Chrysemys</i> sp. (Ey), ?Ocadia sp. (Gey), <i>Emydoidea tasbaka</i> (Ey), <i>Kazakhemys zaisanensis</i> (Pl), ? <i>Chelydropsis</i> sp. (Cy)	** <i>Chrysemy</i> sp. (Ey), <i>Ocadia</i> sp. (Gey)	
			low.	* <i>Pelodiscus jakhimovitchae</i> (Ty)			
		Zaisan	up.	** <i>Baicalemys moschifera</i> (Ey)			
			low.	(**) <i>Baicalemys</i> sp. (Ey)			
	Early	Akzhar	up.	<i>Protestudo</i> sp. (Ts)			
			midd.		*- <i>Chelydropsis poena</i> (Cy)		
					* <i>Pelodiscus</i> sp. (Ty)		
					*- <i>Kazakhemys zaisanensis</i> (Pl)		
				** <i>Baicalemys jegalloi</i> (Ey)			
				** <i>Ocadia iliensis</i> (Gey)			
low.			* <i>Protestudo alba</i> (Ts)				
	Emydidae gen. indet. (Ey)						