New record of a haplocyonine amphicyonid in Early Miocene of Nei Mongol fills a long-suspected geographic hiatus

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Abstract We place on the record a newly discovered amphicyonid (beardogs) upper molar from the Early Miocene Lower Red Mudstone Member of Aoerban Formation in central Nei Mongol. This molar is highly diagnostic of European haplocyonine or North American temnocyonine, two subfamilies of beardogs that have long been known in those continents but notably absent in Asia. The new molar is strikingly similar to Haplocyonoides mordax and Temnocyon percussor with its dumbbell-shaped M1 outline, reduced parastyle, isolated protocone by a surrounding cingulum, and extreme reduction of pre- and postprotocristae. Given the limited material at hand, we tentatively refer the new Chinese fossil to the European Haplocyonoides cf. H. mordax because of their similar size and age relationship. If this identification is correct, our new record thus fills a large gap in the geographic distribution of the haplocyonines and represents an excursion of this rare subfamily from Europe.

Key words Aoerban, Nei Mongol; Early Miocene; Amphicyonidae; zoogeography


1 Introduction

Straddling between Europe to the west and North America to the east, Asia has long been suspected to be either a source of certain Cenozoic mammals that, thus far, are only found in Europe and/or North America, or at the very least, must have been a “jumping board” to and from the continents of destination as they disperse. When taxa are found in both Europe and North America, their absence in Asia thus fuels speculation of a Holarctic continuity,
and the expectation that filling in the Asian gap may be a matter of time. Temnocyonine and haplocyonine amphicyonids are such an example of a long-suspected geographic hiatus waiting to be filled.

Temnocyoninae and Haplocyoninae are peculiar lineages of amphicyonids, or beardogs, with hypercarnivorous dentitions and in some species, digitigrade posture (Hunt, 2011). Members of these groups typically have a very trenchant lower molar battery that align their main cusps in a single row to facilitate shearing function and associated narrow, high-crowned premolar series (e.g., Bonis, 1973; Hunt, 2011; Peigné and Heizmann, 2003). Asian records of amphicyonids have generally lagged behind those of Europe and North America, partly because of a shorter history of field explorations associated with a late start of scientific enterprise in almost all Asian countries. Occasionally, field records in Asia do catch up, and we have the satisfaction of reporting a new fossil record of the haplocyonines from the Early Miocene of Nei Mongol (Inner Mongolia) that confirms, once again, Asia has much to offer in our understanding of Holarctic zoogeography.

**Abbreviations**  ACM, Amherst College Museum of Natural History, Amherst, Massachusetts; AMNH, Division of Paleontology, American Museum of Natural History, New York; F:AM, Frick Collection, American Museum of Natural History, New York; IVPP, Institute of Vertebrate Paleontology and Paleoanthropology, Chinese Academy of Sciences, Beijing; IM, IVPP fossil localities from Nei Mongol; LACM, Natural History Museum of Los Angeles County, Los Angeles; NM, Northwest Museum, Portland State University, Portland, Oregon; SMF, Senckenberg Forschungsinstitut und Naturmuseum, Frankfurt; TRO, Timberlane Research Organization, Lake Wales, Florida; UCMP, Museum of Paleontology, University of California, Berkeley; UNSM, Vertebrate Paleontology, University of Nebraska State Museum, Lincoln; USNM, Department of Paleobiology, Smithsonian Institution, Washington, D.C.; YPM-PU, Princeton Collection, Peabody Museum, Yale University, New Haven.

2 Systematic paleontology

**Class Mammalia Linnaeus, 1758**

**Order Carnivora Bowdich, 1821**

**Family Amphicyonidae Haeckel, 1866**

**Subfamily Haplocyoninae Bonis, 1966**

**Genus Haplocyonoides Hürzeler, 1940**

*Haplocyonoides* cf. *H. mordax* Hürzeler, 1940


**Locality, stratigraphy, and age** IVPP IM1530 locality (XWIM04), N43°20’34’’, E113°54’30’’, elevation 1050 m, Sonid Zuoqi, Xilinhot League, central Nei Mongol (Fig. 1).
Locality IM1530 is stratigraphically straddling the boundary of the Lower Red Mudstone and Middle Green Mudstone members of Aoerban Formation (Fig. 2). Locally the two members transition into each other, possibly due to diagenetic effects of the green coloration. IVPP

Fig. 1 Location of Aoerban area in central Nei Mongol (upper) and IVPP IM1530 locality (red star) and updated geologic map of Aoerban strata (lower)
Modified from Wang et al. (2009). Note that distribution of the Bilutu bed (light grey) differs substantially from that of Wang et al. (2009:fig. 1).
V 22639 thus belongs to the upper-most Lower Aoerban Fauna (Wang et al., 2009). Based on biochronology of small mammals, Qiu et al. (2013a) estimated that the upper part of the Lower Aoerban Fauna was approximately 19 Ma in the late Xiejian or possibly early Shanwangian Chinese land mammal ages (see Qiu et al., 2013b).

![Photograph of the IVPP IM1530 locality (marked by geologic hammer) at a transitional zone between the Lower Red Mudstone and Middle Green Mudstone members of Aoerban Formation. Stratigraphic scheme follows that of Wang et al. (2009:fig. 3). Photo looking south.](image)

**Description**  With the exception of the missing roots, IVPP V 22639 is almost perfectly preserved, missing only the anterior rim of the enamels at the waist. Fresh breakage at the base of the roots indicates that the roots were broken after re-emergence from sediment encasement during the erosion. V 22639 shows signs of only minor wear at the apex of the metacone and thus represents that of a young adult.

V 22639 is almost twice as transversely wide as anteroposteriorly long (see measurements in Table 1). Overall it is dumbbell-shaped with a distinctly constricted waist at the junction of the protocone on the lingual side and paracone-metacone on the labial side. In occlusal view, paracone and metacone are about the same size, although the paracone is taller; this is especially true when viewed from the lingual side, but less so from the labial side. A parastyle is poorly developed, indicated by a slightly thickened and higher cingulum surrounding the parastyle. A distinct notch separates the paracone and metacone. Both paracone and metacone have an anterior and posterior ridge, which are aligned in a single line, and are completely encircled by a thin cingulum.

The protocone has a rounded base and located slightly anterior to the paracone-metacone notch. The low-crowned protocone is about half as tall as the metacone and about 1/3 of paracone. An extremely faint preprotocrista is present on its anterolabial face and is
pointed to the narrowest point (anterior waist) of the tooth. A postprotocrista is slightly more distinct than the preprotocrista. It is anteroposteriorly oriented and ends posteriorly at the posterior cingulum. Both paraconule (protoconule) and metaconule are absent and this gives the protocone an isolated appearance. The protocone is nearly completely encircled by a cingulum of various thickness. Lingually this cingulum forms an almost perfect half circle and has similar crown-heights throughout the protocone. The cingulum, however, thickens toward the posterolingual aspect of the protocone, more than twice as thick as its anterior segment, and along the medial and

Table 1 Measurements of upper first molars of Temnocyoninae and Haplocyoninae (mm)

<table>
<thead>
<tr>
<th>Taxa</th>
<th>Specimen</th>
<th>n</th>
<th>L1</th>
<th>L2</th>
<th>L3</th>
<th>L4</th>
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Note: Those for Temnocyoninae are adopted from Hunt (2011: tables 3, 5) and those for Haplocyoninae are means (when numbers of specimens are greater than one) from Peigné and Heizmann (2003: table 11). Since the lengths of M1s are defined somewhat differently between those in Hunt (2011: fig. 46) and those in Peigné and Heizmann (2003:8, fig. 3), we used the ratio of these two measurements (see Fig. 3 for definition of measurements) in IVPP V 22639 to estimate those of the haplocyonines (designated with an “e” after each measurement).
posterior sides of the protocone the cingulum has some minor wrinkles on its crest.

In anterior and posterior views, V 22639 has a distinctly concave appearance (Fig. 4A,
C) due to a deep trigon basin that occludes with a high-crowned m1 hypoconid. The high-
crowned paracone-metacone crest shears with the labial face of the m1 hypoconid.

3 Comparisons

Despite the meager material from Aoerban, there is no doubt that V 22639 belongs to
Amphicyonidae, although a dumbbell-shaped M1 is found in other similar-sized carnivorans,
such as Enhydrocyon and Paraenhydrocyon of hesperocyonine canids (Wang, 1994),
Megalictis of basal musteloids (Matthew, 1907), and Eomellivora of mellivorine mustelids
(e.g., Valenciano et al., 2015). However, in all of the latter three cases, a prominently
developed M1 parastyle dominates the labial half of the tooth with a much larger paracone
and very reduced metacone, in sharp contrast to a more evenly developed paracone and
metacone (with a more quadrate outline for the labial half of the tooth) in most amphicyonids.

The Aoerban tooth is easily distinguished from all known amphicyonids in Asia.
Excluding small, primitive forms in the Late Eocene through Early Miocene, such as
Guangxicyon sinoamericanus from the Late Eocene of Guangxi Province (Zhai et al., 2003),
unnamed species from Late Eocene of Mongolia (Egi et al., 2009), Cynodictis elegans from
Middle Oligocene of Nei Mongol (Huang, 1982), and the questionable Ictiocyon cf. I. socialis
from Early Miocene of Gansu (Wang et al., 2005), almost all Asian Miocene forms are large
amphicyonine amphicyonids. Well known examples include the Early Miocene Amphicyon
confucianus and Ysengrinia sp. from Shanwang, Shandong Province (Qiu and Qiu, 2013; Qiu
et al., 1986; Young, 1937), Middle Miocene Ysengrinia sp. of Japan (Kohno, 1997; Kohno et
al., 1997), Middle Miocene A. tairumensis of Nei Mongol (Colbert, 1939) and Gansu (Deng
et al., 2013), Middle Miocene A. ulungurensis of Xinjiang (Qi, 1989), and Late Miocene A.
palaeoindicus and Vishnucyon cf. V. chinjiensis from Yunnan Province (Qi, 2006), although
cautions must be exercised in some of these generic assignments because many Asian taxa of
large size, often based on fragmentary materials, tend to be placed in Amphicyon as tentative
assignments (Hunt, 2003). From Southeast and South Asia, several taxa are assignable to
amphicyonines, such as Middle Miocene Maemohcyon potisati of Thailand (Peigné et al.,
2006), Late Miocene Amphicyon of Myanmar (Egi et al., 2010; Sein and Thein, 2011), A.
sindiensis, A. palaeoindicus, A. pithecophilus, Arctampicyon lydekkeri, and Vishnucyon
chinjiensis from Early to Late Miocene Siwalik strata of India and Pakistan (Colbert, 1935;
Lydekker, 1884; Pilgrim, 1932). M1 in V. chinjiensis has a markedly narrowed lingual half
with a constricted appearance, but its protocone retains distinct pre- and postprotocristae
(Pilgrim, 1932:plate II, fig. 9). When present, the M1s in these Asian forms have triangular
outlines with distinct pre- and postprotocristae typical of amphicyonines, and are easily
distinguishable from the dumbbell-shaped Aoerban M1. Even without an upper molar, these
taxa have the reduced, low-crowned lower premolars or basined lower molars in contrast to
the high-crowned premolars and trenchant molars in haploxyonines and temnocyonines.

Aktaucyon brevifacialis from the Early Miocene of Kazakhstan shows a hint of waist
constriction in M1 (Kordikova et al., 2000). However, its pre- and postprotocristae are still distinct and it has a large M2 relative to M1, features that suggest that its M1 constriction may be a result of convergence. The Middle Miocene *Gobicyon macrognathus* from Nei Mongol (Colbert, 1939; Zhai, 1964), and possibly also from Xinjiang (Wang et al., 1998) and Gansu (Deng et al., 2013), may be an aberrant form of haplocyonines because of its high-crowned, piercing premolars, although its fragmentary m1 talonid appears to be not quite as trenchant as its European relatives (Hunt, 1998:208). Unpublished upper M1s from Hezheng area, Gansu Province, also exhibit a dumbbell shape (Qi Zhanxiang pers. comm.). A Serbian (formerly Yugoslavian) partial mandible previously referred to *G. macrognathus* by Pavlovic and Thenius (1959) has since become the basis of a new species *Haplocyonoides serbiae* (Ginsburg, 1999a), although its diagnostic characters are still in dispute (Peigné and Heizmann, 2003).

The distinctly constricted waist with a dumbbell outline in V 22639 is the most salient feature of this specimen, seen only in European haplocyonines and North American temnocyonines. Haplocyoninae is a lineage of hypercarnivorous amphicyonids from Late Oligocene to Middle Miocene of Europe, commonly including genera such as *Haplocyon, Haplocyonoides*, and *Haplocyonopsis* (Ginsburg, 1999a; Peigné and Heizmann, 2003). Dentally, it is characterized by high-crowned premolars and highly trenchant lower molars that may lose metaconids and entoconids (Bonis, 1966, 1973; Helbing, 1928; Peigné and Heizmann, 2003; Rothausen, 1988). Most haplocyonine species with a known M1 (including *Haplocyon crucians, Haplocyon elegans, Haplocyon dombrowskii, Haplocyonopsis crassidens, Haplocyonoides mordax, Haplocyonoides suevicus*) tend to be somewhat more primitive than V 22639 in their stronger parastyle, larger paracone relative to metacone with more acute angle of labial border of M1 with that of P4, M1 protocone with at least a remnant pre- and postprotocristae, and not very constricted waist area.

**Advanced haplocyonines** may have a dumbbell-shaped upper M1s, such as in *Haplocyonoides mordax* (Hürzeler, 1940). However, published figures of the only M1 of *H. mordax* (SMF-M1653e; Hürzeler, 1940:figs. 3-5) from the type locality in Hessler, Mainz Basin show several differences from that of V 22639: a relatively larger protocone, narrow cingulum surrounding the protocone, and a stronger preprotocrista with a distinct paraconule at the end (see Fig. 4F). Peigné and Heizmann (2003), however, referred a second M1 (SMF-M6000) from Budenheim, Rhineland-Palatinate, Germany, which has never been figured thus far. S. Peigné has graciously provided a photograph of this specimen to us that permits a much needed sense of variation (Fig. 4G, H). This referred specimen is strikingly similar to V 22639 in its much reduced protocone as a result of a widened surrounding cingulum, greatly reduced pre- and postprotocristae, and a nearly complete loss of a paraconule. As a result, lingual half of SMF-M6000 assumes a distinct appearance of an isolated protocone, not only because of its smaller protocone footprint relative to the broad surrounding cingulum but also its isolation from the trigon basin due to the reduction of pre- and postprotocristae. Details on the labial half of this tooth are also remarkably consistent with
that of V 22639. The labial side of SMF-M6000 is almost symmetrical with nearly identical size of paracone and metacone to the point that it is difficult to ascertain which side this tooth belongs without associated P4 or M2. Nevertheless judging by the slightly higher-crowned paracone and parastyle (Fig. 4H), we interpret this tooth to be a right M1, which, if correct, would be slightly more derived than V 22639 because of the general morphcline among European haplocyonines toward decreased parastyle and symmetrical paracone and metacone. Minor differences between the European and Chinese forms include a more anteriorly located protocone, a slightly more bulging parastyle, and a less distinct (not very ridge-like) lingual cingulum in V 22639.

_Haplocyonoides mordax_ was initially established from Hesseler near Wiesbaden in Mainz Basin, Germany in the Early Miocene (MN2) based on isolated upper and lower teeth (Hürzeler, 1940) that may or may not belong to a single individual (Rothausen, 1988). Bonis (1973:pl. 4, fig. 5) referred an additional right maxillary fragment with P4-M1 from Laugnac (MN2, Lot-et-Garonne) as _?H. mordax_, although he noted that its M1 lacks a median constriction. More recently Ginsburg (1999b:119) also listed _H. cf. H. mordax_ from “faluns of Anjou (MN3 or 5)” in addition to recognizing a new species, _H. serbiae_ (formerly _Gobicyon macrognathus_ by Pavlovic and Thenius, 1959), plus the previously described _H. ponticus_ (Kuss, 1960). While describing a new German species, _H. suevicus_, Peigné and Heizmann (2003:appendix) provided a complete list of hypodigm for _H. mordax_. In addition to materials from the type locality and Laugnac mentioned above, they included one specimen from Weisenau (MN1) and three specimens from Budenheim (MN2), both from Mainz Basin in Rhineland-Palatinate, Germany. Furthermore, they (Peigné and Heizmann, 2003:68) listed several sites from the MN3 of France and M2 of Spain, but preferred to treat _H. mordax huerzeleri_ (Rothausen, 1988) as _nomen dubium_ as well as casting doubt over generic assignments of _H. serbiae_ and _H. ponticus_.

V 22639 is also quite comparable to some North American temnocyonines, which share considerable similarities with European haplocyonines, interpreted to be convergences by Hunt (2011). In several respects temnocyonines have become even more hypercarnivorous than haplocyonines with short and robust crushing P4 in advanced forms, M1 protocone isolated by a flat platform of surrounding cingulum, substantial reduction of M2 and loss of M3, and digitigrade posture (Hunt, 2011). The M1 morphology is particularly diagnostic for the temnocyonines and clear trends can be observed to help delineate lineages. The smallest and most basal species, _Temnocyon altigenis_, has an enlarged parastyle, a protocone at the beginning stage of isolation but still with a preprotocrista connecting to a small paraconule, and a lingual cingulum slightly widened. Successively larger and more derived species, such as _T. subferox_ and _T. ferox_, have a more isolated protocone, lost preprotocrista (in _T. subferox_ but not in _T. ferox_) and paraconule, and more expanded lingual cingulum. _T. fingeruti_, apparently a side branch from the main chronospecies series (_T. altigenis-subferox-ferox_) (Hunt, 2011:fig. 70), does not quite follow this same trend, such as its retaining a strong
pre- and postprotocristae and a lack of middle constriction. Large, bone-crushing genera, *Mammacyon* and *Delotrochanter*, further elaborate this theme by widening the cingulum surrounding protocone, among other specializations.

In light of above trends within the main *Temnocyon* chronospecies series (Hunt, 2011:fig. 70), V 22639 falls somewhere close to *T. subferox* and *T. ferox* both in metric plot (Fig. 5) and some qualitative characters. The Chinese form has extremely reduced pre- and postprotocristae, not quite the complete loss in *T. subferox* (Hunt, 2011:fig. 10A) but certainly exceeded that condition in *T. ferox* (Hunt, 2011:fig. 12B). However, neither of the two North American species has reduced their parastyles quite to the extent as in V 22639. The degree of protocone area enlargement in V 22639 is also somewhat more pronounced than the above two American species. We also note that some characters are not strictly in a linear sequence, such as the precocious reduction of pre- and postprotocristae in *T. subferox*, although current knowledge about intraspecific variations is too limited to permit evaluation of morphological outliers.

A poorly known large species, *Temnocyon percussor* Cook, 1909, is also intriguingly close to V 22639. An isolated M1 (AMNH 81047) from the type locality (Cook Quarry, Fig. 5  Two dimensional plots of M1 anteroposterior length vs. M1 protocone length (see Fig. 3 for definitions) of Haplocyoninae and Temnocyoninae, as originally devised by Hunt (2011:fig. 46) to illustrate relative enlargement of the protocone area (Data from Table 1)
Anderson Ranch Formation, Sioux County, Nebraska, late Arikareean) was referred to this species (Fig. 4I). As described by Hunt (2011), this tooth is remarkably similar to V 22639. Its large size aside, AMNH 81047 has almost identical morphology of V 22639 in its protocone area enlargement: widening of lingual cingulum, reduction of pre- and postprotocristae, and reduction of parastyle. The only subtle difference is a slightly more convex labial border, compared to a very faint concave border in V 22639.

Although we cannot completely rule out the new Aoerban material being related to the North American temnocyonine, the balance of evidence, as shown in a single M1, seems to tip toward something close to *Haplocyonoides mordax*. In favor of such an assignment is the relatively closer age relationship of the Chinese and European forms (see Zoogeographic remarks below). If such a taxonomic assessment is correct, future discoveries of more complete materials of this Nei Mongol species should bear additional haplocyonine characters.

4 Zoogeographic remarks

In his study of Aquitanian mammals, Bonis (1973) commented the striking morphological similarities between European haplocyonines and North American temnocyonines, as was also noted earlier by Viret (1929). Bonis regarded these similarities as parallel developments by groups that are confined to their respective continents, a conclusion also maintained by Hunt (1996, 1998, 2011), who further explicitly derived his Temnocyoninae from within an early stock of Haplocyoninae (Hunt, 1998:fig. 11.3). Additionally, European haplocyonines and North American temnocyonines seem to have their independent evolutionary trajectory and chronological sequences, at least in the case of more derived temnocyonines (Fig. 5). If that is the case, it implies a single immigration event during the Early Oligocene that brought a primitive haplocyonine to North America to give rise to the temnocyonines, as has been postulated by Hunt (1998). However, *Haplocyonoides mordax* is much too late and too derived to be ancestral to temnocyonine.

If Peigné and Heizmann’s (2003) concept of *Haplocyonoides mordax* is to be followed, the age range for *H. mordax* extends to European Neogene mammal zones 1-3. If we take this at its face value, i.e., a species range of as long as ~5 million years (23-18 Ma), the later part of the European records would be equivalent in age to the new Nei Mongol record. However, based on actual M1s from MN2 of Germany that we can compare, the Chinese form is likely an eastward expansion of this lineage. If so, the Aoerban form would be the first occurrence of this lineage in Asia, possibly followed by later *Gobicyon* in the Middle Miocene.

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**内蒙古敖尔班地区早中新世犬熊类的新记录**

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**摘要**: 犬熊科(Amphicyonidae)的4个亚科(Amphicyoninae, Daphoeninae, Haplocyoninae和Temnocyoninae)中, Haplocyoninae (简齿犬熊亚科)和Temnocyoninae (剪切犬熊亚科)是两类比较特殊的、高度食肉化的姊妹群。Haplocyoninae仅出现在欧洲的晚渐新世到早中新世,而Temnocyoninae则局限在北美的早渐新世到早中新世。目前亚洲虽还没有记录,但新近纪的亚洲无疑是欧洲与北美之间迁徙途中的必经之路。因此本文记录的新材料既是意外发现又是意料之中。2015年我们在内蒙古中部早中新世敖尔班组下红层中首次发现一枚犬熊类的左上第一臼齿。这颗臼齿虽然零星,但非常特征,属Haplocyoninae或Temnocyoninae无疑,值得记述。该牙齿由于中间的收缩及原尖周边齿带的变宽而具有明显的哑铃型轮廓。另外其原尖上的前、后脊也几乎消失，形成一种原尖缩小而且孤立的形态。上述特征与欧洲早中新世的Haplocyonoides mordax（咬合似简齿犬熊）似乎最接近,但由于材料稀少敖尔班犬熊类与北美的Temnocyon percussor更为接近的可能也不能完全排除。如果上述判断正确的话，新发现的Haplocyonoides则是目前发现的少数几例早中新世从欧洲迁移到亚洲的种类之一。这一支系在亚洲有可能延续到中中新世的Gobicyon (戈壁犬熊)。

关键词: 内蒙古敖尔班, 早中新世, 犬熊亚科, 地理分布

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