



## THE EOMYIDAE IN ASIA: BIOGEOGRAPHY, DIVERSITY AND DISPERSALS

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**Abstract:** In Asia, the first find of an eomyid rodent was reported almost one century after the first studies of the family Eomyidae in North America and Europe. Since then, eomyid rodents have been increasingly found in Asia particularly over the past two decades. Here, we review the Asian record of this family at the genus level. Currently, 22 species within 14 genera were reported from Asia, including seven endemic genera and rare materials of apeomyine eomyids. Eomyids emphasize the palaeogeographic importance of Asia in considering intercontinental dispersal events of small mammals. With newly compiled data for Asian eomyids, we also compare genus-level diversity trends through time among North America, Europe, and Asia. Despite data standardizations limited with respect to potential biases in the fossil record, we found that the Asian eomyid diversity closely follows ecological shifts induced by climate changes. In general, Asian eomyid genera disappeared earlier than their European counterparts. We suggest that this pattern is not dictated by differences in the quality of the fossil record and is related to the expansion of drier habitats over large areas of Asia.

**Key words:** Rodentia, Eomyidae, palaeobiogeography, intercontinental dispersal, Valley of Lakes, Nei Mongol, Inner Mongolia, Junggar Basin, endemism

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## Introduction

The Eomyidae is an extinct family of small to medium-sized rodents, which are presumed to be phylogenetically close to geomyoid rodents (gophers, kangaroo rats) based on shared cranial characters (Wahlert 1978, Flynn 2008). Many of them are considered to have occupied similar niches to extant dormice, preferably inhabiting warm-temperate forest environments (Engesser 1999). However, eomyids probably occupied more ecological niches than dormice, as suggested by their more diverse dental morphology from bunodont to lophodont with increasing crown heights (Engesser 1999). Although most species of eomyid rodents are known based on isolated teeth only and a few mandible and skull fragments, two beautifully preserved specimens unearthed from the late Oligocene locality Enspel (Germany) evidence the great ecological disparity of this family. One of

them, *Eomys quercyi*, preserves the outline of a patagium supported by a cartilage originating from the elbow, much like that of scaly-tailed flying squirrels (Anomaluridae), revealing that this rodent was a glider (Storch et al. 1996). The other specimen preserves soft body parts of *Eomyodon volkeri*, showing that the species is a generalised terrestrial form (Engler and Martin 2015).

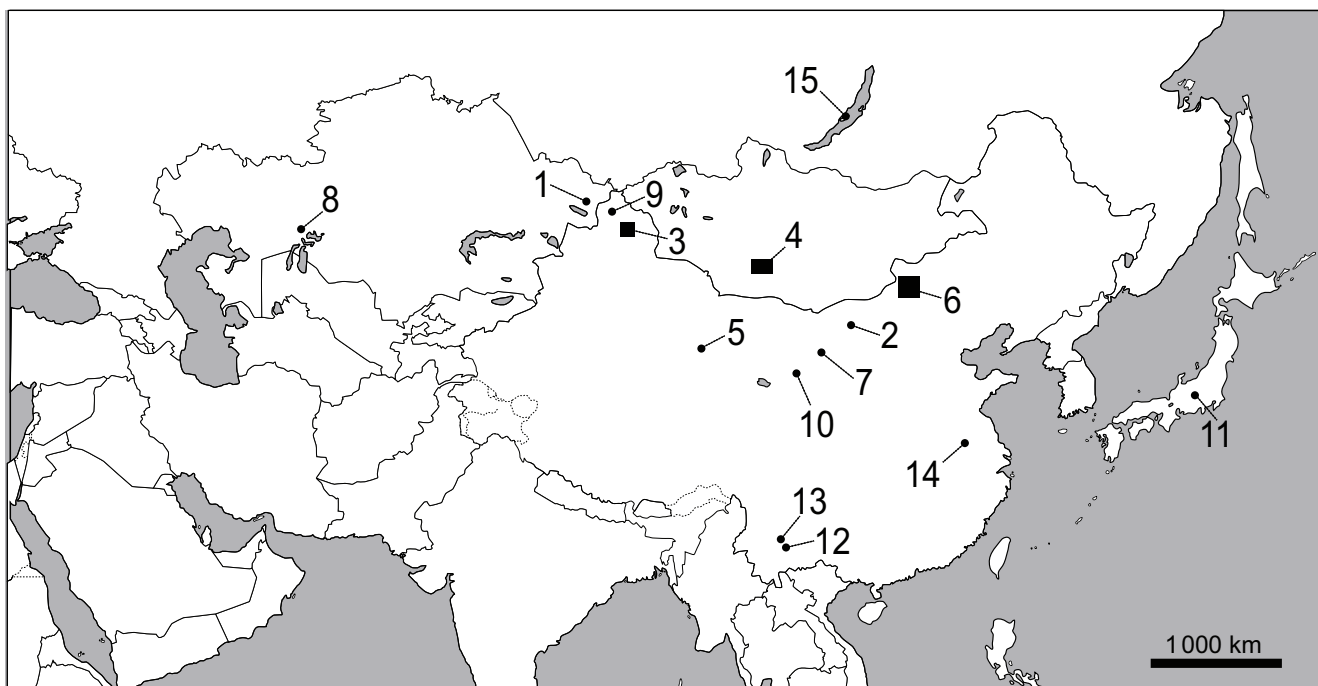
In Asia, the first reported eomyid fossil is a right jaw belonging to a new species of the American genus *Leptodontomys* from the late Miocene of North China (Zheng and Li 1982). Almost one century had passed since the first studies of the Eomyidae in North America and Europe (Cope 1884, Schlosser 1884). In the 1990s, a handful of eomyid fossils had been found in North China and Kazakhstan, including *Symplokeomys zaysanicus*, which marks the oldest occurrence (late Eocene) of the family in Asia (Emry et al. 1997), and Asian species of *Eomys*, *Pseudotheridomys*,

and *Eomyodon* (Wang and Emry 1991, Lopatin 2000, Wang 2002). Meanwhile, in Japan, a small eomyid fossil was discovered in the early Miocene deposits of Gifu and identified as *Pseudotheridomys* sp. (Tomida and Setoguchi 1994). In the 2000s, Wu et al. (2006) erected the Asian genus *Asianeomys* and transferred the following species to it: the Asian species of *Pseudotheridomys* (*P. asiaticus* from Inner Mongolia, China, *P. yanshini* from Kazakhstan) and *Eomyodon dangheensis* from Gansu Province, China. To date, six species of *Asianeomys* have been recorded from North China (Wu et al. 2006, Maridet et al. 2011, Gomes Rodrigues et al. 2014, Qiu and Li 2016), Mongolia (Maridet et al. 2015), and Kazakhstan (Lopatin 2000), with recent finds of the richest material in Inner Mongolia, China (Qiu and Li 2016) and Central Mongolia (Maridet et al. 2015). Most recently, the Japanese material ascribed to *Pseudotheridomys* sp. was transferred to a new genus, *Japaneomys*, closely related but basal to *Asianeomys* (Kimura et al. 2019), and *Omboomys* was erected for a small-sized bunodont eomyid lacking mesoloph(id) from Mongolia (Maridet et al. 2015). Likewise, apeomyine eomyids were recently discovered in Asia although the reported materials include only one tooth of *Apeomys* from Sihong, Jiangsu Province, central East China (Qiu 2017) and a single tooth of *Megapeomys* from Kani, Gifu, central Japan (Tomida 2011). Discoveries of eomyid rodents have been centered in mid-high latitudes, which have been more intensively studied, but they also occur in southern regions. Currently, up to three endemic genera

(*Heteroeomys*, *Plesieomys*, *Yuneomys*) were discovered from the hominoid-bearing sites in Lufeng, Yunnan Province, southwestern China (Qiu 2006, 2017). As summarised above, the fossil records in continental Asia and a locality in Japan have remarkably increased our understanding of the evolution and biogeography of the Eomyidae for the last two decades. Yet, no updated account has been published since the last review of Chinese eomyids by Qiu (1994). Here, we review the Asian record of eomyids at the genus level. For each genus, we provide a list of the included Asian species and main sites as well as a brief description of their dental characteristics including further considerations and remarks. Furthermore, we compiled taxonomic data for each species from a given locality based on original references and compared genus-level diversity trends of the family Eomyidae in North America, Europe, and Asia.

## Material and methods

The first and last occurrences of all eomyid rodents were compiled for North America, Europe, and Asia in order to explore diversity dynamics of the family in each of these continents (Tab. 1). For North American taxa, data were taken from the Paleobiology Database (PBDB, downloaded on July 3, 2017 from <https://paleobiodb.org/classic/displayDownloadGenerator>) and updated to include the recent new findings reported by Flynn (2008), Korth (2008), and Korth and Samuels (2015). For European



**Text-fig. 1. Distribution of fossil localities where eomyid rodents were discovered.** 1: Sunduk, Kazakhstan, 2: Saint Jacques, Inner Mongolia, China; 3: Tieersihabahe in the Junggar Basin, Xinjiang, China, 4: 11 localities (Taatsiin Gol (right), Hasanda Gol, Tatal Gol, Taatsiin, Del, Huch Teeg, Unkheltseg, Luuny Yas, Hotuliin Teeg, Ulaan Tolgoi, Builstyn Khudag) in the Valley of Lakes, Mongolia, 5: Yandantu (Danghe area), Gansu, China, 6: 5 localities (Lower Aoerban, Gashunyinadeg, Upper Aoerban, Balunhalagen, Bilutu) in Sonid Zuoqi, 3 localities (Amuwusu, Shala, Loc. 346) in Sonid Youqi, 2 localities (Ertemte2, Harr Obo2) in Huade, Huitenghe in Abag qı, Inner Mongolia, China, 7: Uulantatal, Inner Mongolia, China, 8: Altynshokysu, Aktyubinsk, Kazakhstan, 9: Burqin, Habahe in the Junggar Basin, Xinjiang, China, 10: Songshan, Gansu, China, 11: Dota, Gifu, Japan, 12: Shihuiba, Lufeng, Yunnan, China, 13: Leilao, Yuanmou, Yunnan, China, 14: Sihong, Jiangsu, China, 15: Tagay in Olkhon Island, Lake Baikal, Siberia, Russia. More detailed information is provided in Appendix.

**Table 1. First and last occurrences of eomyid rodents at the genus level in Asia, Europe, and North America, corresponding to Text-fig. 10 and Appendix.**

Genus	First occurrence (Ma)	Last occurrence (Ma)	Region	Reference for min age	Reference for max age
<i>Apeomys</i>	16.9	16.9	Asia	Qiu (2017)	Qiu (2017)
<i>Asianeomys</i>	26.5	18.4	Asia	Qiu et al. (2013); Qiu and Li (2016)	Wang (2002); Wang et al. (2008); PBDB
<i>Eomys</i>	25.4	25.4	Asia	Wang and Emry (1991)	Wang and Emry (1991)
<i>Heteroeomys</i>	8.5	8.5	Asia	Qiu (2006)	Qiu (2006)
<i>Japaneomys</i>	18.5	18.5	Asia	Kimura et al. (2019)	Kimura et al. (2019)
<i>Keramidomys</i>	19.5	6.4	Asia	Qiu et al. (2013); Qiu and Li (2016)	Qiu et al. (2013); Qiu and Li (2016)
<i>Leptodontomys</i>	19.5	6.2	Asia	Qiu et al. (2013); Qiu and Li (2016)	Qiu et al. (2013); Qiu and Li (2016)
<i>Ligerimys</i>	19.5	18.4	Asia	Qiu et al. (2013); Qiu and Li (2016)	Qiu et al. (2013); Qiu and Li (2016)
<i>Megapeomys</i>	18.5	18.5	Asia	Takeuchi (1992); Tomida (2011); Mörs et al. (2016)	Takeuchi (1992); Tomida (2011); Mörs et al. (2016)
<i>Omboomys</i>	9.2	9.2	Asia	Maridet et al. (2015)	Maridet et al. (2015)
<i>Pentabuneomys</i>	19.5	9.4	Asia	Qiu et al. (2013); Qiu and Li (2016)	Qiu et al. (2013); Qiu and Li (2016)
<i>Plesieomys</i>	8.5	8.5	Asia	Qiu (2006)	Qiu (2006)
<i>Symplokeomys</i>	35.6	35.6	Asia	Emry et al. (1997); PBDB	Emry et al. (1997); PBDB
<i>Yuneomys</i>	8.5	8.5	Asia	Qiu (2006, 2017)	Qiu (2006, 2017)
<i>Apeomys</i>	23.2	16.8	Europe	Mörs and Flink (2018)	Maridet et al. (2013)
<i>Eomyodon</i>	24.3	22.4	Europe	Maridet et al. (2013)	Maridet et al. (2013)
<i>Eomyops</i> ( <i>Leptodontomys</i> )	16.8	4.2	Europe	Maridet et al. (2013)	Maridet et al. (2013)
<i>Eomys</i>	33.2	23.1	Europe	Maridet et al. (2010)	Maridet et al. (2013)
<i>Estramomys</i>	5.2	2.2	Europe	Maridet et al. (2013)	Maridet et al. (2013)
<i>Keramidomys</i>	15.3	5.2	Europe	Maridet et al. (2013)	Maridet et al. (2013)
<i>Ligerimys</i>	20.9	16.8	Europe	Maridet et al. (2013)	Maridet et al. (2013)
<i>Megapeomys</i>	18.4	16.8	Europe	Mörs and Flink (2018)	Maridet et al. (2013)
<i>Pentabuneomys</i>	18.4	15.3	Europe	Maridet et al. (2013)	Maridet et al. (2013)
<i>Pseudotheridomys</i>	25.0	16.8	Europe	Maridet et al. (2013)	Maridet et al. (2013)
<i>Rhodanomys</i>	23.7	20.9	Europe	Maridet et al. (2013)	Maridet et al. (2013)
<i>Ritteneria</i>	22.4	20.1	Europe	Maridet et al. (2013)	Maridet et al. (2013)
<i>Adjidaumo</i>	38.8	33.6	North America	PBDB	PBDB
<i>Arikareeomys</i>	22.6	22.6	North America	PBDB	PBDB
<i>Aulolithomys</i>	38.8	35.6	North America	PBDB	PBDB
<i>Centimanomys</i>	35.6	35.6	North America	PBDB	PBDB
<i>Cristadjidaumo</i>	38.8	35.6	North America	PBDB	PBDB
<i>Cupressimus</i>	35.6	35.6	North America	PBDB	PBDB
<i>Kansasimys</i>	7.6	7.6	North America	PBDB	PBDB
<i>Leptodontomys</i>	28.6	7.6	North America	PBDB	Korth (2008)
<i>Litoyoderimys</i>	38.8	35.6	North America	PBDB	PBDB
<i>Megapeomys</i>	18.2	18.2	North America	PBDB	PBDB
<i>Metadjidaumo</i>	33.6	28.6	North America	PBDB	PBDB
<i>Metanoiamys</i>	43.3	35.6	North America	PBDB	PBDB
<i>Namatomys</i>	35.6	35.6	North America	PBDB	PBDB
<i>Orelladjidaumo</i>	33.6	33.6	North America	PBDB	PBDB
<i>Paradjidaumo</i>	38.8	32.1	North America	PBDB	PBDB
<i>Paranamatomys</i>	35.6	35.6	North America	PBDB	PBDB
<i>Protadjidaumo</i>	43.3	38.8	North America	PBDB	PBDB
<i>Pseudotheridomys</i>	25.6	14.8	North America	PBDB	PBDB
<i>Viejadjidaumo</i>	38.8	38.8	North America	PBDB	PBDB
<i>Yoderimys</i>	35.6	35.6	North America	PBDB	PBDB
<i>Zemiodontomys</i>	35.6	35.6	North America	PBDB	PBDB
<i>Montanamus</i>	35.3	35.3	North America	Flynn (2008)	Flynn (2008)
<i>Aguafriamys</i>	38.5	38.5	North America	Flynn (2008)	Flynn (2008)
<i>Apeomyoides</i>	15.4	15.4	North America	Flynn (2008)	Flynn (2008)
<i>Comanchoeomys</i>	7.1	7.1	North America	Flynn (2008)	Flynn (2008)
<i>Ronquillomys</i>	6.0	6.0	North America	Flynn (2008)	Flynn (2008)
<i>Zetamys</i>	25.6	25.6	North America	PBDB	PBDB

taxa, we utilized the dataset of Maridet et al. (2010, 2013), supplemented by Mörs and Flink (2018). We follow Engesser (1979) in considering *Leptodontomys* and *Eomyops* to be distinct genera although the validity of *Eomyops* is currently debated (Qiu 1994, Ruiz-Sánchez et al. 2009, Prieto 2012). For the Asian taxa, we compiled a detailed dataset based on the original literature (see Appendix). Age for upper and lower boundaries of MP and MN zones are taken from the Geologic Time Scale 2012 (Hilgen et al. 2012, Vandenberghe et al. 2012).

The median of the estimated range was used for the age of each fossil specimen. We excluded any entry whose age estimates range over 6 million years or identification is indeterminate at the species level. The choice of 6 million years for the quality control of the data is rather arbitrary to include the single occurrence of Asian *Apeomys* although age estimates are less than 3 million years for most of the localities. All genera but *Zetamys*, for which the age uncertainty of the only known locality is over 10 million years (Martin 1974), met the above criteria and were therefore included in the analysis. Generic richness herein is represented by actual occurrences using a range-through approach in one million-year time bins. The time bins start at 45.5 Ma and end at 0.5 Ma. The upper limit of a bin is included, and the lower limit is excluded. For more sophisticated data standardisations, readers are directed to an excellent review about sampling bias and data standardisations for biodiversity by Aberhand and Kiessling (2012) and references therein. Tooth terminology follows Emry et al. (1997).

## Abbreviations

MN	Mammal Neogene zone for European land mammal age
MP	Mammal Paleogene zone for European land mammal age
M or m	Molar, the upper case for the upper dentition and the lower case for the lower dentition
P or p	Premolar, the upper case for the upper dentition and the lower case for the lower dentition
IVPP	Institute of Vertebrate Paleontology and Paleoanthropology, Beijing, China
IZ NAS RK	Institute of Zoology, National Academy of Sciences, Republic of Kazakhstan
NHMW	Museum of Natural History Vienna, Vienna, Austria
NMNS	National Museum of Nature and Science, Tokyo, Japan

## Systematic palaeontology

A total of 47 eomyid genera have been described to date. Of these, fourteen genera are reported from Asia, and half of them are endemic to Asia or only known from the type locality. In this section, we review the Asian eomyids, especially focusing on the endemics with a brief description of diagnostic dental traits, locality occurrences (Text-fig. 1), and stratigraphic ranges (Text-fig. 2). The taxonomic data were newly compiled along with geological and biostratigraphical information in Appendix, which were also

used to explore the eomyid diversity patterns in Asia in the last part of this study.

### Family Eomyidae WINGE, 1887

#### Subfamily Eomyinae WINGE, 1887

##### Tribe Eomyini WINGE, 1887

#### Genus *Symplokeomys* EMRY, WANG, TJUTKOVA et LUCAS, 1997

Text-figs 1–3

Type species. *Symplokeomys zaysanicus* EMRY, WANG, TJUTKOVA et LUCAS, 1997.

Included Asian species. The type and only species.

Dental characteristics. Medium-sized eomyid, apparently retaining P3 as in Yoderiminae; branching mesoloph(id) and interconnecting lophules in synclines/synclinids II and III, forming complex bunolophodont patterns; no lingual part of the anteroloph; hypolophid connecting to posterolophid; presence of spurs in transverse loph(id)s.

Asian occurrences (age). For the type and only species, locality K10 in Zaysan Basin, Sunduk, Kazakhstan (Aksyr Formation, Ergilian, late Eocene).

Comments. The genus is peculiar compared to any other known eomyids in having extremely complex occlusal patterns with interconnecting lophules in synclines/synclinids II and III as well as retaining P3 as in the subfamily Yoderiminae (Emry et al. 1997). The authors note that the presence of P3 is a primitive condition in the Eomyidae and was independently retained in *Symplokeomys*. Phylogenetic relationships of *Symplokeomys* with other eomyids have not been resolved.

#### Genus *Asianeomys* WU, MENG, YE et NI, 2006

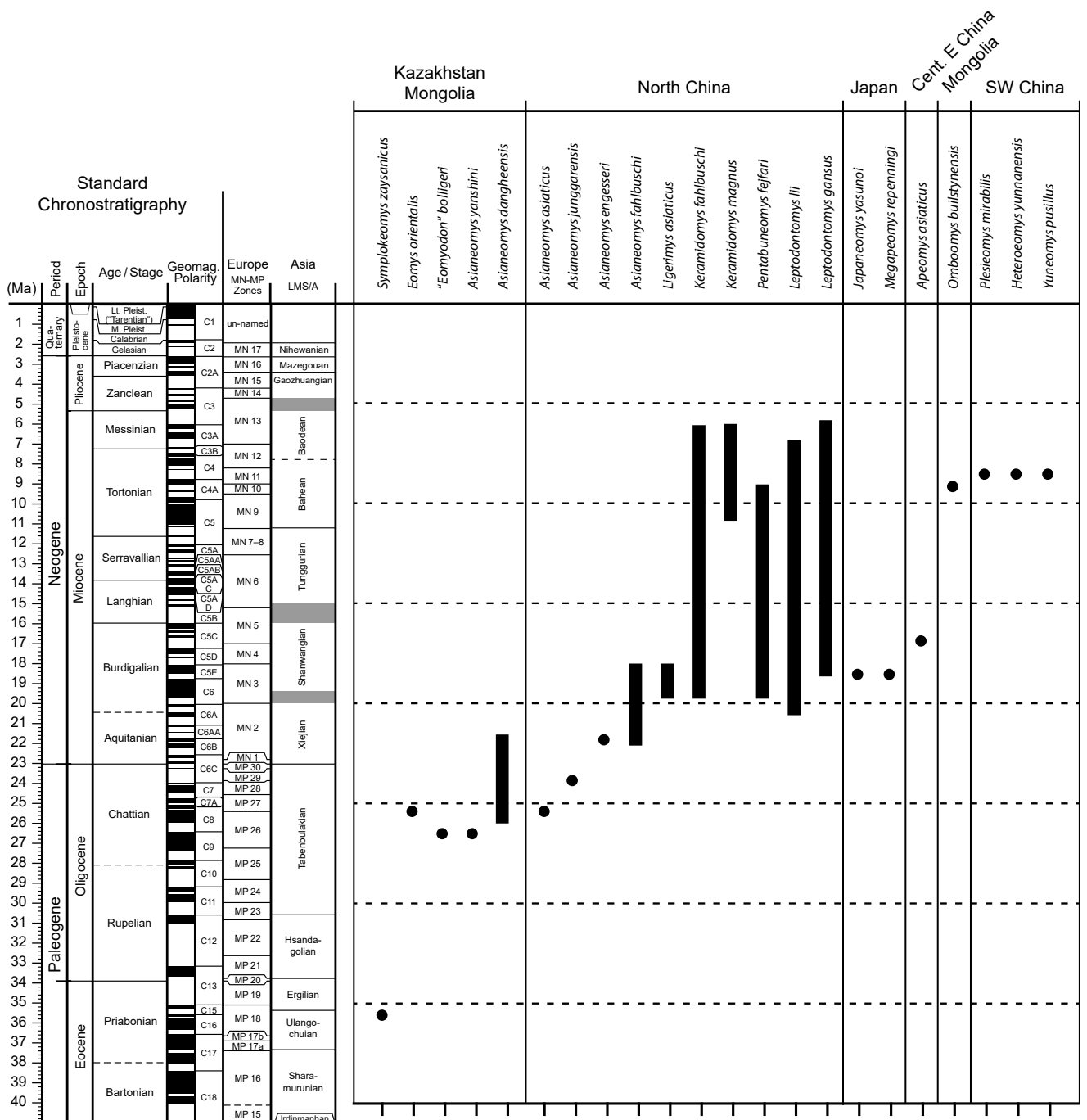
Text-figs 1, 2, 4

Type species. *Asianeomys junggarensis* WU, MENG, YE et NI, 2006.

Included Asian species. *Asianeomys junggarensis*, *Asianeomys asiaticus*, *Asianeomys dangheensis*, *Asianeomys fahlbuschi*, *Asianeomys yanshini*, *Asianeomys engesseri*. *Asianeomys engesseri* could be a junior synonym of *A. fahlbuschi* according to Qiu and Li (2016). Note that “*Eomyodon*” *bolligeri* from Kazakhstan possibly belongs to the genus *Asianeomys* (Maridet et al. 2015).

Dental characteristics. Bunolophodont; small to medium-sized eomyid; P4 with or without anteroloph; entoloph connecting to the posterior end of the protocone rather than protoloph, differing from *Pseudotheridomys*, so that the syncline II is longer than the syncline I; three-rooted p4 lacking anteroconid; four roots on m1–m2; hypolophid connecting to the anterior arm of the hypoconid, differing from more posterior connection in *Eomys*; mesoloph(id) usually long; hypolophid present on m3 (Wu et al. 2006, Qiu and Li 2016).

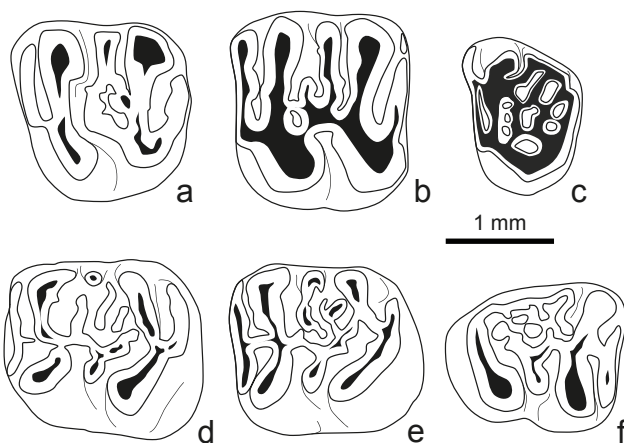
Asian occurrences (age). For *A. junggarensis*, localities XJ98035, XJ98024, XJ200209, Tieersihabahe of North Junggar Basin, China (Tieersihabahe mammal assemblage zone I, late Oligocene, 24.4 to 23.2 Ma by Meng



**Text-fig. 2.** Stratigraphic range of Asian eomyids based on “range-through” occurrences. Standard chronostratigraphy was built in TimeScale Creator version 7.3 (<https://engineering.purdue.edu/Stratigraphy/tscreator/index/index.php>), which is based on the GTS 2016 age model. For mammalian biochronology, European Neogene Mammal (MN) unit and Paleogene Mammal (MP) unit as well as the Asian land mammal ages in the Paleogene were also downloaded from TimeScale Creator version 7.3. Neogene Chinese land mammal stage/ages (LMS/A) are based on Qiu and Li (2016).

et al. 2006) (Wu et al. 2006). For *A. dangheensis*, locality XJ99005, North Junggar Basin, China (Suosuoquan Mammal assemblage zone II, early Miocene, 21.9 to 21.7 Ma by Meng et al. 2006) (Wu et al. 2006); 6 different sections (Del, Huch Teeg, Tatal Gol, Unkheltseg, Luun Yas, Hotuliin Teeg) within 660 km<sup>2</sup> in Central Mongolia (Valley of Lakes) (Loh Formation, local biozones C1, C1/D, D, late Oligocene to early Miocene, 25.6 to 21 Ma by Daxner-Höck et al. 2017) (Maridet et al. 2015); IVPP Loc. DH199904, Aksay Kazak Autonomous County, Gansu, China (Paoniuzhan Formation, Tabenbulukian, 28.4 to 23 Ma) (Wang 2002, Wang et al.

2008). For *A. fahlbuschi*, locality XJ99005, North Junggar Basin, China (Suosuoquan mammal assemblage zone II, early Miocene, 21.9 to 21.7 Ma by Meng et al. 2006) (Wu et al. 2006); two sites located 40 km apart in central Inner Mongolia, China, Aoerban and Gashunyinadege in stratigraphic order (the lower red member of the Aoerban Formation to Gashunyinadege bed, late Xiejian to early Shanwangian, correlated to MN 2 – MN 3, 21.7 to 17.2 Ma) (Qiu and Li 2016). For *A. asiaticus*, IVPP Loc. 77048, Saint Jacques, Inner Mongolia, China (Wulangulage Formation, late Oligocene, 27.8 to 23 Ma) (Wang and Emry 1991). For



**Text-fig. 3.** *Symplokeomys zaysanicus* EMRY, WANG, TJUTKOVA et LUCAS, 1997 from type locality. a: left P4 (IZ NAS RK 35 (10)/1829), b: left M1 or M2 (IZ NAS RK 35 (10)/1831), c: left M3 (IZ NAS RK 35 (10)/1836), d: left m1 (IZ NAS RK 35 (10)/1843, holotype), e: left m2 (IZ NAS RK 35 (10)/1845), f: right m3 (IZ NAS RK 35 (10)/1853). A scale equals to 1.0 mm.

*A. yanshini*, Altynshokysu locality, Aktyubinsk, Kazakhstan (Aral Formation, correlated with the Central Mongolian local biozone C, ~27.4 to ~25.6 Ma, by Bendukidze 2009) (Lopatin 2000). For *A. engesseri*, locality XJ99005, North Junggar Basin, China (Suosuoquan mammal assemblage zone II, early Miocene, 21.9 to 21.7 Ma by Meng et al. 2006) (Wu et al. 2006). For *A. aff. engesseri*, locality XJ200604, northern Xinjiang province, China (age close to

Sihong and Gashunyinadege localities, late early Miocene, ~17 to ~18 Ma, Deng 2006) (Maridet et al. 2011). For cf. *Asianeomys bolligeri*, 2 different sections within 10 km in central Mongolia (Valley of Lakes) (Local biozone C, late Oligocene, ~27.4 to ~26.5 Ma by Daxner-Höck et al. 2017) (Maridet et al. 2015). For “*Eomyodon*” *bolligeri* from Kazakhstan, North Aral Region (Altyn shokysu locality, bone bed I and II, Aral Formation, late Oligocene, correlated with the Central Mongolian local biozone C, ~27.4 to ~25.6 Ma by Bendukidze 2009) (Lopatin 2000).

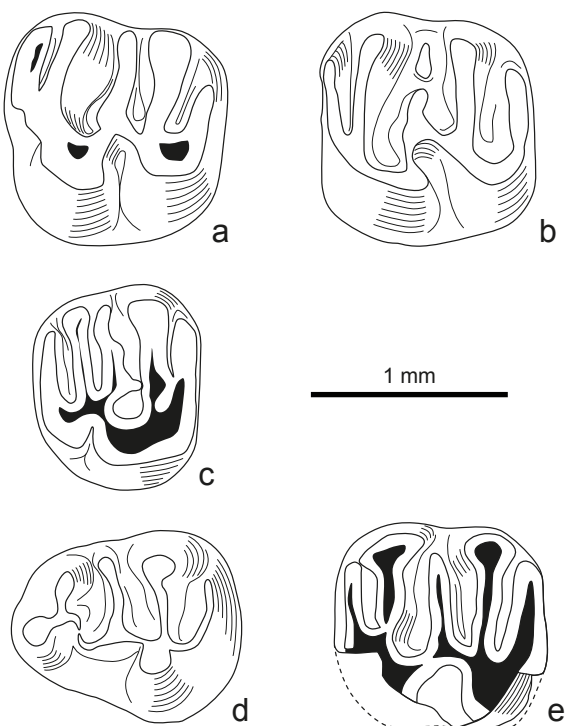
**Comments.** Wu et al. (2006) erected the genus *Asianeomys* for material from the Junggar Basin in Xinjiang, North China. They transferred all Asian species of eomyid rodents known at the time to the new genus: two species of *Pseudotheridomys* (“*P.*” *asiaticus*, “*P.*” *yashini*) and one species of *Eomyodon* (“*E.*” *dangheensis*). The Asian occurrence of *Pseudotheridomys* (particularly, “*P.*” *asiaticus*, slightly earlier than the European occurrence of the genus) was once considered to be possible evidence for Asian origin of the genus (Korth 2008), but currently no valid species of *Pseudotheridomys* is reported from Asia. Because *Asianeomys* has *Eomys*-like upper dentition and *Pseudotheridomys*-like lower dentition, it is difficult to properly identify the genus when only small samples are available. To date, *Asianeomys* has been reported from North China, Mongolia, and Kazakhstan. The genus is more closely related to *Keramidomys* than *Pseudotheridomys* based on dental similarity to a basal species *Keramidomys*, *K. fahlbuschi*, recently discovered in Inner Mongolia, China (Qiu and Li 2016, Kimura et al. 2019).

#### Genus *Eomys* SCHLOSSER, 1884

**Comments.** *Eomys orientalis* is the only Asian species of *Eomys*. It was first discovered from the Oligocene of Inner Mongolia, China (Wang and Emry 1991). Wang and Emry (1991) pointed out that *E. orientalis* is similar to European *Eomys* in the more lingually extended synclines, whereas the species is also similar to North American *Adjidaumo* in having a weakly developed synclinid IV. Overall, *Eomys orientalis* is most similar to *E. antiquus*, which represents the earliest occurrence (early Oligocene, MP 21) of the family in Europe (Maridet et al. 2010). Until recently, the record of Asian *Eomys* had been limited to the type locality with a handful of specimens described in Wang and Emry (1991). However, Maridet et al. (2015) recently reported new material provisionally assigned to *Eomys* cf. *orientalis* from the early Oligocene of Central Mongolia.

#### Genus *Keramidomys* HARTENBERGER, 1966

**Comments.** Two species are known from Asia: *K. fahlbuschi* QIU, 1996 and *K. magnus* QIU et LI, 2016 (see Appendix). *Keramidomys* has been reported from multiple localities within an area of 13,000 km<sup>2</sup> in central Inner Mongolia, China. In addition, few specimens were found in Siberia (Russia), Valley of Lakes (Mongolia), and Xinjiang (China) (Appendix 1). The stratigraphic range of the genus is as long as that of cosmopolitan *Leptodontomys* in Asia, ranging from the Xiejian (correlated to MN 2 – MN 3, 21.7 to 17.2 Ma) to Baodean (correlated to MN 12 – MN 13,



**Text-fig. 4.** *Asianeomys junggarensis* WU, MENG, YE et NI, 2006 from type locality. a: left P4 (IVPP V14452.2, paratype), b: left M1/2 (IVPP V 14452.1, holotype), c: right M3 (IVPP V14454.1), d: left p4 (IVPP V14452.5, paratype), e: left m1/m2 (IVPP V14453.2). A scale equals to 0.5 mm.

7.4 to 5.3 Ma), geologically from the Aoerban Formation and Gashunyinadege bed to the Bilutu bed, represented by *K. fahlbuschi* (Qiu and Li 2016). The other species known from the region is *K. magnus*, which presents a shorter range than *K. fahlbuschi*, recorded in late Miocene local faunas equivalent in age to MN 9 to MN 13. *Keramidomys* is thought to have originated in East Asia and later dispersed into Europe (Mein 2009), considering the earlier occurrence of Asian *K. fahlbuschi*, which is generally less lophodont than European species (Qiu and Li 2016, Kimura et al. 2019). It first appears in the European record at MN 5 (Mein 2009), which postdate the regional climax of the eomyid diversity.

#### **Genus *Leptodontomys* SHOTWELL, 1956 or *Eomyops* ENGESSER, 1979**

**Comments.** The genera *Eomyops* and *Leptodontomys* may be synonyms. Many different European species had been long assigned to the North American genus *Leptodontomys* until Engesser (1979) erected the new genus *Eomyops* for the European forms which differ from *Leptodontomys* in details of the mandible morphology and the presence of two ridges on the enamel of the lower incisors. Yet, the validity of *Eomyops* has been questioned and is still debated because cheek teeth of both genera are remarkably similar (Qiu 1994, de Bruijn et al. 2012). Generally, Asian species are ascribed to *Leptodontomys* (Qiu and Li 2016) or *Leptodontomys/Eomyops* (Maridet et al. 2015) until lower incisors of Asian species are found to clarify their taxonomic attribution.

Two species are known from Asia: *Leptodontomys gansus* ZHENG et LI, 1982 and *L. lii* QIU, 1996 (Text-fig. 2, Appendix). *Leptodontomys gansus* was the first eomyid to be discovered in Asia (Zheng and Li 1982) and is represented by a right mandible with a complete set of cheek teeth. Later on, more specimens of the same species were discovered from nine localities in large areas within 32,500 km<sup>2</sup> in central Inner Mongolia, China (Qiu and Li 2016), stratigraphically ranging from the Aoerban Formation and the Gashunyinadege bed (correlated to MN 2 – MN 3, 21.7 to 17.2 Ma) to the Ertemte Formation (correlated to MN 13, 7.1 to 5.3 Ma). *Leptodontomys lii* was first found from the middle Miocene Moergen II locality in central Inner Mongolia, China (Qiu 1996), and later additional materials were found from older and younger deposits in the same areas, mostly overlapping with the stratigraphic occurrences of *L. gansus* except for the late late Miocene Ertemte2 and Harr Obo2 localities, where the latter species are present. *Leptodontomys lii* is similar in dental morphology to *L. gansus* but is distinguishable by larger tooth size with more bulbous main cusps and more pronounced mesoloph(id) with slightly higher crown (Qiu and Li 2016). A fossil of *Leptodontomys/Eomyops* sp. was also found in the local biozone D1/2 (late early Miocene) in Mongolia (Maridet et al. 2015). A small m1/m2 fossil found in Olkhon Island (Lake Baikal, Siberia) was assigned to *Eomyops oppligeri* ENGESSER, 1990 based on its size and morphology (Daxner-Höck et al. 2013). However, the taxonomic assignment should be reconsidered because the fossil is also within the size range of and is comparable in morphology to *Leptodontomys gansus* as shown in fig. 72: I of Qiu and Li (2016).

Qiu (1994) noticed that middle to late Miocene Chinese *Leptodontomys* are intermediate between North American *Leptodontomys* and European *Eomyops* in dental traits on the lower molars and mandible shape. Therefore, Qiu (1994) suggested European *Eomyops* was derived from North American *Leptodontomys* species that dispersed through Asia although the first appearance of Chinese *Leptodontomys* in Tunggur (late middle Miocene), Inner Mongolia, was younger than the first appearance of European *Eomyops* (then, dated to the earliest middle Miocene, MN 5) when his study was published. Currently, fossils of Asian *Leptodontomys* have been discovered from early Miocene deposits such as the lower red member of the Aoerban Formation and the Gashunyinadege bed, correlated to MN 2 – MN 3, in Inner Mongolia, China (Qiu and Li 2016), and thus significantly older than the first European record of *Eomyops*, MN 4 of Mortera-20 A in Valencia, Spain (Ruiz-Sánchez et al. 2009). Owing to the series of discoveries in Inner Mongolia, *Leptodontomys* along with *Keramidomys* are the longest-ranging eomyids in Asia (and the same is true for Europe; see Engesser 1999).

#### **Genus *Pentabuneomys* ENGESSER, 1990**

**Comments.** The genus *Pentabuneomys* included a single species from Europe (*Pentabuneomys rhodanicus* (HUGENEY and MEIN 1968); MN 3 to MN 4, early Miocene) and another from North America (*P. engesseri* KORTH, 2008, early Arikareean, late Oligocene) until the discovery of *Pentabuneomys fejsari*, which is currently its only Asian representative (Qiu and Li, 2016). *Pentabuneomys fejsari* was described based on the total of nearly 40 specimens from the early Miocene Lower Aoerban fauna/Gashunyinadege fauna (correlated to MN 2 – MN 3) and the late Miocene Amuwusu fauna (correlated to MN 9). *Pentabuneomys* was erected by Engesser (1990) to separate European “*Eomys*” *rhodanicus* from other *Eomys* species. *Pentabuneomys* differs from *Eomys* in having bulbous cusps with a large circular mesocone/mesoconid and the presence of a deeper valley between the hypolophid and the posterior cingulum (Engesser 1990, Korth 2008). Korth (2008) reported the first record of *Pentabuneomys* from North America. The North American species, *P. engesseri* is not only chronologically older but also more plesiomorphic than European *P. rhodanicus* in completely lacking the mesolophid and the ectolophid. Thus, it could be possible that the genus originated in North America and later dispersed into Europe through Asia in the late Oligocene to early Miocene. Korth (2008) suggested that *Pentabuneomys* was derived from a *Leptodontomys*-like ancestor with reduced lophs in cheek teeth. Qiu and Li (2016) also discussed the similarity between *Leptodontomys/Eomyops* and *Pentabuneomys* to suggest that Asian *P. fejsari* is intermediate in occlusal morphology between European *Eomyops hebeiseni* and *Pentabuneomys rhodanicus*. The phylogenetic relationships of *Pentabuneomys* in relation to the genera mentioned above is still uncertain.

#### **Genus *Ligerimys* STEHLIN et SCHAUB, 1951**

**Comments.** Three specimens from early Miocene localities (equivalent to MN 2 – MN 3) in Inner Mongolia

(China) were assigned to the first Asian species of *Ligerimys*: *L. asiaticus*, which is larger than other species of *Ligerimys* (Qiu and Li 2016). In addition to the diagnostic traits mentioned by Qiu and Li (2016), the Asian species is characterized by the synclinid II closed by the posterior extension of longitudinally elongated metaconid. *Ligerimys* is considered to be derived from *Pseudotheridomys* (Escarguel and Aguilar 1997, Engesser 1999) and was considered to be endemic to Europe before *L. asiaticus* was discovered in Inner Mongolia. More specimens are necessary to understand the phylogenetic relationships of this species and closely related taxa.

**Genus *Japaneomys* KIMURA, TOMIDA, KALTHOFF,  
CASANOVAS-VILAR et MÖRS, 2019**

Text-figs 1, 2, 5

**Type species.** *Japaneomys yasunoi* KIMURA, TOMIDA, KALTHOFF, CASANOVAS-VILAR et MÖRS, 2019.

**Included Asian species.** The type and only species.

**Dental characteristics.** Characters shared with *Asianeomys*, *Keramidomys*, and *Estramomys* are four roots in m1 and the hypolophid anteriorly connecting to the anterior arm of hypoconid. More bunodont with lower lophids than the genera listed above, yet transverse lophids complete; small eomyid, similar in size to *Keramidomys*; two rooted p4 (rather than three roots as in all species of *Asianeomys* except *A. junggarensis*); anterior lobe narrower than posterior lobe on m1 (correspondingly, synclinid I of m1 much shallower and shorter than half the length of synclinid IV); hypolophid anteriorly concave on m1 (Kimura et al. 2019).

**Asian occurrences (age).** For the type and only species, Dotsu Town, Kani City, Gifu Prefecture, central Japan (near the uppermost level of the Nakamura Formation, Mizunami Group; age estimated to be around 18.5 Ma) (Kimura et al. 2019).

**Comments.** The genus *Japaneomys* was erected based on a right mandible with p4 and m1 discovered from the early Miocene of central Japan (Kimura et al. 2019). An m1 of *Megapeomys* was also discovered in the same locality (Tomida 2011). The jaw was originally ascribed to *Pseudotheridomys* sp. (Tomida and Setoguchi 1994) but later Tomida (2011) noted its morphological similarities with *Keramidomys*. However, *Japaneomys* retains plesiomorphic dental traits compared to contemporaneous eomyids such as *Keramidomys* (Kimura et al. 2019). Phylogenetic analyses have revealed that *Japaneomys* diverged earlier than *Asianeomys* and *Keramidomys* in the late Oligocene, when Japan was part of the coastal margin of East Asia. We suggest that some degree of geographic isolation existed at that time, thus separating certain lineages of small mammals in the coastal region from those in inner-continental regions of Asia.

**Genus *Omboomys* MARIDET, DAXNER-HÖCK, BADAMGARAV  
et GÖHLICH, 2015**

Text-figs 1, 2, 6

**Type species.** *Omboomys builstynensis* MARIDET, DAXNER-HÖCK, BADAMGARAV et GÖHLICH, 2015.

**Included Asian species.** The type and only species.

**Dental characteristics.** Small-sized bunodont eomyid. It is clearly smaller than other Asian endemic genera such as *Asianeomys* and *Heteroeomys* and slightly smaller than *Japaeomys*. Characterized by a slender morphology and a low crown, but with cusp(id)s noticeably higher than the mid-portion of the tooth in antero-posterior view. Other noticeable characteristics are upper teeth missing mesoloph, whereas in lower teeth the mesolophid is rare and weakly developed (thus, the molars only have three syncline(id)s with the central one being much wider and larger than the others). M1/2 with trapezoidal outline and oblique, short longitudinal crest joining directly the metaloph to the posterior end of the protocone. p4s with oblique longitudinal crest joining the hypoconid with the metalophid or with the metaconid (but never with the protoconid).

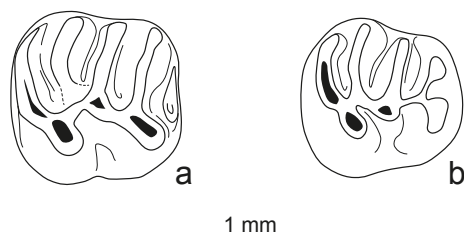
**Asian occurrences (age).** For the type and only species, locality Builstyn Khudag, Mongolia (Tuyn Gol Formation, local biozone E, late Miocene, younger than 13 Ma).

**Comments.** Late Miocene *Omboomys* is an endemic eomyid genus in the late Miocene of northeastern Asia. However, the origin of *Omboomys* is unknown due to the lack of an intermediate fossils between the late early Miocene and the late Miocene in the Valley of Lakes in Mongolia (Daxner-Höck et al. 2017). The general bunodont morphology, the small size and low crown along with the trapezoidal shape of the M1/2, and the curved longitudinal crest connecting to the protocone in P4–M1/2 suggest its closer phylogenetic affinity to *Leptodontomys/Eomyops* than any other genus of Eomyidae known in the Miocene of Eurasia. The disappearance of the mesoloph(id)s is also observed in other eomyid clades in the Miocene of Europe such as the *Rhodanomys-Ritteneria* clade or the *Keramidomys-Estramomys* clade (Engesser 1990, 1999). Considering the more likely affinity of *Omboomys* with *Leptodontomys/Eomyops*, the disappearance of the mesoloph[id]s seems to be a convergent evolution, resulting from a local evolutionary history leading to *Omboomys* in northeastern Asia.

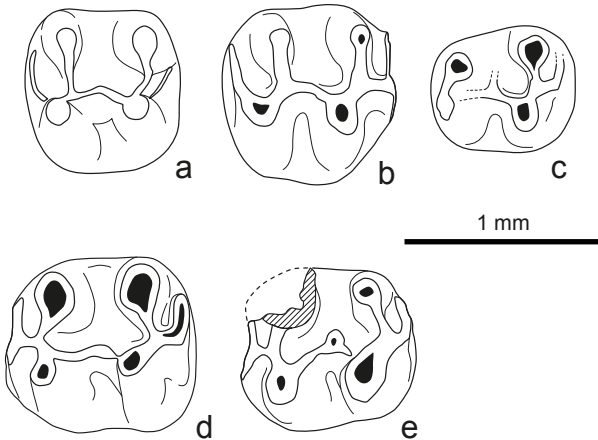
**Genus *Heteroeomys* QIU, 2006**

Text-figs 1, 2, 7

**Type species.** *Heteroeomys yunnanensis* QIU, 2006.



**Text-fig. 5.** *Japaneomys yasunoi* KIMURA, TOMIDA, KALTHOFF, CASANOVAS-VILAR et MÖRS, 2019 from type locality. a: right m1 (NMNS-PV19995, holotype), b: right p4 (NMNS-PV19994, paratype). Note that specimens were labelled reversely by mistake in the original literature. A scale equals to 0.5 mm.



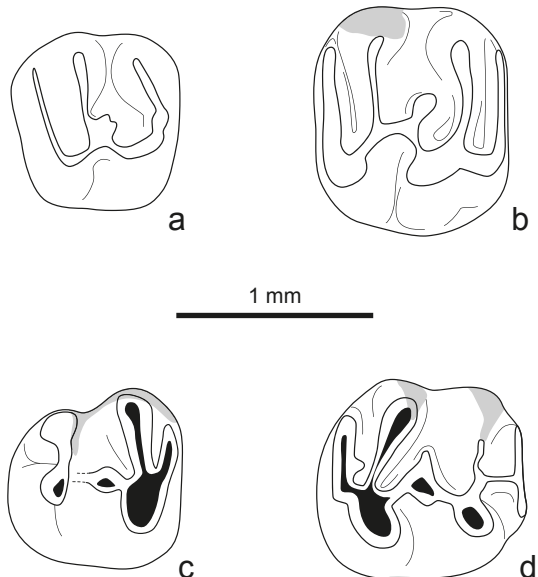
**Text-fig. 6.** *Omboomys builstynensis* MARIDET, DAXNER-HÖCK, BADAMGARAV et GÖHLICH, 2015 from type locality. a: right P4 (NHMW 2013/0076/0002), b: right M1/2 (NHMW 2013/0076/0004, holotype), c: left p4 (NHMW 2013/0076/0009), d: left m1/2 (NHMW 2013/0076/0013), e: left m3 (NHMW 2013/0076/0014). A scale equals to 0.5 mm.

Included Asian species. The type and only species.

**Dental characteristics.** Bunodont; medium-sized eomyid, similar in size to *Asianeomys*, slightly larger than *Plesioeomys* and within large size variation of *Eomyops*. Plesiomorphic features include mesoloph(id) of moderate length; main cusps asymmetrically developed between the labial and lingual sides; weak anterolophid shorter than metalophid; weak ectolophid; protoloph and metaloph directed more perpendicular than in *Pseudadjidaumo*. Derived features comprise relatively higher crown, anteroloph extending far lingually and connecting directly to the protocone without the lingual part of protocone, which is unique among any known eomyid genera (modified from Qiu 2006).

**Asian occurrences (age).** For the type and only species, locality Shihuiba, Lufeng, Yunnan Province, China (Shihuiba Formation, late Miocene, not well dated but see Flynn and Qi 1982, Ni and Qiu 2002, Appendix); Leilao, Yuanmou, Yunnan Province, China (Xiaohe Formation, early Baodean, biostratigraphically around 9 Ma by Flynn and Qi 1982, Ni and Qiu 2002).

**Comments.** Eomyid rodents were only reported from the modern Palaearctic realm until a series of finds in the late Miocene hominoid localities, Shihuiba and Leilao, Yunnan Province in southwestern China, which is part of the modern Oriental realm (Qiu 2006) (Text-fig. 1). The Yunnan eomyids are represented by three endemic genera with bunodont tooth patterns: *Heteroeomys*, *Plesioeomys*, *Yuneomys*. They not only demonstrate that eomyids were distributed in tropical forest environments of Asia but also suggest regionally unique evolutionary histories of Asian eomyid rodents with several intercontinental dispersals. Qiu (2006) pointed out that Asian eomyids in the northern region have more lophodont tooth patterns than those in the southern region, reflecting different vegetation types with more open environments in the north. The Yunnan eomyids are similar in dental traits to North American *Pseudadjidaumo* and *Leptodontomys*/



**Text-fig. 7.** *Heteroeomys yunnanensis* QIU, 2006 from type locality. a: right P4 (IVPP V 14730.3), b: right M1/2 (IVPP V 14729, holotype), c: left p4 (IVPP V 14731.5), d: right m1/2 (IVPP V 14731.6). A scale equals to 0.5 mm.

*Eomyops*. However, phylogenetic relationships among these genera remain unknown.

#### Genus *Plesioeomys* QIU, 2006

Text-figs 1, 2, 8

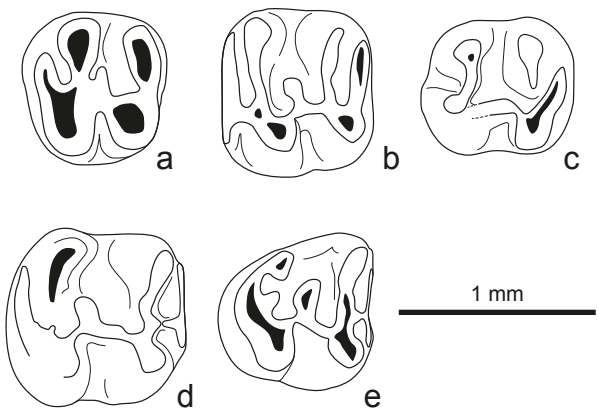
**Type species.** *Plesioeomys mirabilis* QIU, 2006.

**Included Asian species.** The type and only species.

**Dental characteristics.** Bunodont; relatively small eomyid, similar in size to *Pseudadjidaumo* and within large size variation of *Eomyops*; main cusps symmetrically developed; lingual portion of the anteroloph greatly reduced; anteroloph weakly developed; syncline II longer than other synclines; mesolophid of medium length with minute terminal cuspule present; metaloph poorly developed; lingual syncline symmetrical; short anterolophid connecting to metalophid through a longitudinal crest; hypolophid variably absent or interrupted (modified from Qiu 2006).

**Asian occurrences (age).** For the type and only species, locality Shihuiba, Lufeng, Yunnan Province, China (Shihuiba Formation, late Miocene, not well dated but see Flynn and Qi 1982, Ni and Qiu 2002, Appendix); Leilao, Yuanmou, Yunnan Province, China (Xiaohe Formation, early Baodean, biostratigraphically approximately 9 Ma after Flynn and Qi 1982, Ni and Qiu 2002).

**Comments.** Qiu (2006) pointed out that *Plesioeomys* is morphologically similar to *Pseudadjidaumo* in cusp size, weakly developed loph(id)s, and closed synclines I and IV. The Yunnan specimens were assigned to a new genus because loph(id)s are less developed than in *Pseudadjidaumo*, which is known from the Mojave Desert in North America (Lindsay 1972). In *Plesioeomys*, a mesoloph enlarged at the terminal point is considered to be a diagnostic trait.



**Text-fig. 8.** *Plesiomys mirabilis* QIU, 2006 from type locality. a: right P4 (IVPP V 14727.1), b: left M1/2 (IVPP V 14726, holotype), c: left p4 (IVPP V 14727.4), d: right m1/2 (IVPP V 14728.3), e: right m3 (IVPP V 14727.11). A scale equals to 0.5 mm.

Prieto (2012) noted that the enlarged termination of the mesoloph is present in European species of *Eomyops* such as *E. noelinae* (Ruiz-Sánchez et al. 2009) and also variably present in *E. catalaunicus* from German localities (Prieto 2012). In fact, the upper dentition of *Plesiomys* seems to be indistinguishable from that of *Eomyops*. Maridet et al. (2015) also pointed out this similarity and briefly stated the possible synonymy of *Plesiomys*, as well as *Heteroeomys* from the same locality, to *Leptodontomys/Eomyops*. Here, we retain the genus *Plesiomys*, considering the hypolophid being absent or indistinct as a diagnostic trait at the generic level. However, a detailed review of Asian material in comparison with European *Eomyops* is required to clarify the validity of this genus.

#### Genus *Yuneomys* QIU, 2017

Text-figs 1, 2, 9

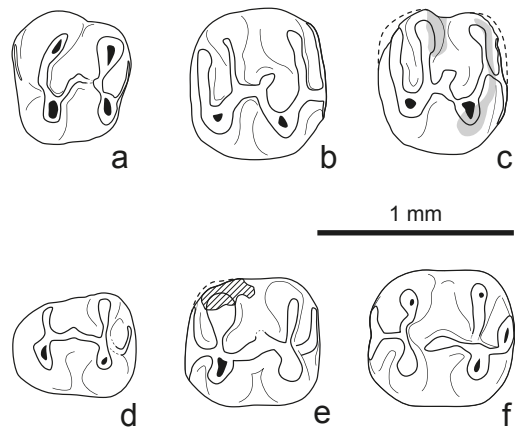
Type species. *Yuneomys pusillus* (QIU, 2006).

Included Asian species. The type and only species.

**Dental characteristics.** Bunodont; small eomyid, smaller or equivalent in size to *Leptodontomys* and *Eomyops*; short mesoloph; complete entoloph; small mesoconid; anterolophid connecting to the metalophid through a longitudinal crest at the midline of the tooth. Differing from the above genera in having m1/2 with metalophid and hypolophid less developed, short, and directed transversely; symmetrical arrangement of cusps and lophids between the anterior and posterior lobes (i.e., metaconid, protoconid, anterolophid vs. entoconid, hypoconid, posterolophid); mesolophid absent (modified from Qiu 2017; this study).

**Asian occurrences (age).** For the type and only species, locality Shihuiba, Lufeng, Yunnan Province, China (Shihuiba Formation, late Miocene, not well dated but see Flynn and Qi 1982, Ni and Qiu 2002, Appendix).

**Comments.** The genus was originally described as a new species of *Leptodontomys* (Qiu 2006). However, it was



**Text-fig. 9.** *Yuneomys pusillus* (QIU, 2006) from type locality. a: left P4 (IVPP V 14733.4), b: right M1/2 (IVPP V 14732, holotype), c: right M1/2 (IVPP V 14733.1), d: left p4 (IVPP V 14733.5), e: left m1/2 (IVPP V 14733.2), f: left m1/2 (IVPP V 14733.6). A scale equals to 0.5 mm.

later transferred to a newly erected genus, *Yuneomys* (Qiu 2017). Thus, all of the Yunnan eomyid genera (*Heteroeomys*, *Plesiomys*, *Yuneomys*) are endemic to southwestern China although they are morphologically similar to *Eomyops* and *Leptodontomys*. Scarce specimens of *Yuneomys* preclude accurate differential diagnosis, particularly in comparison with *Eomyops* and *Leptodontomys*, which show considerable variation in cheek tooth morphology. The upper dentition of *Yuneomys* is in fact identical to that of the two genera. Here, we tentatively retain *Y. pusillus* encased in *Yuneomys*. Nevertheless, as stated above, we suggest that the validity of *Yuneomys* and *Plesiomys* should be reexamined, particularly taking into account the large variation in *Eomyops* and *Leptodontomys*.

#### Subfamily Apeomyinae FEJFAR, RUMMEL et TOMIDA, 1998

**Comments.** The subfamily Apeomyinae, which contains *Apeomys*, *Megapeomys*, *Apeomyoides*, and *Arikareeomys* is known based on extremely scarce materials and is restricted to the late Oligocene and early Miocene (Fahlbusch 1968, Korth 1992, Engesser 1999, Smith et al. 2006, Flynn 2008, Mörs and Flink 2018). Nevertheless, apeomyine eomyids are Holarctic in distribution, occurring in Europe (*Apeomys*, *Megapeomys*), North America (*Megapeomys*, *Apeomyoides*, *Arikareeomys*), and Asia (*Apeomys*, *Megapeomys*).

#### Genus *Megapeomys* FEJFAR, RUMMEL et TOMIDA, 1998

**Comments.** *Megapeomys* is phylogenetically closest to *Apeomys* (Engesser 1999). In the Asian record *Megapeomys* is as scarce as *Apeomys* and may predate it slightly. A single species of *Megapeomys*, *M. repenningi*, was described in Asia based on a single tooth from the early Miocene of central Japan (Fejfar et al. 1998, Tomida 2011). Tomida (2011) noted that *M. repenningi* resembles contemporaneous European *M. lavocati* in occlusal morphology, but the former species is more basal in having a distal root unsplit. According to Mörs and Flink (2018), *M. repenningi* is similar in size to a second and larger European species, *M. lindsayi*, which is morphologically

similar to *M. lavocati*. The single North American species of *Megapeomys*, *M. bobwilsoni*, is larger and more hypsodont, and the distal root is completely split into two separate roots.

### Genus *Apeomys* FAHLBUSCH, 1968

**Comments.** In Asia, a high-crowned P4 specimen was discovered in 1994 from the late early Miocene Xiacaowan Formation in Sihong, Jiansu Province, central East China. It was recently identified as the only Asian species of *Apeomys*, *A. asiaticus* QIU, 2017 (Text-fig. 2, Appendix). Compared to European *A. tuerkheimae* and *A. oldrichi*, Asian *A. asiaticus* has more derived dental features such as the anteroloph fused with the protoloph, absence of the mesoloph, and the closed syncline III (Qiu 2017).

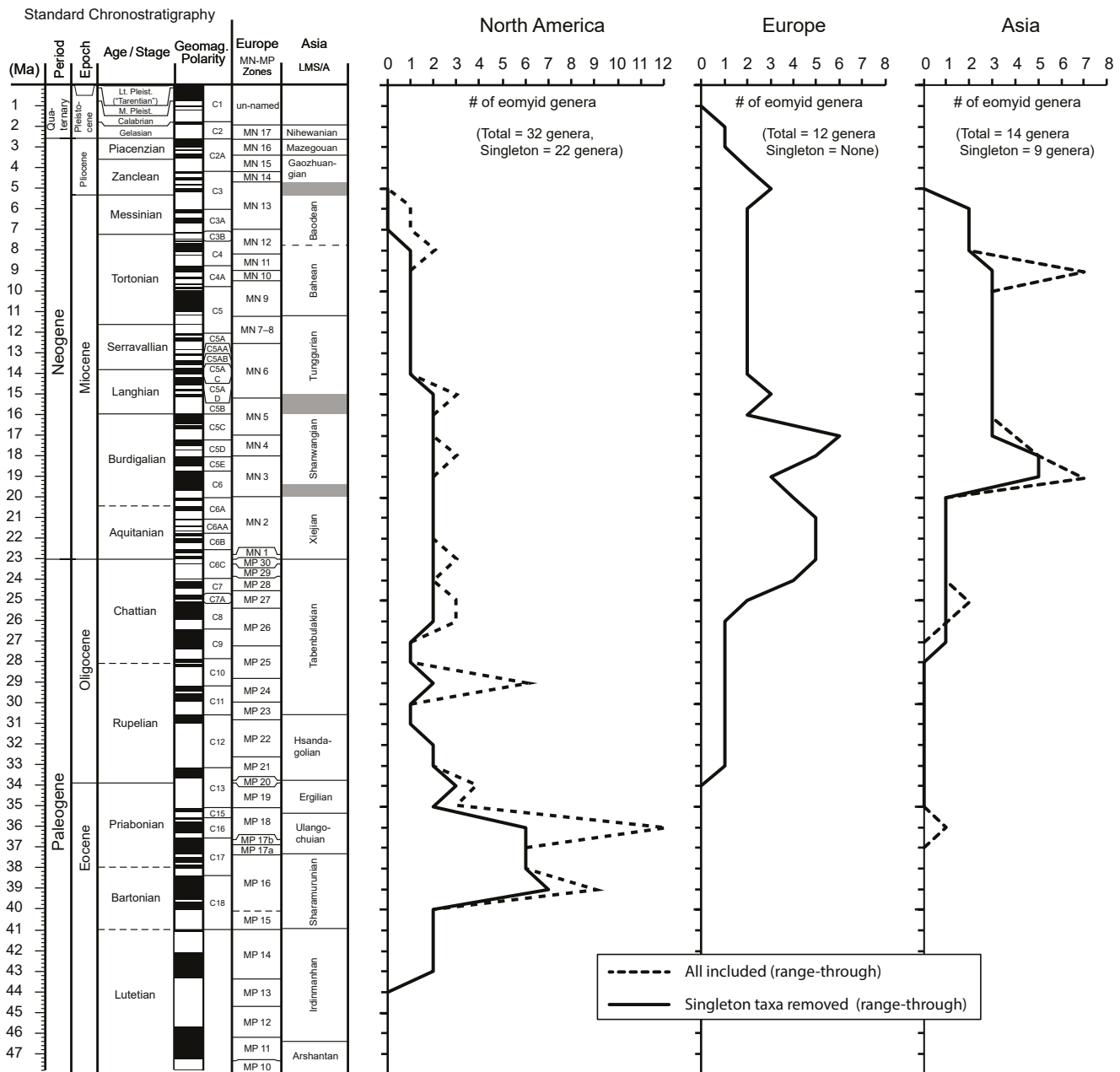
### Biogeographic significance of Asian eomyids

Over the past two decades, more and more eomyid fossils have been discovered in East Asia and the Asian part of Russia. To date, 22 species within 14 genera were reported from Asia, including seven endemic genera and rare material of apeomyine eomyids from southwestern China and Japan (Text-fig. 2). The total number of eomyid genera in Asia is now equivalent to that in Europe. For most of the endemic genera, phylogenetic relationships and their evolutionary history are of high interest but remain uncertain. On the other hand, the other seven non-endemic genera emphasize biogeographic importance of Asia in intercontinental dispersal events. Interestingly, four of these genera (*Pentabuneomys*, *Leptodontomys/Eomyops*, *Apeomys*, *Megapeomys*) show Holarctic distributions. Based on chronological occurrence and dental morphology, *Pentabuneomys* and *Leptodontomys/Eomyops* show a unidirectional migration, dispersing from North America into Eurasia probably by the early Miocene. Apeomyine eomyids likely had higher dispersal ability, showing at least two intercontinental dispersal events. *Apeomys* most likely originated in North America and then migrated into Eurasia during the late Oligocene given that two more basal apeomyine eomyids (*Zophoapeomys*, *Proapeomys*) predate *Apeomys whistleri* in North America (Korth 2008, Korth and Samuels 2015). Later, the apeomyine *Megapeomys* dispersed into North America, as suggested by the occurrence of a more basal species of *Megapeomys* in Japan than in North America (Tomida 2011) and the presence of more derived apeomyine taxa (*Apeomyoides savagei*) in North America (Korth 2008, Korth and Samuels 2015). The remaining three genera of the Asian eomyids are *Eomys*, *Keramidomys*, and *Ligerimys*, all of which occurred in Europe as well. In Eurasia, *Eomys* represents the oldest-known eomyid genus (locality Möhren 19, Germany, MP 21; locality Ronzon, France, MP 21, earliest Oligocene; Maridet et al. 2010) and is therefore associated with the first dispersal event of eomyid rodents from North America into Europe (Fahlbusch 1973, 1979, Engesser 1999). In Asia, scarce material of *E. orientalis* has been recovered from the early to late Oligocene of Central Mongolia and Inner Mongolia, China (Wang and Emry 1991, Maridet et al. 2015). More evidence of early dispersals of eomyid rodents is expected from these areas. Given that recent phylogenetic analysis indicates that

*Keramidomys* is related to the Asian endemics *Asianeomys* and *Japaneomys* (Kimura et al. 2019), *Keramidomys* likely originated in Asia and later dispersed westward. Asian *Ligerimys* is unique compared to other eomyids because its geographic range was mainly limited to western Europe before the discovery of early Miocene *L. asiaticus* (MN 2 – MN 3) in Inner Mongolia (Qiu and Li 2016). The occurrence of Asian *Ligerimys* suggests that this genus opportunistically dispersed eastward from Europe, seemingly without leaving diverse local descendants.

### Eomyid diversity through time with emphasis on Asia

Eomyid rodents originated in North America during the middle Eocene (Korth 1994) and then dispersed into Asia during the late Eocene (Emry et al. 1997) and into Europe by the early Oligocene (Engesser 1999). They were remarkably successful and widely distributed in the Holarctic but disappeared by the end of the Miocene in North America and by the early Pleistocene in Eurasia (Engesser 1999, Flynn 2008). Text-fig. 10 compares the occurrences of eomyid genera in the fossil record of North America, Europe and Asia (Tab. 1). After the earliest occurrence of eomyid rodents in the middle Eocene (~43.3 Ma) of North America, eomyid rodent diversity steadily increased and achieved a high plateau by 39 Ma, lasting for ~3 million years. The increased diversity occurred during progressive global cooling in the late Eocene (Zachos et al. 2001). Taking singleton taxa into consideration, the highest peak of eomyid diversity occurred in the late Eocene (~36 Ma) (updated in this study; see also Korth 1994, Flynn 2008). Even though eomyids dispersed from North America during the late Eocene (Asia) and earliest Oligocene (Europe), their diversity was initially low outside North America (Text-fig. 10). In North America, eomyid diversity progressively declined across the Eocene–Oligocene boundary, coinciding with the onset of Antarctic glaciation and the first occurrence of permanent ice-sheets (Zachos et al. 2001). This was a time of major turnover events in mammal faunas, the so-called “Grande Coupure” (= the Great Cut) in Europe and Mongolian Remodelling in Asia (see Meng and McKenna 1988, Costa et al. 2011). Eomyid diversity continuously decreased and reached a low level in the early Oligocene, but it slightly bounced back in the late early Oligocene, counting singleton taxa. However, North American eomyids hit their lowest diversity during the major expansion of the Antarctic glaciation (i.e., Oi-2b glacial event) in the late Oligocene. Only a single genus, the ubiquitous *Leptodontomys*, survived the cooling event. After the Oi-2a and Oi-2b glaciation events at 28 Ma – 26.7 Ma (Wade and Pälike 2004), eomyid diversity in North America remained at a low level and did not recover even during the mid-Miocene Climatic Optimum (17 Ma to 15 Ma), whereas eomyids flourished both in Europe and Asia at that time (Text-fig. 10). They were apparently diverse and successful in the warm (tropical to subtropical) dense forest environments in the late Eocene of North America, and their rapid decline during the Oligocene coincides with the expansion of open habitats including dry woodlands and grasslands (Retallack 2001, Strömberg 2011).



**Text-fig. 10.** Diversity of eomyid rodents in North America, Europe, and Asia, counted as “range-through” occurrences of eomyid genera. Two datasets were used to generate the diversity curves. All taxa were included in one dataset (solid line), and in the other dataset (dashed line), singleton taxa, were removed (any genera whose first and last occurrences are restricted to a single time bin; Sepkoski 1996). In each time bin, a data point is placed in its median age. References for standard chronostratigraphy and mammalian biostratigraphy follow Text-fig. 2.

In Eurasia, eomyid rodents diversified somewhat but were never as diverse as the late Eocene taxa of North America. They diversified sufficiently after their diversification and subsequent decline in North America. A singleton genus, *Symplokeomys*, from Kazakhstan is the only Eurasian representative contemporaneous with the highest eomyid diversity in North America (Text-figs 2, 10). *Eomys* is the earliest taxon to appear more persistently in the fossil record of Eurasia. Although the genus is more commonly known from the Oligocene of Europe (e.g., Comte and Vianey-Liaud 1987, Engesser 1990, Maridet et al. 2010), *Eomys* fossils, including those under provisional taxonomic assignments, have been reported from Mongolia (Daxner-Höck 2010, Maridet et al. 2015) in addition to the

type locality of the only Asian species, *E. orientalis*, from Inner Mongolia (Wang and Emry 1991) (see Appendix). Eomyid rodents experienced their highest generic diversity during the middle to late Burdigalian stage in Eurasia. A remarkable peak is observed in Europe at 17 Ma (Engesser 1999), which corresponds to the beginning of the mid-Miocene Climatic Optimum (Zachos et al. 2001), whereas a slight increase of generic diversity predated the warming event in Asia (Text-fig. 10). In Europe, the subsequent decline in eomyid diversity coincides with the end of the mid-Miocene Climatic Optimum (Zachos et al. 2001), but complex biotic factors may also have played an important role. Particularly, “modern” cricetid rodents like *Democricetodon* and *Megacricetodon* dispersed into Europe

at ~18 Ma and soon became major components of rodent faunas (Daams et al. 1988, Aguilar et al. 1999). These early cricetids, showing a generalized bunolophodont dentition, are considered to have been opportunists adapted to diverse environments including more arid habitats (Gómez Cano et al. 2017); however, they may also have competed with eomyids in more forested environments.

On the other hand, Asian eomyid diversity remained rather static during most of the Miocene, especially after the middle Burdigalian diversity peak (Text-fig. 10). The genera present in this interval are the long-ranging *Asianeomys*, *Keramidomys*, and *Leptodontomys*, which are well-represented in the intensively surveyed sections of Inner Mongolia (e.g., Qiu and Li 2016). Ecometric studies show that central East Asia was consistently more arid than Southern Europe throughout the Neogene (Eronen et al. 2010, Fortelius et al. 2014), which could have limited eomyid diversity.

By the end of the Pliocene, eomyid rodents disappeared in North America and Asia, whereas they survived until the early Pleistocene in northern Europe. Global climatic cooling resulted in the extension of grass-dominated ecosystems in the late Miocene (Quade and Cerling 1995, Cerling et al. 1997, Tipple and Pagani 2007), whereas in Europe, warm-temperate forest environments persisted in many areas and at higher latitudes although arid landscapes existed in the Mediterranean regions since the middle Miocene (Jiménez-Moreno and Suc 2007, but see Quan et al. 2014). The persistent warm-temperate forest environments may explain the much later extinction of eomyids in Europe. In Asia, even after extensive field efforts, no eomyid fossils have been found in Pliocene localities in Inner Mongolia, China (Qiu and Storch 2000). It seems reasonable that the earlier regional disappearance is a true signal in Inner Mongolia due to the ongoing aridification in central Asia linked to the intensification of the monsoon climate (e.g., An et al. 2001, Liu and Yin 2002, Biasati et al. 2010). However, considering the discovery of late Miocene eomyids (Qiu 2006, Qiu 2017) from Yunnan Province (southwestern China), it is possible that eomyid refuges existed during a certain time period in southern China or continental Southeast Asia. The exploration of new regions and the intensive sampling of known fossiliferous areas may eventually test such a hypothesis.

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## Appendix

Species occurrences of all eomyid rodents in Asia. Each entry represents a species from a single locality.

Entry	Scientific Name	Country	Province/Region/Prefecture	County/City/Town/Geographic Region)	Fossil Locality	Lithostratigraphic Unit	Upper boundary		Lower boundary		References
							Biosтратigraphic Zone	Regional Biozone	Age (Ma)	Rationale for the age assignment	
1	<i>Aprozomys astutus</i>	China	Jiangsu	Shiogang	Zhengji	Xiaocawayan Formation	Shanwangjian		14.2	Upper boundary of MN 5	Shanwangjian
2	<i>Asiameomys astutus</i>	China	Inner Mongolia	Saint Jacques	IVPP Loc. 77048	Wulangula Formation			23.0	Upper boundary of Chattian (= the Oligocene/Miocene boundary)	19.5 MN 3
3	<i>Asiameomys boligeri</i>	Kazakhstan	Aktyubinsk	Chelkar (North Aral Region)	Altynshoksu	Aral Formation	correlated to Central Mongolian local biozone C	Upper boundary of Central Mongolian local biozone C	25.6	correlated to Central Mongolian local biozone C	Lopatin (2000); Bendukidze (2009); Daxner-Höck et al. (2017) for correlated age
4	<i>Asiameomys dangheensis</i>	China	Xinjiang	(Junggar Basin)	X199005	Suoququan Formation	Xiejian	Suoququan mammal assemblage zone II	21.7	Magnetostratigraphy	27.4 Lower boundary of Central Mongolian local biozone C
5	<i>Asiameomys dangheensis</i>	Mongolia	(Valley of Lakes)	Del	Loh Formation	local biozone C1	Magnetostratigraphy, faunal comparison with biozone C1	local biozone C1	24.2	Magnetostratigraphy, faunal comparison with biozone C1	21.9 Suoququan mammal assemblage zone II
6	<i>Asiameomys dangheensis</i>	Mongolia	(Valley of Lakes)	Huch Teeg	Loh Formation	local biozone C1	Magnetostratigraphy, faunal comparison with biozone C1	local biozone C1	24.2	Magnetostratigraphy, faunal comparison with biozone C1	25.6
7	<i>Asiameomys dangheensis</i>	Mongolia	(Valley of Lakes)	Tatal Gol	Loh Formation	local biozone D	Magnetostratigraphy, faunal comparison with biozone C1	local biozone D	24.2	Magnetostratigraphy, faunal comparison with biozone C1	25.6
8	<i>Asiameomys dangheensis</i>	Mongolia	(Valley of Lakes)	Urkhetseg	Loh Formation	local biozone D	Faunal comparison with biozone D (the occurrence of <i>Democricetodon sui</i> )	local biozone D	21.0	Faunal comparison with biozone D (the occurrence of <i>Democricetodon sui</i> )	23.0
9	<i>Asiameomys dangheensis</i>	Mongolia	(Valley of Lakes)	Lunuy Yas	Loh Formation	local biozone D	Faunal comparison with biozone D (the occurrence of <i>Democricetodon sui</i> )	local biozone D	21.0	Faunal comparison with biozone D (the occurrence of <i>Democricetodon sui</i> )	23.0
10	<i>Asiameomys dangheensis</i>	Mongolia	(Valley of Lakes)	Houlin Teeg	Loh Formation	local biozone D	Faunal comparison with biozone D (the occurrence of <i>Democricetodon sui</i> )	local biozone D	21.0	Faunal comparison with biozone D (the occurrence of <i>Democricetodon sui</i> )	23.0
11	<i>Asiameomys dangheensis</i>	China	Aksay Kazak Autonomous County, Gansu	Yandantu, Minzhu Xiang	IVPP Loc. DH199904	Paoniuhuan Formation	Tabenbulukian		23.0	Upper boundary of Tabenbulukian	28.4
12	<i>Asiameomys engesseri</i>	China	Xinjiang	(Junggar Basin)	X199005	Suoququan Formation	Xiejian	Suoququan mammal assemblage zone II	21.7	Magnetostratigraphy	21.9 Suoququan mammal assemblage zone II
13	<i>Asiameomys fahibuschi</i>	China	Inner Mongolia	Sonid Zuoqi	Lower Aoerhan	Lower Red Mudstone Member, Aoerhan Formation	Xiejian (correlated to MN 3)		17.2	Upper boundary of MN 3	21.7
14	<i>Asiameomys fahibuschi</i>	China	Inner Mongolia	Sonid Zuoqi	Gashunyinadege bed	Shanwangjian (correlated to MN 3)	Xiejian (correlated to MN 3)		17.2	Xiejian (correlated to MN 3)	19.5 Lower boundary of MN 3
15	<i>Asiameomys fahibuschi</i>	China	Xinjiang	(Junggar Basin)	X199005	Suoququan Formation	Xiejian	Suoququan mammal assemblage zone II	21.7	Magnetostratigraphy	21.9 Suoququan mammal assemblage zone II
16	<i>Asiameomys jungarensis</i>	China	Xinjiang	Tierishababa (Junggar Basin)	X198035	Tierishababa Formation	Tabenbulukian	Tierishababa mammal assemblage zone I	23.2	Magnetostratigraphy	24.4 Magnetostratigraphy
17	<i>Asiameomys jungarensis</i>	China	Xinjiang	Tierishababa (Junggar Basin)	X198024	Tierishababa Formation	Tabenbulukian	Tierishababa mammal assemblage zone I	23.2	Magnetostratigraphy	24.4 Magnetostratigraphy
18	<i>Asiameomys jungarensis</i>	China	Xinjiang	Tierishababa (Junggar Basin)	XJ200209	Tierishababa Formation	Tabenbulukian	Tierishababa mammal assemblage zone I	23.2	Magnetostratigraphy	24.4 Magnetostratigraphy

Entry	Scientific Name	Country	Province/Region/Prefecture	County/City/Geographic Region)	Fossil Locality	Lithostratigraphic Unit	Upper boundary			Lower boundary			References	
							Biostratigraphic Zone	Regional Biozone	Age assignment (Ma)	Biostratigraphic Zone	Regional Biozone	Age (Ma)		
19	<i>Asiameomys yunnanensis</i>	Kazakhstan	Aktubinsk	Chelkar (North Aral Region)	Altynshoksu	Aral Formation	correlated Central Mongolian local biozone C	25.6	Upper boundary of Central Mongolian local biozone C	correlated Central Mongolian local biozone C	27.4	Lower boundary of Central Mongolian local biozone C	Lopatin (2000); Bendukdzee (2009); Daxner-Höck et al. (2017)	
20	<i>Asiameomys</i> sp.	China	Xinjiang	(Junggar Basin)	XJ99005	Suo suoquan Formation	Xiejian	Suo suoquan mammal assemblage zone II	21.7	Magnetostratigraphy	Xiejian	Suo suoquan mammal assemblage zone II	Meng et al. (2006); Wu et al. (2006); Bi et al. (2013)	
21	<i>Asiameomys</i> sp.	China	Inner Mongolia	Ulanatal	UTL6	Ulanatal Formation			24.2	Upper boundary of Central Mongolian local biozone C1	correlated to Mongolia local biozone C1	25.6	Lower boundary of Central Mongolian local biozone C1	Gomes Rodrigues et al. (2014) and references therein; Daxner-Höck et al. (2017)
22	<i>Asiameomys aff. engesserri</i>	China	Xinjiang	Burqin, Habae (Junggar Basin)	XJ 200604				18.05	Taxonomic comparison with the Sihong locality (the occurrence of <i>Microdromys orientalis</i> )		21.9	Taxonomic comparison with the Suosuoquan mammal assemblage zone II (the occurrence of <i>Asiameomys engesserri</i> )	Maridet et al. (2011)
23	<i>Asiameomys</i> sp.	China	Xinjiang	Burqin, Habae (Junggar Basin)	XJ 200604				18.05	Taxonomic comparison with the Sihong locality (the occurrence of <i>Microdromys orientalis</i> )		21.9	Taxonomic comparison with the Suosuoquan mammal assemblage zone II (the occurrence of <i>Asiameomys engesserri</i> )	Maridet et al. (2011)
24	cf. <i>Asiameomys</i> Mongolia <i>bolidigeri</i>	Mongolia		(Valley of Lakes)	Taatsiin Gol (right)	Loh Formation	local biozone C	25.6	Upper boundary of Central Mongolian local biozone C	local biozone C	27.4	Lower boundary of Central Mongolian local biozone C	Maridet et al. (2015)	
25	cf. <i>Asiameomys</i> Mongolia <i>bolidigeri</i>	Mongolia		(Valley of Lakes)	Taatsiin Gol (west)	Loh Formation	local biozone C	25.6	Upper boundary of Central Mongolian local biozone C	local biozone C	27.4	Lower boundary of Central Mongolian local biozone C	Maridet et al. (2015)	
26	<i>Eomyodon</i> sp.	China	Inner Mongolia	Qianlishan District	IVPP Loc. 78018	Wulangulage Formation			23.0	Upper boundary of Chattian (= the Oligocene/Miocene boundary)		27.8	Lower boundary of Chattian	Wang and Emry (1991)
27	" <i>Eomyops opuligerae</i> "	Russia	Siberia	Olkhon Island, Lake Baikal	Tagay				-13	Taxonomic comparison (roughly correlated to MN 7+8)		-13	Taxonomic comparison (roughly correlated to MN 7+8)	Daxner-Höck et al. (2013)
28	<i>Eomyops/Leptodontomys</i> sp.	Mongolia	(Valley of Lakes)	Ulaan To goi	Loh Formation		local biozone D1/2	13.0	<sup>40</sup> Ar/ <sup>39</sup> Ar age on basalt III layer	local biozone D1/2	15.97	early/middle Miocene boundary	Maridet et al. (2015)	
29	<i>Eomyops</i> <i>orientalis</i>	China	Inner Mongolia	Saint Jacques	IVPP Loc. 77049.2	Wulangulage Formation			23.0	Upper boundary of Chattian (= the Oligocene/Miocene boundary)		27.8	Lower boundary of Chattian	Wang and Emry (1991)
30	<i>Eomyops</i> <i>orientalis</i>	China	Inner Mongolia	Qianlishan District	IVPP Loc. 78018	Wulangulage Formation			23.0	Upper boundary of Chattian (= the Oligocene/Miocene boundary)		27.8	Lower boundary of Chattian	Wang and Emry (1991)
31	<i>Eomyops</i> aff. <i>orientalis</i>	Mongolia	(Valley of Lakes)	Taatsiin Gol (right)	Hasanda Gol Formation		local biozone B	28.0	<sup>40</sup> Ar/ <sup>39</sup> Ar age on basalt II layer	local biozone B	31.5	<sup>40</sup> Ar/ <sup>39</sup> Ar age on basalt II layer	Maridet et al. (2015)	
32	<i>Eomyops</i> aff. <i>orientalis</i>	Mongolia	(Valley of Lakes)	Hasanda Gol Formation	Hasanda Gol Formation		local biozone B	28.0	<sup>40</sup> Ar/ <sup>39</sup> Ar age on basalt II layer	local biozone B	31.5	<sup>40</sup> Ar/ <sup>39</sup> Ar age on basalt II layer	Maridet et al. (2015)	
33	<i>Eomyops</i> ct. <i>orientalis</i>	Mongolia	(Valley of Lakes)	Taatsiin Gol	Hasanda Gol Formation		local biozone A	31.5	<sup>40</sup> Ar/ <sup>39</sup> Ar age on basalt II layer	local biozone A	33.9	Oligocene boundary	Maridet et al. (2015)	
34	<i>Eomyops</i> sp.	Mongolia	(Valley of Lakes)	Taatsiin Gol (right)	Hasanda Gol Formation		local biozone B	28.0	<sup>40</sup> Ar/ <sup>39</sup> Ar age on basalt II layer	local biozone B	31.5	not well dated	Maridet et al. (2015)	
35	<i>Heteromys yunnanensis</i>	China	Yunnan	Lufeng	Shihuihua	Shihuihua Formation			not well dated	not well dated		not well dated	well biostatigraphic comparison with Siwalk rhizomyid rodents by Flynn and Qi (1982) but not well dated	Flynn and Qi (1982); Ni and Qiu (2002); Qiu (2006)
36	<i>Heteromys yunnanensis</i>	China	Yunnan	Yuanmou	Leilaotao	Xiaohao Formation			not well dated	not well dated		not well dated	well biostatigraphic comparison with Siwalk rhizomyid rodents by Flynn and Qi (1982) but not well dated	Flynn and Qi (1982); Ni and Qiu (2002); Qiu (2006)
37	<i>Japanomeomys yasumoti</i>	Japan	Gifu	Kani	Data	Nakanura Formation			17.2	Upper boundary of MN 3		19.8	K-Ar dating in upper Hachiya Formation	Kimura et al. (2019)

Entry	Scientific Name	Country	Province/Prefecture	County/City/Town (Geographic Region)	Fossil Locality	Lithostratigraphic Unit	Biostratigraphic Zone	Upper boundary		Lower boundary		References
								Age (Ma)	Rationale for the age assignment	Regional Biozone	Biostratigraphic Zone	
38	<i>Keranidomys fahibuschi</i>	China	Inner Mongolia	Sonid Zuqi	Lower Aoerban	Lower Red Mudstone Member, Aoerban Formation	Xiejian (correlated to MN 3)	17.2	Upper boundary of MN 3	Xiejian (correlated to MN 2)	21.7	Lower boundary of MN 2
39	<i>Keranidomys fahibuschi</i>	China	Inner Mongolia	Sonid Zuqi	Gashunyinadege bed	Shanwangjian (correlated to MN 3)		17.2	Upper boundary of MN 3	Xiejian (correlated to MN 3)	19.5	Lower boundary of MN 3
40	<i>Keranidomys fahibuschi</i>	China	Inner Mongolia	Sonid Zuqi	Upper Aoerban	Upper Red Mudstone Member, Aoerban Formation	Shanwangjian (correlated to MN 5)	14.2	Upper boundary of MN 5	Shanwangjian (correlated to MN 3)	19.5	Lower boundary of MN 3
41	<i>Keranidomys fahibuschi</i>	China	Inner Mongolia	Sonid Youqi	H-X Highway Road original mark 346	Tunggur Formation		11.2	Upper boundary of MN 7/8	Tunggurian (correlated to MN 7/8)	12.9	Lower boundary of MN 7/8
42	<i>Keranidomys fahibuschi</i>	China	Inner Mongolia	Sonid Zuqi	Balunhalagen	Balunhalagen bed	Bahean (correlated to MN 11)	7.6	Upper boundary of MN 11	Bahean (correlated to MN 9)	11.2	Lower boundary of MN 9
43	<i>Keranidomys fahibuschi</i>	China	Inner Mongolia	Sonid Zuqi	Biitu	Biitu bed	Baodean (correlated to MN 13)	5.3	Upper boundary of MN 13	Baodean (correlated to MN 12)	7.6	Lower boundary of MN 12
44	<i>Keranidomys magnus</i>	China	Inner Mongolia	Sonid Youqi	Arnuwusu	Arnuwusu bed	Bahean (correlated to MN 9)	9.9	Upper boundary of MN 9	Bahean (correlated to MN 9)	11.2	Lower boundary of MN 9
45	<i>Keranidomys magnus</i>	China	Inner Mongolia	Sonid Youqi	Shala	Shala bed	Bahean (correlated to MN 11)	7.6	Upper boundary of MN 11	Bahean (correlated to MN 10)	9.9	Lower boundary of MN 10
46	<i>Keranidomys magnus</i>	China	Inner Mongolia	Sonid Zuqi	Balunhalagen	Balunhalagen bed	Bahean (correlated to MN 11)	7.6	Upper boundary of MN 11	Bahean (correlated to MN 9)	11.2	Lower boundary of MN 9
47	<i>Keranidomys magnus</i>	China	Inner Mongolia	Sonid Zuqi	Biitu	Biitu bed	Bahean (correlated to MN 13)	5.3	Upper boundary of MN 13	Baodean (correlated to MN 12)	7.4	Lower boundary of MN 12
48	<i>Keranidomys aff. mohleri / fahibuschi</i>	Russia	Siberia	Olkhon Island, Lake Baikal	Tagay						~13	Taxonomic comparison (roughly correlated to MN 7+8)
49	<i>Keranidomys sp.</i>	Mongolia	(Valley of Lakes)	Ulaan Tolgoi	Loh Formation		local biozone D1/2	13.0	<sup>40</sup> Ar/ <sup>39</sup> Ar age on basalt III layer			Taxonomic comparison (roughly correlated to MN 7+8)
50	<i>Keranidomys</i> sp.	China	Xinjiang	Burjin, Habale (Junggar Basin)	XJ 200604					Taxonomic comparison with the Sihong locality ( <i>Microtromys orientalis</i> )	15.97	Taxonomic comparison (roughly correlated to MN 7+8)
51	<i>Leptadontomys gansus</i>	China	Inner Mongolia	Sonid Zuqi	Gashunyinadege bed	Shanwangjian (correlated to MN 3)					21.9	Taxonomic comparison with the Suosuoqian mammal assemblage zone II (the occurrence of <i>Asiaceomys engesserii</i> )
52	<i>Leptadontomys gansus</i>	China	Inner Mongolia	Sonid Youqi	H-X Highway Road original mark 346	Tunggur Formation (correlated to MN 7/8)		17.2	Upper boundary of MN 3	Xiejian (correlated to MN 3)	19.5	Lower boundary of MN 3
53	<i>Leptadontomys gansus</i>	China	Inner Mongolia	Sonid Zuqi	Balunhalagen	Balunhalagen bed	Bahean (correlated to MN 11)	11.2	Upper boundary of MN 7/8	Tunggurian (correlated to MN 7/8)	12.85	Lower boundary of MN 7/8
54	<i>Leptadontomys gansus</i>	Inner Mongolia	Sonid Youqi	Arnuwusu	Arnuwusu bed	Bahean (correlated to MN 9)						Lower boundary of MN 9
55	<i>Leptadontomys gansus</i>	Inner Mongolia	Abag	Huitenghe	Huitenghe bed	early Baodean (correlated to M 11)		7.6	Upper boundary of MN 11	Bahean (correlated to MN 10)	11.2	Lower boundary of MN 10
56	<i>Leptadontomys gansus</i>	Inner Mongolia	Sonid Youqi	Shala	Shala bed	Bahean (correlated to MN 11)		7.6	Upper boundary of MN 11	Bahean (correlated to MN 9)	11.2	Lower boundary of MN 9
57	<i>Leptadontomys gansus</i>	Inner Mongolia	Sonid Zuqi	Biitu	Biitu bed	Baodean (correlated to MN 13)		5.3	Upper boundary of MN 13	Baodean (correlated to MN 12)	7.4	Lower boundary of MN 12
58	<i>Leptadontomys gansus</i>	Inner Mongolia	Huade	Eritmete2	Eritmete Formation (correlated to MN 13)			5.3	Upper boundary of MN 13	Baodean (correlated to MN 13)	7.1	Upper boundary of MN 13
59	<i>Leptadontomys gansus</i>	Inner Mongolia	Huade	Hair Obo2	correlated to Eritmete Formation (correlated to MN 13)			5.3	Upper boundary of MN 13	Baodean (correlated to MN 13)	7.1	Upper boundary of MN 13

Entry	Scientific Name	Country	Province/Region/Prefecture	County/City/Town (Geographic Region)	Fossil Locality	Lithostratigraphic Unit	Biostratigraphic Zone			Upper boundary			Lower boundary			References
							Regional Biozone	Age assignment (Ma)	Rationale for the age assignment	Biostatigraphic Zone	Regional Biozone	Age (Ma)	Rationale for the age assignment			
60	<i>Leptidontomys ganensis</i>	China	Gansu	Songshan-Tianzu Xian	Loc. 1	Baodean (correlated to MN 13)	5.3	Upper boundary of MN 13 (correlated to MN 12)	Baodean	Xiejian (correlated to MN 3)	7.4	Lower boundary of MN 12	Zheng and Li (1982); Deng (2006)			
61	<i>Leptidontomys ili</i>	China	Inner Mongolia	Sonid Zuoqi	Lower Aoerban	Lower Red Mudstone Member, Aoerban Formation	17.2	Upper boundary of MN 3 (correlated to MN 2)	Xiejian (correlated to MN 3)	Tungurian (correlated to MN 3)	21.7	Lower boundary of MN 2	Qiu et al. (2013); Qiu and Li (2016)			
62	<i>Leptidontomys ili</i>	China	Inner Mongolia	Sonid Zuoqi	Gashunyinadege bed	Shanwangian (correlated to MN 3)	17.2	Upper boundary of MN 3 (correlated to MN 3)	Xiejian (correlated to MN 3)	Tungurian (correlated to MN 3)	19.5	Lower boundary of MN 3	Qiu et al. (2013); Qiu and Li (2016)			
63	<i>Leptidontomys ili</i>	China	Inner Mongolia	Sonid Youqi	H-X Highway Road original mark 346	Tungur Formation (correlated to MN 7/8)	11.2	Upper boundary of MN 7/8	Bahean (correlated to MN 7/8)	Tungurian (correlated to MN 7/8)	12.85	Lower boundary of MN 7/8	Qiu et al. (2013); Qiu and Li (2016)			
64	<i>Leptidontomys ili</i>	China	Inner Mongolia	Sonid Youqi	Anmuusu bed	Bahean (correlated to MN 9)	9.9	Upper boundary of MN 9 (correlated to MN 9)	Bahean (correlated to MN 9)	Bahean (correlated to MN 9)	11.2	Lower boundary of MN 9	Qiu et al. (2013); Qiu and Li (2016)			
65	<i>Leptidontomys ili</i>	China	Inner Mongolia	Sonid Zuoqi	Balunhalagen bed	Balunhalagen bed (correlated to MN 11)	7.6	Upper boundary of MN 11 (correlated to MN 9)	Bahean (correlated to MN 9)	Bahean (correlated to MN 9)	11.2	Lower boundary of MN 9	Qiu et al. (2013); Qiu and Li (2016)			
66	<i>Leptidontomys ili</i>	China	Inner Mongolia	Abag	Huitenghe bed	Huitenghe bed (correlated to MN 11)	7.6	Upper boundary of MN 11 (correlated to MN 10)	Baodean (correlated to MN 11)	Bahean (correlated to MN 10)	9.9	Lower boundary of MN 10	Qiu et al. (2013); Qiu and Li (2016)			
67	<i>Leptidontomys ili</i>	China	Inner Mongolia	Sonid Yeqi	Shala	Shala bed (correlated to MN 11)	7.6	Upper boundary of MN 11 (correlated to MN 10)	Baodean (correlated to MN 11)	Bahean (correlated to MN 10)	9.9	Lower boundary of MN 10	Qiu et al. (2013); Qiu and Li (2016)			
68	<i>Leptidontomys ili</i>	China	Inner Mongolia	Sonid Zuoqi	Bilutu bed	Bilutu bed (correlated to MN 13)	5.3	Upper boundary of MN 13 (correlated to MN 12)	Baodean (correlated to MN 13)	Baodean (correlated to MN 12)	7.4	Lower boundary of MN 12	Qiu et al. (2013); Qiu and Li (2016)			
69	<i>Ligerimys asiaticus</i> sp.	China	Yunnan	Yuannou	Loc. 9905	not well dated			not biostratigraphic well dated comparison with Siwalik rhizomyid rodents by Flynn and Qi (1982) but not well dated	not well dated	not well dated		not biostratigraphic well comparison with dated Siwalik rhizomyid rodents by Flynn and Qi (1982) but not well dated	Flynn and Qi (1982); Ni and Qiu (2002); Qiu (2006, 2017)		
70	<i>Ligerimys asiaticus</i>	China	Inner Mongolia	Sonid Zuoqi	Lower Aoerban	Lower Red Mudstone Member, Aoerban Formation	17.2	Upper boundary of MN 3 (correlated to MN 3)	Xiejian (correlated to MN 3)	Xiejian (correlated to MN 2)	21.7	Lower boundary of MN 2	Qiu et al. (2013); Qiu and Li (2016)			
71	<i>Ligerimys asiaticus</i>	China	Inner Mongolia	Sonid Zuoqi	Gashunyinadege bed	Shanwangian (correlated to MN 3)	17.2	Upper boundary of MN 3 (correlated to MN 3)	Xiejian (correlated to MN 3)	Xiejian (correlated to MN 3)	19.5	Lower boundary of MN 3	Qiu et al. (2013); Qiu and Li (2016)			
72	<i>Megapeconymus trapezumius</i>	Japan	Giftu	Kani	Data	Nakanura Formation	17.2	Upper boundary of MN 3 (correlated to MN 3)					K-Ar dating in Takeuchi (1992), Tomida (2011)			
73	<i>Omboomys buitynensis</i>	Mongolia	(Valley of Lakes)	Bulstyn Khudag	Tuyu Gol Formation	local biozone E	5.3	Miocene/Pliocene boundary					$^{40}\text{Ar}/^{39}\text{Ar}$ dating in local biozone E	Marinet et al. (2015)		
74	<i>Pentabacconomys jeffari</i>	China	Inner Mongolia	Sonid Zuoqi	Lower Aoerban	Lower Red Mudstone Member, Aoerban Formation	17.2	Upper boundary of MN 3 (correlated to MN 3)	Xiejian (correlated to MN 3)	Xiejian (correlated to MN 2)	21.7	Lower boundary of MN 2	Qiu et al. (2013); Qiu and Li (2016)			
75	<i>Pentabacconomys jeffari</i>	China	Inner Mongolia	Sonid Zuoqi	Gashunyinadege bed	Shanwangian (correlated to MN 3)	17.2	Upper boundary of MN 3 (correlated to MN 3)	Xiejian (correlated to MN 3)	Xiejian (correlated to MN 3)	19.5	Lower boundary of MN 3	Qiu et al. (2013); Qiu and Li (2016)			
76	<i>Pentabacconomys jeffari</i>	China	Inner Mongolia	Sonid Zuoqi	Upper Aoerban	Shanwangian (correlated to MN 5)	14.2	Upper boundary of MN 5 (correlated to MN 5)	Shanwangian (correlated to MN 5)	Shanwangian (correlated to MN 5)	19.5	Lower boundary of MN 3	Qiu et al. (2013); Qiu and Li (2016)			
77	<i>Pentabacconomys jeffari</i>	China	Inner Mongolia	Sonid Youqi	Anmuusu bed	Bahean (correlated to MN 9)	9.9	Upper boundary of MN 9 (correlated to MN 9)	Bahean (correlated to MN 9)	Bahean (correlated to MN 9)	11.2	Lower boundary of MN 9	Qiu et al. (2013); Qiu and Li (2016)			
78	<i>Pentabacconomys jeffari</i>	China	Inner Mongolia	Sonid Zuoqi	Balunhalagen bed	Balunhalagen bed (correlated to MN 11)	7.6	Upper boundary of MN 11 (correlated to MN 9)	Bahean (correlated to MN 11)	Bahean (correlated to MN 9)	11.2	Lower boundary of MN 9	Qiu et al. (2013); Qiu and Li (2016)			
79	<i>Plestoleomys mirabilis</i>	China	Yunnan	Lufeng	Shihuihua	Shihuihua Formation	not well dated		not biostratigraphic well dated comparison with Siwalik rhizomyid rodents by Flynn and Qi (1982) but not well dated	not well dated	not well dated		not biostratigraphic well comparison with dated Siwalik rhizomyid rodents by Flynn and Qi (1982) but not well dated	Flynn and Qi (1982); Ni and Qiu (2002); Qiu (2006)		
80	<i>Symplocomys zyzanicus</i>	Kazakhstan	(Basin)	Sunduk (Zaysan Basin)	Loc. K10	Aksyr Formation	Ergilian	33.9	Upper boundary of Ergilian	Ergilian	Ergilian	37.2	Lower boundary of Ergilian	Emry et al. (1997); PBDB		
81	<i>Yaneomys pusillus</i>	China	Yunnan	Lufeng	Shihuihua	Shihuihua Formation	not well dated		not biostratigraphic well comparison with Siwalik rhizomyid rodents by Flynn and Qi (1982) but not well dated	not well dated	not well dated		not biostratigraphic well comparison with dated Siwalik rhizomyid rodents by Flynn and Qi (1982) but not well dated	Flynn and Qi (1982); Ni and Qiu (2002); Qiu (2006, 2017)		