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Rhinocerotidae (Mammalia) from the Late Miocene of Bulgaria

DENIS GERAADS, Paris, and NIKOLAÏ SPASSOV, Sofia
With 3 plates, 3 figures and 7 tables.

Zusammenfassung

Abstract
We describe the remains of Rhinocerotidae from about 10 Late Miocene localities of Bulgaria, the main ones being Kalimantsi and Hadjidimovo, and the newly discovered site of Strumyani. At least seven species are represented, mostly by cranial elements. A skull is assigned to Chilotherium kiliiasi, a species for which a new sub-genus, Eochilotherium, is established. Some fragmentary remains are referred to Ch. (Chilotherium), which reaches in Bulgaria the westernmost limit of its range. Acerorhinus is well represented at Kalimantsi, but this genus probably deserves revision. An unexpected discovery is that of Brachypotherium at Ahmatovo, its latest known occurrence in Europe. The horned rhinos, Ceratotherium and Dihoplus, co-occur at two localities. This relatively great diversity points to a variety of environments, as it is unlikely that the main fossil faunas are very different in age. It also reflects the geographic position of Bulgaria, where the ranges of the mainly Asiatic Chilotherium, of the Pontic Acerorhinus and of the Balkano-Iranian Ceratotherium overlap, in addition to a possible immigration of Brachypotherium from Africa into Europe.
Key words: Rhinocerotidae, Mammalia, Late Miocene, Bulgaria, Balkans

Schlüsselwörter: Rhinocerotidae, Mammalia, Obermiozän, Bulgarien, Balkan

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Introduction

Until now, the Miocene Rhinocerotidae of Bulgaria have been known only through a few outdated works, such as the early papers of BAKALOV (1933, 1939), a brief description by NIKOLOV & KOVACHEV (1966) and the monograph of BAKALOV & NIKOLOV (1962). Since then, more discoveries, and large-scale excavations conducted mainly in the areas of Hadjidimovo (Mesta River valley), Kalimantsi and recently at Strumyani (Struma River valley), all in south-western Bulgaria, have significantly improved the collections. A summary of our present knowledge of the main faunal associations is given in SPASSOV et al. (2006). Furthermore, several recent papers dealing with upper Miocene rhinos of the area (HEISSIG 1975, 1996, 1999; KAYA & HEISSIG 2001, GERAADS 1988, 2005; GERAADS & KOUFOS 1990; FORTELIUS et al. 2003; GIAOURTSAKIS 2003; GIAOURTSAKIS & HEISSIG 2004; GIAOURTSAKIS et al. 2006) have shed new lights on the evolution of this family. The present paper is an attempt to revise all available material, stored in the National Museum of Natural History, Sofia (NMNHS), in its Assenovgrad Museum branch (NMNHA), and in the Department of Geology, University of Sofia (DGUS). It is compared with other Late Miocene to Recent rhinos, stored in the Natural History Museum, London (BMNH); Faculté des Sciences, Lyon (FSL); Hessisches Landesmuseum, Darmstadt (HLMD); National Museums of Kenya, Nairobi (KNM); Musée Géologique et Minéralogique, Lausanne (MGML); Macedonian Museum of Natural History, Skopje (MMNH); Muséum National d'Histoire Naturelle, Paris (MNHN); Maden Tektik ve Arama Museum, Ankara (MTA); Naturhistorisches Museum Basel (NHB); Naturhistorisches Museum, Wien (NMW); National Museum of Ethiopia, Addis Ababa (NME);
Fossil localities (Text-fig. 1)


Hadjidimovo

Hadjidimovo is situated in the Mesta river valley, East of the Pirin ridge, not far from the Greek border. Four fossiliferous sites are known, the most important of which being Hadjidimovo-1 (Hadjidimovo-Girizite), with light clayey sands belonging to the Nevrokop formation (VATSEV 1980). It yielded more than 20000 fossils (NMNHA) belonging to about 30 mammal species, including the rhinos described below, making it one of the richest Late Miocene sites of the Eastern Mediterranean and peri-Pontic areas. Although the Hadjidomovo site (unknown locality) was first mentioned by NIKOLOV (1973, 1985), this huge collection was mostly accumulated and cared for, thanks to the efforts of D. KOVACHEV, mainly between 1985 and 1998. Hadjidimovo-1 could be earlier than Pikermi, as indicated by the evolutionary stages of *Mesopithecus* (KOUFOS et al. 2003) *Hipparion (Cremohipparion) mediterraneum* (HRISTOVA & SPASSOV 2005), *Adcrocuta eximia* (pers. obs. N.S.), and the presence of *Tragoportax rugosifrons* (SPASSOV & GERAADS 2004). SPASSOV (2002) suggested that the locality could be dated to the MN11/MN12 boundary.

All materials from Hadjidimovo are stored in NMNHA.

Kalimantsi

The Late Miocene deposits around the Kalimantsi village, on the western side of the Pirin ridge, in the Middle Struma basin, cover a large area (ca. 6 km2) with more than 10 known fossiliferous spots, scattered over a vertical distance of about 100 m. The recent revision of their biochronology demonstrates that all Kalimantsi sites are of Turolian age. Kalimantsi-1 (the river bank), belonging to the newly established Gradishte genetic lithocomplex, is more likely to yield a fauna of early Turolian age. *Hipparion cf. macedonicum* is known only from this level. The other Kalimantsi faunas (referred to the Strumyani genetic lithocomplex) are considered to belong to the middle Turolian (SPASSOV et al. 2006), probably slightly older than the Pikermi fauna, according to the evolutionary stages of *Hipparion (Cremohipparion) mediterraneum* and
H. (Hippotherium) brachypus (HRISTOVA & KOVACHEV 2005). Gazella sp., H. (Cremohipparion) mediterraneum and H. (Hippotherium) brachypus dominate in these levels. Tragoportax cf. amalthea, Palaeoreas lindermayeri, Helladotherium duvernoyi, Bohlinia attica, Adrocuta eximia, Mesopithecus pentelicus are also well represented (SPASSOV et al. 2006).

Presently, at least 34 species are known from Kalimantsi (all localities combined). Some specimens of Rhinocerotidae from Kalimantsi were mentioned and illustrated by BAKALOV & NIKOLOV (1962); they are stored in DGUS. New materials collected by D. KOVACHEV and co-workers, mostly from Kalimantsi-1, are stored in NMNHA.

**Strumyani**

The new rich locality Strumyani-2, near Sandanski in the Struma River valley, that we excavated in 2002-2003, has not been published in detail, but is likely to be of the same broad age (Turolian), probably early Turolian or first half of the middle Turolian, from the Hipparion assemblage (L.HRISTOVA, pers. comm.).

Some other localities have yielded only a few fossils.

**Central part of Western Bulgaria, Sofia region**

A quarry near Staniantsi in the Godech region (N.-W. Bulgaria), near the Serbian border, yielded only a fragment of incisor. The fauna might be Turolian (NIKOLOV 1985; SPASSOV 2002; MARKOV 2004b). It yielded unpublished mandibular teeth similar to those of **Tapirus balkanicus** from Balsha (SPASSOV & GINSBURG 1999) that can be referred to the **T. pannonicus** - **T. balkanicus** group and also indicate an ante-Pliocene age for the locality.

**Struma river basin, S-W Bulgaria**

Some localities yielded fragmentary rhino remains, but very little is known about the geologic context. A rhino incisor comes from the coal-rich levels, at a depth of about 200 m, of the Oranovo quarry near Simitli (south of Blagoevgrad), eponym locality of the newly established Oranovo genetic lithocomplex, probably of early Upper Miocene age.

Two rhinoceros finds were discovered near Slatino, north of Blagoevgrad. The deposits of the Slatino genetic lithocomplex are also considered as belonging to the lower part of the Upper Miocene (TZANKOV et al. 2005). Their precise provenance (most probably two different fossiliferous spots) is unknown. Their matrix is a grey-green gravellite clay concretion most probably washed by fluent waters. Such kind of clays, typical for the Slatino genetic lithocomplex (SPASSOV et al. 2006) crop out in several places around Slatino village.

At Kromidovo, not far from Kalimantsi, the fossils come from a spot in the village, but we could not locate it during our 2005 survey. The presence of **Mesopithecus pentelicus** (KOUFOS et al. 2003; SPASSOV et al. 2006), would point to an age close to MN12, but the
measurements of the specimens do not quite match those of the Pikermi sample, and this age estimate is far from secure.

Maritsa and Tundja river basins, middle part of Southern Bulgaria

A number of Upper Miocene fossiliferous spots are known from the Cherkezitsa River valley near Plovdiv, between the villages Seltsi, Bogdanits and Ahmatovo (NIKOLOV & KOVACHEV 1966). All of them could be approximately contemporaneous. The richest mammal fauna is from the sand quarry of Ahmatovo, the most important locality in the Maritsa River basin (BAKALOV & NIKOLOV 1962; NIKOLOV & KOVACHEV 1966). These authors reported Gomphotherium angustidens, but it is in fact Choerolophodon (TASSY 1983). DRAGOMANOV et al. (1981) referred the fossiliferous spots of the quarry to three different stratigraphic levels, but our observations do not confirm this stratigraphic subdivision, and age differences must be minimal. The revised list of the proboscideans from Ahmatovo is: Deinotherium gigantissimum, “Mammut” cf. borsoni, Choerolophodon pentelici, Tetralophodon atticus, Anancus sp. (MARKOV 2004b). As the genus Anancus appears in Bulgaria and Europe probably in the middle Turolian but in post-Pikermi time (SPASSOV et al. 2006) and Choerolophodon is not definitely known in the late Turolian, the age of Ahmatovo could be the second half/the end of the middle Turolian (MN12). The Brachypotherium skull and isolated teeth of an old individual described below are from the Ahmatovo quarry itself, while the isolated P4 of the same genus is from the deposits of the river-bank of Tcherkezitsa between Seltsi and Ahmatovo, stratigraphically slightly lower, but probably not much older.

In 1988 some small bovid, small giraffid and rhinos remains were discovered in the yard of the coachbuilder in the town of Yambol (on Tundja river). The Neogene deposits along the Tundja River belong to the Elhovska formation (KOJUMDJIEVA et al. 1984), the age of which ranges from the Maeotian to the Dacian-Romanian.

Systematic Palaeontology

Order Perissodactyla OWEN, 1848

Family Rhinocerotidae GRAY, 1821

We must stress again that the systematics of fossil rhinos is strongly defective, mainly due to the lack of comparative descriptions and diagnoses. We urge future workers to follow the recommendations of CIFELLI & KIELAN-JAVOROWSKA (2005), and give differential diagnoses, of which we tried to provide some drafts, in the hope that these might serve as a basis.

Uppercase denotes upper teeth, lowercase denotes lower teeth. Measurements are in mm, and follow GUÉRIN (1980). Measurements of tooth rows are at occlusal level, measurements of isolated teeth are maximum.
Genus *Chilotherium* RINGSTRÖM, 1924

**Type-species:** *Chilotherium anderssoni* RINGSTRÖM, 1924: 26. Late Miocene of China.

**Diagnosis:** Both sexes hornless. Straight nasals. Frontal region depressed, forming a shallow fossa. Parietal crests wide apart. Maxilla flat or slightly depressed, facial crest and nasal notch far from orbit. Premaxillae forming two thin vertical walls, lacking incisors. Lower jaw with widened symphysial part and large tusk-like i2, separated by a broad diastema. Permanent dental formula: I 0/1(2) C0/0 P 3/3 M3/3; deciduous dental formula DI 0/2 DC 0/0 DP 4/3(4). Antecrochet long. Body low and stout, belly low above ground. Limbs much shortened. Manus and pes tridactyl with short, diverging metapodials. Lateral metapodials obliquely directed backwards (mostly translated from RINGSTRÖM 1924).

Sub-genus *Eochilotherium* nov. subgen.

**Type-species:** *Aceratherium kiliasi* GERAADS & KOUFOS, 1990.

**Diagnosis:** supra-orbital process somewhat inflated, post-orbital process weak. Antecrochet moderate. Mandibular symphysis slightly broadened, incisors of plesiomorphic condition, close in structure to the i2 of *Aceratherium* / *Acerorhinus* (see below), not strongly enlarged, with subtriangular cross-section, with rounded border between the lateral and the ventral surfaces, which are subequal in width.

We reluctantly create this new sub-genus, because there are already too many names for European hornless rhinos, but it lacks several of the derived features which define the nominal sub-genus, *C. (Chilotherium)*, which includes e.g., *C. anderssoni* from China, *C. persiae* from Maragha, and *C. schlosseri* from Samos. These apomorphic characters are: 1) lack of upper incisors; 2) enlarged flattened lower incisors, inserted wide apart in a broadened symphysis, with an upturned mesial (internal) edge - this is perhaps the most important apomorphic feature, unique to this group (see below for a more detailed description); 3) no supra-orbital process, and a well-marked postorbital one; 4) short premolars; 5) very long antecrochet; 6) brachypodial limbs. Most of these characters were already listed in the original diagnosis (RINGSTRÖM 1924:26). In *C. (Eochilotherium)*, in which we include *C. (Eochilotherium) kiliasi* and *C. (E.) samium* (see below), the premolars are also short, but (1) and (6) are unknown, the lower incisors are medium-sized, with plesiomorphic characters (GERAADS & KOUFOS 1990, pl. 2, fig. 5), the symphysis is only moderately broadened, there is a strong supra-orbital process but no post-orbital one, and antecrochet is moderate. We agree with HEISSIG (1996, 1999) and FORTELIUS et al (2003) that these species can best be taken as primitive *Chilotherium*, but we prefer to keep them as taxonomically separate from the highly derived members of this group.
Chilotherium (Eochilotherium) cf. kiliasi GERAADS & KOUFOS, 1990

Type locality: Pentalophos, Greece, late Vallesian (MN 10) (see GERAADS & KOUFOS 1990: figs. 1-7; pl. 2, figs. 1-5; pl. 3, figs. 1 & 4; non pl. 3, figs 2, 3, 5).


Material: Unnumbered partial skull, with M3 erupting, lacking the cranial vault and pre-dental portion, from Kromidovo, DGUS (Pl. 1, Figs. A-B).

Description: Only the ventral part of the skull is preserved. The zygomatic arches form an even curve, without sharp bend at the level of the glenoid fossa. The cranial basis is narrow. The premolars are small relative to the molars (measurements: table 1). All teeth have a long crochet, and a weak antecrochet, but small tubercles in the median valley, best expressed on M1, may represent its lingual extensions. The premolars have a moderate lingual cingulum, and the hypocone of P3 has a small mesial spur, but the median valley remains wide open. There is a distinct paracone fold, and the labial wall of the ectoloph is less flattened than in typical Chilotherium.

Comparison and discussion: We refer this specimen to C. (E). cf. kiliasi on the basis of the close similarity in size and morphology of the teeth with those of a maxilla from Pentalophos (GERAADS & KOUFOS 1990, fig. 2), but we are well aware that cheek-teeth alone are not a very sound basis for species identification. GERAADS & KOUFOS (1990) described as A. kiliasi a sample of skulls and mandibles that FORTELIUS et al. (2003) showed to be heterogeneous. Indeed, the mandible PNT-142 has large premolars and is not anteriorly broadened, and must belong to a different species (see below). A. kiliasi, of which the best specimen from Pentalophos is PNT 12 (GERAADS & KOUFOS 1990, pl. 3, fig. 3-4), was referred to Chilotherium by HEISSIG (1996, 1999) and FORTELIUS et al (2003), who also referred some material from Sinap to this species. They considered it to be close to C. samium (WEBER, 1905), from Samos, a species that was excluded from Chilotherium by GERAADS & KOUFOS (1990) on the basis of WEBER's description. Since the type material of C. samium is lost, there will indeed always remain a doubt about the identity of the species, but sound hypotheses must be based upon actual specimens. Thus, our understanding of C. samium is based upon a skull from Samos, SMF-M3601, labelled as such by K. HEISSIG. Both species lack the apomorphic features of C. (Chilotherium) listed above, and are better referred to a separate sub-genus.

FÖRTELIUS et al. (2003) thought that C. samium is close to C. wimani RINGSTRÖM, 1924, from China, but the published material of C. wimani does show the apomorphic features of the
nominotypical subgenus, especially the huge lower incisors inserted on a much broadened symphysis (RINGSTRÖM 1924, fig. 28 and pl. 8, fig. 1).

Sub-genus **Chilotherium** RINGSTRÖM, 1924

**Type-species:** *Chilotherium anderssoni* RINGSTRÖM, 1924. Late Miocene, China.

**Diagnosis:** Supra-orbital process may be present but weak; post-orbital process well-marked. Lower jaw with strongly widened symphysial part. Antecrochet very long. Lower incisors (i2) long and very large, with triangular cross-section. The flat narrow latero-ventral face (lateral in other rhinos) is slightly broadened towards the base, and separated from the medio-ventral one by a sharp edge. The large dorso-lateral and medio-ventral ones are strongly broadened towards the base; their common edge forms the cutting edge, kept sharp by attrition with the upper lip, and dorso-medially oriented.

*Chilotherium* *cf.* *sarmaticum* KOROTKEVICH, 1958

**Type locality:** Berislav, Ukraine, late Vallesian or early Tuolian.

**Diagnosis:** A small and less cursorial chilothere; parietal crests wider apart than in other chilotheres; nasals short; antecrochet moderate; metapodials less massive and limbs more shortened than in other species; talus usually with only two calcaneal facets (simplified from KOROTKEVICH 1958b and 1970).

**Material:** i2 from Oranovo; perhaps a fragment of incisor from the quarry near Staniantsi.

**Description:** The crown of the Oranovo incisor is slightly curved. The tooth is flattened, of regular width. The cross section is triangular, even at the base. The dorso-lateral surface lacks enamel, as in other *Chilotherium* s. str., and is slightly concave, abruptly becoming narrower towards the apex. It is separated from the slightly convex medio-ventral surface by a dorso-medial edge much sharpened by wear, but with a distinct prominence at its base. A third, ventro-lateral surface is flat or even slightly concave, and delimited by two clear keels from the other surfaces. It is of almost even width towards the base. This third surface is not so sharply delimited and becomes broader towards the base in the tusks of the larger species of *Chilotherium*. Measurements: max. basal width of the wear surface ~ 39 mm., crown length = 78 (restored ~ 86 mm.), max. root diameter = 35 mm., width of ventro-lateral surface ~12 mm.

*Chilotherium sarmaticum* was first described from Berislav, Ukraine (KOROTKEVICH 1958a & b; 1970). Although not discussed in the synonymy lists of the chilotheres published by HEISSIG (1975, 1999), this species is smaller than the currently accepted taxa from the Turkey and Eastern Europe. The Oranovo incisor is also smaller than the large chilothere species from Grebeniki and Maragha (SPASSOV et al. 2006). Berislav has been referred to MN11a by
Korotkevich (1988; Vallesian for this author), to the late Vallesian by De Bruijn et al. (1992), or to the early Turolian by Krakmálnaya (1996). The morphology described by Korotkevich (1970) indicates a still less cursorial locomotory apparatus for *Ch. sarmaticum* than for other true chilotheres of the open spaces and the faunal analysis made by this author shows a more forested habitat in Berislav than in other localities of the same region such as Grebeniki and Novaya Emetovka. This would also be in good agreement with a Vallesian age. The Oranovo incisor is also virtually identical in shape and size with *Ch. cf. sarmaticum* described from Reghiu, Romania, a locality that was assigned to the beginning of MN11 (Stiuca 2003), but could be older. Oranovo could be of similar age. Codrea (1996) noted that it apparently never crossed the Carpathians. Oranovo marks together with Staniantsi and Kromidovo (Bulgaria), Pentalophos-1 (Greece) and Marievo region (Rep. of Macedonia: a molar fragment with preserved protoloph, coll. NMNHS) the western limit of the range of *Chilotherium*.

The tooth from Staniantsi is the apex of the left incisor, with a black to brown-grayish color, indicating that it comes from the coal levels of the quarry. It has all the characteristics mentioned in the diagnosis of *Ch. (Chilotherium)*. The tip of the tusk (as in the Slatino i2) has a small abrasion facet indicating that it was also perhaps used for digging; from what remains of the tooth, it looks somewhat larger than the i2 of *Ch. cf. sarmaticum* from Oranovo; thus, we refer it only doubtfully to the same species.

*Chilotherium* cf. *kowalevskii* Pavlow, 1913

**Type-locality:** Grebeniki, Ukraine, probably first half of the Turolian.

**Diagnosis:** dorsal skull profile gently concave, zygomatic arch very robust, paroccipital process rather short; upper premolars large, with strong crochet, and long antecrochet connecting the metaloph, thus closing the medisinus almost up to the top of the crown; i1 shed late in life; limb bones stocky.

**Material:** FM 2700 unworn left P3 and FM 2699 right P4 (Pl. 1, Fig. C); FM 2698 fragment of left mandible with m1-m3; FM 2697 incomplete mandible with left i2 and left and right p2-m2 (coll. NMNHS), coachbuilder yard, Yambol.

**Description:** besides the slight difference in L/W proportions (P3: 34.2 x 42.6; P4: 38.5 x 45), the two upper teeth are remarkably alike and are certainly from the same individual. They are moderately hypsodont; the labial wall is rather flat, with only a small paracone fold which vanishes towards the base of the crown. There is a complete lingual cingulum around P3, and probably also around P4, but it is much higher anteriorly and posteriorly, where it closes a antero-posteriorly elongated deep post-fossette. The protoloph forms a regular curve from the paracone to the hypocone, completely closing lingually the medisinus up to the top of the crown.
on P3, whereas on P4 its lingual end (antecrochet) descends somewhat, to meet the hypocone lower down. The protocone is only apparent as a distinct entity in the lower half of the crown, a deep pinching separating it from the rest of the loph. Both teeth have strong crochet and crista, which probably met to close a medifossette after heavy wear.

The mandible FM 2697 is of a subadult specimen. It shows the broadened symphysis typical of *Chilotherium*. The right tusk (i2) is missing, and only about 1/3 of the left one is erupted. Its cross-section is clearly triangular, with well expressed edge between the ventrolateral and ventro-medial faces. The paralophids of the cheek-teeth, including p2, are long, the anterior and posterior valleys are short, V-shaped. Measurements are given in Tables 2 & 3

The semimandible FM 2698 is from an old individual and lacks the anterior half. It has a thin corpus and relatively massive ramus, strongly developed and rounded mandibular angle, and large vascular incisure.

Comparisons: This morphology of the upper premolars (similar to the semimolariform pattern of HEISSIG, but probably secondarily acquired) is not common in the Eastern Mediterranean, and HEISSIG took it as characteristic of *Chilotherium kowalevskii*, a species first described from Grebeniki by PAVLOV (1913). The mandibular teeth dimensions fit well those given by KOROTKEVICH (1970) for the type locality, Grebeniki. HEISSIG (1996) listed it also from three Anatolian localities, and from Losengrad (Kirklareli) in European Turkey, from where BAKALOV & NIKOLOV (1962) had reported an upper molar as *Aceratherium angustifrons* ANDREE, 1921, a Samos species that HEISSIG (1999) considered as synonymous with *Ch. kowalevskii*.

Pending detailed study of the material (especially the rich collection from Çorakyerler), we will accept HEISSIG'S conception of the species, which is thus distributed along the Northern, Western and Southern parts of the peri-Pontic region.

*Chilotherium* sp.

**Material:** a right M3, coll. NMNHA – A1030, from Ahmatovo (Pl. 1, Fig. D).

**Description and comparison:** The tooth is in medium wear but was probably rather hypsodont. The labial wall is vertical, rather flat and with smooth relief, with weak paracone fold. There is a strong cingulum along the base of the protoloph base, plus a weak labial cingulum at the base of the ectoloph. The crochet is strong and curved mesially, the crista is vestigial. The antecrochet is characteristic of a chilothere in being very long, with a distal part curved lingually, descending abruptly down and reaching the lingual border of the tooth, leaving a narrow deep fossette between it and the protocone. The hypocone part is missing but was separated down to the bottom of the medifossette from the antecrochet. The protoloph is regularly curved. The protocone becomes abruptly wider towards its base; it is well separated
from the rest of the protoloph by a strong double pinching. The tooth is large: ectoloph length = 54.4 mm (estimated restored length ~ 56 mm); width = 51 mm. It is larger than the teeth of *Ch. sarmaticum* from Berislav and of *Ch. kowalevskii* from Grebeniki (KOROTKEVICH 1970). By its morphology and dimensions it represents a large and evolved species; for geographical reasons, the best choice would be *Ch. schlosseri*, known from Samos, but a specific identification would be harsh.

Genus *Acerorhinus* KRETZOI, 1942

Type-species: *Acerorhinus zernowi* (BORISSIAK, 1914), from Sebastopol (MN10).

Diagnosis: post-orbital process usually small, enlarged supra-orbital process, nasal notch close to orbit, often with a strong facial crest, premolars long, upper teeth with strong crochet, lower teeth with a short paralophid. The symphysis is moderately widened; the lower incisors are large and tusk-like, with subtriangular cross-section, with smooth border between the lateral and the ventral surfaces, which are subequal in width, and with large dorsal surface, moderately enlarged towards the base, limited by sharp medial and lateral cutting edges.

*Acerorhinus* sp.

Materials from Bulgaria: From Kalimantsi, in the NMNHA: K-595, almost complete skull, lacking the premaxillae and most of the nasals (Pl. 2, Figs. A-C); K-596, caudal part of a skull; K-600, fragment of maxilla with M2-M3; K-608, complete mandible (Pl. 2, Figs. D-E); in the DGUS: complete mandible (BAKALOV & NIKOLOV 1962, pl. 6, figs.1-2) and maxilla (BAKALOV & NIKOLOV 1962, pl. 5, fig.1). The mandible in the DGUS was discovered at Kalimantsi-1 (NIKOLOV 1985); all other materials are from higher levels of Kalimantsi. From an unknown locality near Slatino, called Slatino-1, in the NMNHS: fragment of maxilla (SPASSOV et al. 2006, fig.2B).

Description: On skull K-595, the nasal notch is long, reaching the level of M1, and is close to the orbit, the anterior border of which is above the middle of M2. The skull narrows rather abruptly in front of the orbits, but there is no conspicuous facial crest. The orbits are small, their dorsal rim is at the level of the skull roof, and the zygomatic root is deep. There is a long sagittal crest, and the occipital is high and inclined dorso-caudally. The cranial basis is narrow, the post-glenoid process is transversely flattened, the post-tympanic process is poorly expanded, the auditory region is cranio-caudally shortened, and the oval foramen is well-distinct from the anterior lacerated foramen. These latter features, which are also visible on K-596, are well-distinct from those observed in horned rhinos.

Measurements: length condylo-P2 = 463; length condylo-M3 = 216; length from condyle to front of glenoid surface = 125; from condyle to front of orbit = 290; from front of
orbit to nasal notch = 63; maximum supra-orbital width = 190; bizygomatic width = 300; maximum width over post-tympanic processes = 200.

The teeth of K-595 are in medium wear and well-preserved (measurements: table 1). The external wall is convex on the premolars but almost flat on the molars, except for the paracone fold. The protocone is pinched, with a well-marked antecrochet, flattened lingually, and expanded on M2-M3. There is no crista, but the crochet is long on M2 and moderate on the other teeth. The cingulum is continuous lingually on the premolars, but absent on the molars. The premolar row is almost as long as the molar row. The maxilla in the DGUS, referred to Aceratherium zernowi by BAKALOV & NIKOLOV (1962), is similar, except that the antecrochet is stronger on M2 but weaker on the premolars, not blocking the entrance to the medisinus.

The mandible K-608 (Pl. 2, Figs. D-E; measurements: table 2) has an almost straight ventral border, a ramus inclined dorso-rostrally, and an expanded goniac area. The diastema is short. The incisors are inserted not very wide apart, but the ventral border of the symphysis is concave between them. They are robust (diameters = 33.5 x 22), parallel but strongly curved upwards, with a subtriangular cross-section, with two sharp edges, facing medially and laterally, and a rounded ventro-lateral angle.

The cheek-teeth (measurements: table 3) are narrow and brachyodont, so that the crescents are shorter than in horned rhinos, but the paralophid has the same lingual extent. There is a labial cingulum on all teeth, and the premolars are rather long.

The mandible in the DGUS, which is from an older individual, displays the same features, except that the ventral border is slightly convex, the gonion less expanded, and the ascending ramus still more inclined. The incisors look thicker (diameters = 36.5 x 35) because the measurements are taken at the very base of the crown, and less upright, in part because they are more worn.

The Slatino-I maxilla has already been described by SPASSOV et al. (2006), who referred it to Aceratherium sp., but in light of the other material, it should more probably be included here. The antecrochet is well-marked, especially on the broken P4 (not figured by SPASSOV et al. 2006), the cingulum is very strong, thus resembling A. zernowi from Sebastopol.

Comparisons: The many species that were included in this genus (see the discussion on the genus below) can be compared as follows with the Kalimantsi form.

A. zernowi (BORISSIAK, 1914), the type-species, from the late Vallesian of Sebastopol, is rather distinctive, and differs from the Kalimantsi form by its short broad skull, correlated with an occipital plane inclined dorso-cranially and a short sagittal crest. The dorsal profile is concave, the orbit is less elevated, the protocone less flattened lingually, and the crochet and antecrochet are weak on the molars, but clearly stronger on the premolars, especially on P4. On the mandible, the diastema is long, the incisors are diverging, and they are certainly less upright.
than in the upper levels of Kalimantsi. According to HEISSIG (1999), *A. zernowi* also occurs at Pentalophos and Kayadibi, of late Vallesian to early Turolian age.

CERDEÑO (1996b) referred to the same species a nice sample from the Middle Miocene of Tung Gur, but the skull is much narrower relative to its length, the nasals are much longer (compare BORISSIAK 1914, pl. 6, fig. 1b, and CERDEÑO 1996b, fig. 2), the orbit is much lower than the skull roof. It further differs from the Kalimantsi sample by a shallow nasal notch, and stronger mandibular symphysis and incisors. We believe that the Tung Gur form is only distantly related to *Acerorhinus*.

FORTELIUS et al. (2003) also referred to *A. zernowi* several specimens from the Vallesian of Sinap Tepe, Turkey. Our comparisons are based upon their short description and our own brief examination of some of the Sinap material in MTA, including the skull and mandible illustrated by FORTELIUS et al. (2003, fig. 12.2), and supposed by them to be lost, but which was on display in the MTA museum as late as 2001. Still, our knowledge of this important sample remains poor. The Sinap skull (MTA-1998) has an occiput which is inclined dorso-caudally, a rather shallow nasal notch and a faint facial crest, as at Kalimantsi, but the zygomatic arch is more slender; the crochet is weak and the protocone is not very flattened, but the molars of a maxilla MTA-2021 (FORTELIUS et al. 2003, fig. 12.3) have a strong crochet and a protocone which is still more flattened lingually than in most *Chilotherium* (it is hard to understand why FORTELIUS et al. 2003:286, wrote the opposite). A mandible from Sinap Loc. 49 (FORTELIUS et al. 2003, fig. 12.4), of late Vallesian age, shares with K-608 upturned incisors; this is certainly a difference with the Sebastopol mandible (BORISSIAK 1914, pl. 8, fig. 2), since both are male in a similar wear stage. Thus, despite the approximate contemporaneity, identification of the Sinap material with the Sebastopol species is doubtful, but it does share some similarities with the Kalimantsi fossils.

From Karaslari near Veles, a skull in the Natural History Museum, Skopje, Macedonia, also has large lower incisors curving upwards. Although they are unfortunately mostly reconstructed in plaster, it is likely that they were rather similar to the ones from the upper levels of Kalimantsi.

More recent *Acerorhinus* from Turkey are a skull from Kavakdere (FORTELIUS et al. 2003, fig. 12.5), a skull from Kayadibi (HEISSIG 1999, Fig. 16.11, middle figure), and some limb bones from Akkaşdağ (ANTOINE & SARAÇ 2005). They have a concave cranial profile, and a robust zygoma, but the inflated supra-orbital process and much folded tooth pattern differ from the Kalimantsi skull.

From Chobruchi in Moldavia, a site probably close in age to Kalimantsi, PAVLOV (1914) described as *Aceratherium incisivum* several remains also referred to *Acerorhinus* by FORTELIUS et al. (2003), but the differences between them and *A. incisivum* from Eppelsheim (HLMD) or
Prottes in Austria (NHMW) amount mainly to a shallower nasal notch, deeper choanae, and more convex mandibular border, and generic distinction is perhaps not warranted. The Chobruchi material differs from the Kalimantsi sample by the same features, plus an occiput not stretched caudally, an orbit set less high in the skull, a very long post-glenoid process, a longer mandibular diastema and higher ramus (all these features make the Chobruchi rhino more like \textit{A. incisivum}), and no sagittal crest (perhaps not a very significant feature, as the \textit{A. incisivum} from Eppelsheim may or may not have a sagittal crest).

The skull of \textit{Acerorhinus fuguensis} DENG, 2000 from the latest Miocene of China is similar to that of Kalimantsi in its occiput inclined dorso-caudally, sagittal crest, robust zygoma, moderately deep nasal notch, but the narrowing in front of the orbit is stronger, the mandibular symphysis narrower, and the teeth are well-distinct by their long crochet (closing the medifossette on the premolars) and antecrochet, and the bizarre closure of the anterior valley on the lower teeth.

\textit{Diceratherium tsaidamense} BOHLIN, 1937, from the middle Miocene of China, has been referred to \textit{Acerorhinus} by HEISSIG (1975), and was considered by FORTELIUS et al. (2003) as close to the Sinap \textit{Acerorhinus}. The skull is high and narrow, the nasal notch very deep, the glenoid fossa high above the tooth-row, and the anterior premolars are quite small. The mandibular symphysis is rather narrow, but the mesial edges of the I2s are upturned, as in \textit{Chilotherium}. These are all differences with the Kalimantsi specimens.

These rather inconsistent stratigraphic and geographic patterns of distribution of the species referred to \textit{Acerorhinus} cast some doubts on the contents of the genus. The form from the late Vallesian of Sinap Loc. 49 might be close to the one from Kalimantsi-I and Karaslari, but its referral to \textit{A. zernowi} is disputable. Perhaps both samples could represent two stages of a separate lineage that could be a new species, but we refrain from naming it, pending detailed study of the Turkish material.

\textbf{Discussion}: The generic name \textit{Acerorhinus} has been mostly revived by HEISSIG and co-workers (HEISSIG 1989, 1999; FORTELIUS et al. 2003). According to them, and although a diagnosis of the genus has never been provided, this lineage shares with \textit{Chilotherium} the loss of upper incisors, and flattening of the lower ones, which are kept sharp by wear against the upper lip, all these characters being probably related. However, whereas the i2s of \textit{Aceratherium} and \textit{Acerorhinus} retain the plesiomorphic orientation and morphology of the lower tusks (dorsal occlusal surface limited by two cutting edges), derived \textit{Chilotherium} present an apomorphic condition of the tusks: the occlusal surface turns latero-dorsally and the tooth retains a single very sharp cutting edge, the medial one, which shifts to a dorsal position, in relation to some changes of the function of the food-catching apparatus, presumably becoming more effective for cutting hard vegetation. The nasal notch is deeper than in \textit{Chilotherium}, and thus closer to the
orbit, and the facial crest (the caudal border of the insertion of the narial musculature) is also close to the orbital border. The antecrochet is not lengthened as in *Chilotherium*, but the crochet may be quite long. *Acerorhinus* also lacks the *Chilotherium* specializations of triangular i2s with much upturned mesial (medial) edges, very wide symphysis, short premolars, and elevated orbit. Still, both genera must be closely related, as shown, e.g., by *Chilotherium primigenium* DENG, 2006, from China, which has an *Acerorhinus*-like face, but a very wide symphysis.

Unfortunately, one of the basic features for the definition of *Acerorhinus*, the loss of upper incisors, has not been definitely established for any of the species included in the genus, as no specimen has the premaxillae well-preserved. Lack of upper incisors is likely because the strong facial crest, caudal limit of the origin of the maxillo-labialis muscle, suggests that the latter, hence the upper lip, were quite powerful. Detailed examination of the lower incisors might tell whether they would wear against another tooth or against food and upper lip; but such a study remains to be done. However, in rhinos in which i2 wears against a well-developed I1, such as the modern *Rhinoceros* or *Aprotodon borissiaki*, there is a double attrition facet on i2, the lingual part being slightly concave. Such a distinction cannot be seen on the *Acerorhinus* i2s observed by us, probably showing the lack of the corresponding I1, or at least its very reduced size and function.

Still, the distinction from *Aceratherium* is often difficult, as the type-species *A. incisivum* from the Vallesian of Central Europe; also has, e.g., a deep nasal notch, abruptly narrowing nasals, a low orbit, large premolars, and incisors that are not more rounded than in some species of *Acerorhinus*. In spite of this, we agree with HEISSIG (1989, 1999), FORTELIUS et al. (2003), and GIAOURTSAKIS (2003) that there existed in the upper Miocene of South-Eastern Europe some rhinos matching their conception of *Acerorhinus*, but which were often named *Aceratherium incisivum* by local authors (BAKALOV & NIKOLOV 1962; NIKOLOV 1985; KOROTKEVICH 1988).

Genus *Brachypotherium* ROGER, 1904

*Type-species:* *Rhinoceros goldfussi* KAUP, 1834, from Eppelsheim (MN9).

*Brachypotherium* sp.

*Material:* A-599, anterior part of a skull, broken behind the orbits, and lacking the nasals; the whole bone surface is badly damaged, and the labial part of the ectoloph is missing on all teeth, except a small piece on the right M1 (Pl. 1, Fig. F); A-620, P4 (Pl. 1, Fig. E); A-1031, 1032 right and left P2s and A-1033 right M3, all three teeth probably from the same individual. All these specimens are from Ahmatovo in NMNHA. There is also an unnumbered incomplete juvenile skull from Kalimantsi in DGUS, referred to *Dicerorhinus schleiermacheri* by BAKALOV & NIKOLOV (1962), which formed the basis of the inclusion of the latter species in the faunal list of Kalimantsi 3-4 (Turolian fauna) by NIKOLOV (1985).
Description: The skull fragment A-599 is very large, close to the maximum size of the horned rhinos (*Ceratotherium*, *Dihoplus* and elasmotheres) of the Late Miocene. However, the orbit is not much lower than the cranial roof, the nasal notch is deep (bottom above the rear of P4) and very close to the infra-orbital foramen; these features point unambiguously to a "hornless" rhino. In the Late Miocene, most members of this informal group (*Aceratherium*, *Acerorhinus* and related forms, *Chilotherium*; some of them bearing in fact small horns) are much smaller than A-599, only *Brachypotherium* being comparable in size. Several other features support an assignment to the latter genus. The cranial roof is distinctly convex, not flat or even depressed as it is in the other hornless genera; the palate is broad; the premolars are much shorter than the molars, broad relative to their width, and the remaining fragment of the ectoloph on M1 unambiguously shows that this tooth was very brachyodont; the teeth have no crista, and the crochet is weak, although it is bifid on M2; the protocone and hypocone are only slightly pinched on the molars, and the antecrochet is weak. The cingulum is quite reduced, as it is also often in African forms and in *B. perimense* (cf. HEISSIG 1972, pl. 10, fig. 9), but this contrasts with most European teeth referred to *Brachypotherium*. Measurements of A-599: width over M2s = 245; bizygomatic width = 330 +; from orbit to nasal notch= 104. Estimated L x W of M2 (maximum) = 65 x 73. Estimated L x W of P4 = 46.5 x 67.5.

These measurements are close to those recorded by HOOIJER & PATTERSON (1972) for *B. goldfussi* from Eppelsheim, but it should be noted that they are significantly larger than those given by ROGER (1900) for the same population (e.g., P4 = 37 x 57; M2 = 49 x 56).

An isolated P4 A-620 is also very brachyodont (Pl. 1, Fig. E), and large (L x W = 52.6 x 63.1).

In spite of the strong wear, the convergent walls of teeth A-1031-1033 show that they were brachyodont. On the premolars, the labial wall is oblique and rounded, curving mesio-lingually; the lingual cingulum is well marked; there is a minute crochet and no crista; the protoloph is narrow but the protocone is large. The M3 also has weak external relief, rudimentary antecrochet and with arched external wall. The vestige of the metastyle, resting on the base of this wall, gives the basal outline a trapezoidal shape.

Measurements: P2: A-1031: L x W =37 x 46; P2: A-1032: L x W ~ 34.5 x 45.7; M3: A–1033: L x W = 53.3 x 49.5.

A juvenile skull from Kalimantsi in the DGUS, lacking the occipital, and with the nasal region reconstructed in plaster, was figured by BAKALOV & NIKOLOV (1962, pl. 9 and pl. 10, fig. 3) as *Dicerorhinus schleiermacheri*. It clearly belongs to a hornless rhino, as shown by the upper orbital border reaching the level of the skull roof, slightly inflated supra-orbital area, deep nasal notch, reaching the level of DP3 and almost incorporating the infra-orbital foramen, and
extremely robust rostral root of the zygomatic arch. On the milk premolars, the crochet increases in length from DP2 to DP4, there is a weak crista, the protocone is pinched but not flattened lingually, the labial wall has a well-marked paracone fold and a weaker metacone fold, and the cingulum is restricted to the mesial side. They are about as broad as they are long, much larger than in other hornless rhinos, and even larger than all Late Miocene Ceratotherium milk teeth that we have seen.

Comparisons: Brachypotherium had long been considered as a long-lasting genus, but HEISSIG (1999) referred the early middle Miocene forms to Prosantorhinus, thus restricting Brachypotherium to zones MN6 to MN9 (although FORTELIUS et al. 2003, extend its range to MN5). To our knowledge, there is only one secure mention in the European Turolian, at Eichkogel in Austria, well-dated to MN11 by rodents (DAXNER-HÖCK 1996), from where VACEK (1900) illustrated typical Brachypetherium lower molar and upper teeth. The genus is still present in the latest Miocene of both North and East Africa (HARRIS & LEAKEY 2003, and references therein), but the Bulgarian finds are the latest representatives of the genus in Europe. Two explanations can be offered for this late occurrence. Either, Bulgaria could have acted as a refuge for this genus. Another alternative hypothesis is that of a middle Turolian penetration(s) of Brachypotherium in Bulgaria from Africa via Asia Minor. A trilophodont gomphothere presumably from deposits near Seltsi (very close to Ahmatovo and probably close in age) was referred with caution to Gomphotherium angustidens by MARKOV (in press) who notes that a trilophodont gomphothere is also present in the Upper Nawata (Lothagam, Kenya; TASSY 2003), broadly contemporaneous with the fossiliferous area along Cherkezitsa area, and hypothesizes on possible faunal exchanges between East Africa and the Balkans in the latest Miocene.

RhinocerotinaeGRAY, 1821
Genus Dihoplus BRANDT, 1878
Type-species: Rhinoceros schleiermacheri KAUP, 1832, from Eppelsheim, MN9.

Dihoplus cf. schleiermacheri (KAUP, 1832)
Type locality: Eppelsheim, MN9.
Diagnosis: Nasal notch reaches only the anterior border of P2; temporal lines closely approaching; zygomatic arch robust; paroccipital process long. Upper I1 rather large and functional, I2 and I1 present, i2 very large.
Material: a skull fragment with well-worn right P2-M2, lacking molar ectolophs, from an unknown locality near Slatino, called Slatino-2.

This specimen, already described by SPASSOV et al. (2006), was referred to this species rather than to *D. pikermiensis* because of the position of the nasal notch. As discussed by these authors, this find suggests the occurrence of a Vallesian fauna near Slatino, as the record of *D. schleiermacheri* in the Turolian (including reports by BAKALOV & NIKOLOV 1962, and NIKOLOV 1985) is certainly due to its tooth similarities with *D. pikermiensis* as well as to a kind of traditional “inertia” to refer all Late Miocene "*Dicerorhinus*" s.l. to this species. It also occurs at Yulaflı in European Turkey (KAYA & HEISSIG 2001), but the site is in fact clearly Vallesian (GERAADS et al. 2005b).

*Dihoplus pikermiensis* (TOULA, 1906)

Type-locality: Pikermi

Diagnosis: Nasal notch more caudal than in the type-species, with an oblique ventral border. No sagittal crest. I2 absent, I1 vestigial, i2 reduced.

Material: From Strumyani: FM-2469: crushed but virtually complete skull (Text-fig. 2; Pl. 1, Figs. G-H); FM-2470 (coll. NMNHS): complete mandible, found close to the skull and almost certainly of the same individual (Pl. 1, Fig. I-J). From Hadjidimovo: HD-597: incomplete juvenile skull; HD-598: incomplete skull of old adult (Pl. 3, Fig. C); HD-605: maxilla of an old adult; HD-606: upper tooth-row; HD-607: juvenile maxilla (Pl. 3, Fig. D); HD-616: juvenile mandible; HD-697: juvenile maxilla; HD-671 to HD-674: upper premolars; HD-615: complete mandible (Pl. 3, Figs. E-F); some metapodials listed with their measurements table 6. From Kalimantsi: K-601, K-602, K-604: incomplete sets of upper milk teeth (coll. NMNHA).

Description: The skull from Strumyani is complete, but unfortunately crushed, and the very friable bone made it difficult to prepare. Still, its most important features are clearly visible (Text-fig. 2). It carried two horns, on the frontals and nasals. The nasals are long and form a gentle curve. The premaxillae, rarely preserved in fossil forms, are here intact; they are rectangular, about 60 mm long before P2, and show no trace of an incisor alveolus. The anterior orbital border is above M2-M3; the lower floor is not slanting ventrally. The occipital crest overhangs the condyles, but the outline of the occipital face is deformed. The cranial basis is shortened, and the post-tympanic process comes into contact with the post-glenoid process, although without fusion, but closing ventrally the aural foramen (Pl. 1, Fig. G). The cheek-teeth (Pl. 1, Fig. H) are much worn, but there is a weak metacone fold on P4, the spurs in the medisinus of the molars are rather weak, and the protocone tends to be isolated from the protoloph by vertical furrows. All these features of the skull and teeth point to *Dihoplus pikermiensis* rather than to *Ceratotherium neumayri*, the other two-horned rhino of the Eastern
Mediterranean Late Miocene (GERAADS 1988; GIAOURTSAKIS 2003; GIAOURTSAKIS et al. 2006). Measurements are given in tables 1 and 4.

The adult skull HD-598 is broken behind the orbits and in front of P2. The nasals are quite long and somewhat splayed out anteriorly, and the area of insertion of the posterior horn is rugose, but the upper profile is almost straight. The nasal notch is deep; its bottom is above the middle of P4. The front of the orbit is above the back of M2. The zygomatic arch is deep and robust; the inferior orbital border is not slanting ventrally. The choanae reach the anterior border of M2. The teeth are much worn (Pl. 3, Fig. C). The paracone rib is well marked; the hypocone is connected to the ectoloph by a narrow metaloph bridge. At this wear stage, on the molars, there is no visible crista, the crochet is weak, and on the premolars the lingual tubercles are connected. The protocone is clearly pinched on the molars, and has a small basal cingulum on P3-M3. A few more molars and premolars display the same features. An unnumbered occipital has an almost straight nuchal crest that is narrower than the base of the occipital face, which has an almost straight lateral profile.

The mandible from Strumyani, FM-2470, probably associated with skull FM-2469, has both sides well preserved except for the ramus, and the symphysis is complete. It narrows abruptly in front of p2, but is rather long. There is no i1, but a pair of i2s of unexpectedly large size (transverse diameter = 26mm, dorso-ventral diameter = 21mm). Like the cheek-teeth, they are much worn, and much cracked, so that the precise outline of the crown is hard to reconstruct, but the section was rather oval. The i2 occlusion facet is large, dorso-caudally and somewhat medially oriented and occupies all the width of the tip of the tooth. The ventral border of the mandibular corpus is straight.

HD-615 (Pl. 3, Figs. E-F) is a complete lower jaw with both sides well preserved. The depth of the corpus decreases regularly towards the front, with an almost straight lower border. The symphyseal region is similar to that of the Strumyani mandible. Only one non-functional and unworn, vestigial incisor is present on the right side. It is a minute peg-like tooth, inserted far away from the midline, and thus certainly a di2 or i2 rather than a di1. Although the median part of the symphysis is not perfectly preserved, it is almost certain than there was no di1 or i1. Measurements of the mandibles are given in tables 2 and 3.

Incisors, both deciduous and permanent, are absent in modern Ceratotherium, but they may be present in Diceros bicornis (SCHAUERTE 1966, and our observations), where they are minute, and almost certainly deciduous. Among the fossils, they are lacking in C. neumayri (BMNH M4399; SMF 13260; MGML S1060, all from Samos; also FORTELIUS et al 2003; GIAOURTSAKIS et al. 2006), but they seem to be normally present in Dihoplus. They are quite strong in the vallesian D. schleiermacheri, but reduced in D. pikermiensis.
The occurrence of upper incisors in *D. pikermiensis* is demonstrated by an unpublished skull from Pikermi in the NHMW, definitely of this species, with well-worn cheek-teeth, and which has well preserved long premaxillae bearing small incisors. According to GIAOURTSAKIS et al. (2006), a similar skull is preserved in the University of Athens.

Lower incisors are more common. Very small ones are present on two mandibles from Pikermi (MNHN N° 958; BMNH M10150), but GIAOURTSAKIS et al. (2006) noted that the alveoli are larger on another specimen from Pikermi (IPUW N°342red) and on a mandible from Kerassia, a locality close in space, and probably also in age, to Pikermi. Quite logically; they attributed these differences to sexual dimorphism. However, although these authors provided no measurement of the empty alveoli, they appear from their figure smaller than at Strumyani (GIAOURTSAKIS et al. 2006, fig.3C; the scale is certainly incorrect but the maximum width of the symphysis is given as 76.3mm). Two associated skulls and mandibles from Karaslari and Bashibos in the Natural History Museum, Skopje, Macedonia, also have no upper incisors but relatively large lower incisors, albeit smaller than at Strumyani.

HD-597 preserves the same part of the skull as HD-598, but from a juvenile individual. There is a clear contact between the lachrymal and nasal bones. The deciduous premolars are only slightly longer than broad, they have no crista, DP1 has a long metaloph, and DP2 has a postfossette. Both upper tooth-rows HD-607 (Pl.3, Fig.D) and HD-637/697 display the same features that are similar to those found in *D. pikermiensis* from Pikermi (GERAADS 1988). Some upper milk-teeth suggest that this species occurred at Kalimantsi as well. The rather wide postfossette on DP2, pinched protocone and incipient ante-crochet, are features that fit better this species than *Ceratotherium*, but the weak metacone fold, and presence of crista on some DP3s suggest the reverse identification; as most of these milk teeth are poorly preserved, their identifications are not definite. Measurements of the deciduous teeth are given in table 5.

On the mandible HD-616, the metalophid of dp3 has a posterior flange, which blocks lingually the posterior valley, and dp4 has a hint of anterior third lobe.

A number of post-cranial bones are preserved in the NMNHA, the best sample being that of the metapodials (measurements: table 6). They are slightly larger than in the living *C. simum* and than in the sample from Pikermi, which certainly includes both *C. neumayri* and *D. pikermiensis*. At present, there is no definite evidence that more than one species is present at Hadjidimovo, although the range of variation for some measurements is large (table 6), and we provisionally assign them all to *D. pikermiensis*, pending publication of the distinctive criteria (GIAOURTSAKIS, in prep.).

**Comparisons:** This species had often been confused with *Ceratotherium neumayri*, with which it co-exists at Pikermi and in other sites, but GERAADS (1988) showed that they are well distinct, and GIAOURTSAKIS (pers. comm.) greatly increased the number of post-cranial...
distinctive features. Its generic affinities are debatable. FORTELIUS et al. (2003) included it in *Stephanorhinus*, but the plio-pleistocene forms of this genus (type-species: *S. etruscus*) share at least two synapomorphies, loss of upper incisors and an ossified nasal septum, that are definitely not found in *D. pikermiensis*, and we feel it better to restrict this generic name to these well-defined Plio-Pleistocene forms. Instead, we will follow GIAOURTSAKIS (2003) and GIAOURTSAKIS et al. (2006) who link *Dihoplus pikermiensis* with *D. schleiermacheri*; they share a similar cranial and dental morphology, the main difference being the sharp reduction in incisor size in the Turolian form. The large size of the Strumyani incisors speaks in favour of an earlier age for this locality, and lends some support to the derivation of *D. pikermiensis* from *D. schleiermacheri*.

Tribe Dicerotini RINGSTRÖM, 1924

**Genus Ceratotherium GRAY, 1868**

*Type-species:* *Ceratotherium simum* (BURCHELL, 1817), living white African rhino.


*Ceratotherium neumayri* (OSBORN, 1900)

*Type-locality:* Maragha

*Diagnosis:* GERAADS (2005) considered this species as a stem-form from which both later *Ceratotherium* and *Diceros* arose, thus having no apomorphic features. Furthermore, the species is currently being revised by I. GIAOURTSAKIS, and we will provide no diagnosis.


*Description:* Skull K-594 (Pl. 3, Fig. A) is somewhat crushed dorso-ventrally, but its shape and proportions are quite clear. The dorsal profile is slightly concave. The nuchal crest largely overhangs the condyles, and is deeply notched medially. The skull is slightly broader at this level than over the post-tympanic processes. The impact of crushing on the morphology of this area is certainly weak. The bottom of the nasal notch is above the anterior border of P3, the infra-orbital foramen is above the anterior third of P4, and the anterior border of the orbit above the anterior border of M2. All these positions are quite close to the average for *C. neumayri* (see GERAADS 2005, fig.6). The lower orbital border is inclined ventro-laterally, the zygomatic arch is long and slender, and the lower pterygoid border strongly oblique. The cranial basis is rather
long, and the paroccipital process is well behind the post-glenoid process. Measurements are given in table 7.

The teeth (Pl. 3, Fig. B) are hypsodont, the ectoloph is sinuous but the paracone fold is weak. All teeth have a crochet and a small crista, the protoloph and metaloph are of regular thickness, and not pinched. The cingulum is continuous around the lingual side of the premolars.

All these features allow the distinction from *Dihoplus pikermiensis*, but are identical to those of *C. neumayri* from Pikermi and other eastern Mediterranean sites (GERAADS 1988, 1994), which is a likely ancestor for both modern African species (GERAADS 2005). They definitely show that this ancestor was more like the living "white" rhino than like the more browser *Diceros* in cranial morphology, and probably in ecology too.

K-593 is a juvenile maxilla with DP1-DP4 that can unambiguously be referred to the same species, rather than to *D. pikermiensis*, thanks to the criteria put forward by GERAADS (1988) and GIAOURTSAKIS et al. (2006): DP3 and DP4 are longer than broad; there is a strong crista, and no postfossette on DP2. Another, poorly preserved set of upper milk-teeth, K-603, is perhaps of the same species. Measurements of the milk-teeth are given in table 5.

The Strumyani mandible FM-2468 lacks the right ramus, the left one is damaged. It is not very large, but robust, with deep and strongly convex corpus. The symphysis is somewhat damaged on the right side but it is clear that it is short and very broad, without any trace of incisors. All these mandibular features, also present on a newly discovered (2007) mandible from the same locality, are well distinct from those of *Dihoplus*. The cheek-teeth are rather worn. The molar paralophids are long, better developed for example from the ones of the *Dihoplus* mandible from the same locality, which are in the same stage of wear. Measurements are given in table 2.

Two metapodials, a McII K-629 and an unnumbered McIV, are about as long as the shorter specimens from Hadjidimovo. The McII has a short magnum facet, and a palmar tubercle, like the specimens from Pikermi referred to this species (GERAADS 1988).

*Ceratotherium neumayri* is a common rhino of the Eastern Mediterranean Late Miocene. It has a good record in Pikermi, Samos, and as far East as the Caucasus and Maragha, and has been reported from a number of other sites in Greece and Turkey. Strumyani is its northernmost definite record. The range of this species probably matches that of open savannahs in the Turolian; we may tentatively suggest that its occurrence in the late Vallesian of Pentalophos (GERAADS and KOUFOS 1990) and Sinap (FORTELIUS et al. 2003) is linked with the precocious opening of the landscapes in this area.

It probably is the ancestor of both living African forms (GERAADS 1988; 2005); GERAADS & KOUFOS (1990) failed to recognize any evolutionary trend during the Late Miocene, but it is possible that its size increased towards the end of the Turolian (ANTOINE & SARAÇ 2005).
Conclusions

These new or revised Bulgarian finds greatly improve our knowledge on the Late Miocene Rhinos of the Balkan-Iranian zoogeographic province in several aspects:

Morphology and taxonomy.

The morpho-functional importance of the chilotherium tusks is reiterated, as these teeth have major taxonomic and ecological significance. The existence of forms with plesiomorphic features of i2, mandible, cheek-teeth and cranial features prompts us to separate them in a new subgenus *Chilotherium* (*Eochilotherium*). *Chilotherium* (*Chilotherium*) presents apomorphic features in the tusks. After the loss of I1, the dorsal (occlusal) surface turns latero-dorsally and the tooth retains only the very sharp (by attrition with the upper lip), sickle-like, medial cutting edge, which turns dorsally, presumably more effective for cutting hard vegetation, in relation to some change of the function of the food-catching apparatus and feeding behaviour. The species *Chilotherium sarmaticum*, mostly discussed in the Russian literature, but not in recent taxonomic and faunal analyses, should probably be considered as valid, thus representing the earliest true chilotherium.

*Acerorhinus* retains the plesiomorphic orientation and morphology of the lower tusks: occlusal surface dorso-medially oriented, limited by two functional cutting edges, lateral and medial. Two groups can perhaps be distinguished within this taxon: an early one (Vallesian to ?early Turolian), with broad and forwardly inclined i2, and a more progressive one, with increasingly upright tusks, that reaches at least the middle Turolian.

The discovery of a *Dihoplus pikermiensis* with large tusk-like i2s (Strumyani) is a strong argument supporting the gradual reduction of these teeth during the Turolian evolution of the genus.

Biochronology

The find of *Chilotherium* cf. *sarmaticum* in the Slatino genetic lithocomplex, positioned below the Strumyani lithocomplex with Turolian fauna, confirms that this is the earliest true chilotherium, living most probably at the Vallesian-Turolian boundary.

The discovery of *Brachypotherium* in Ahmatovo and Kalimantsi extends the vertical distribution of the genus in Europe, showing it to be present until the middle Turolian in the southwestern part of the continent.

The strong i2s of the *Dihoplus pikermiensis* mandible from Strumyani, the largest known in this species, supports the pre-Pikermian (early Turolian?) age of this locality.

Ecology

On the basis of the skeletal morphology, especially the shortened metapodials, *Chilotherium* has sometimes been interpreted as a swamp or marsh dweller (e.g., Beliaeva
1954), but RINGSTRÖM (1924) merely regarded it as suid-like, and in fact nothing suggests aquatic habits. Its specialized food catching apparatus made it well-adapted to the vast open landscapes of the central Asiatic province (GERAADS et al. 2002).

*Brachypotherium* has a still more shortened zeugopod, and is often depicted with an hippopotamus-like habitus. However, the stylopod is long (CERDEÑO 1996a; HEISSIG 1999; GERAADS & SARAÇ 2003), as in graviportal tetrapods, and the brachyodont teeth are also unexpected in a river or swamp dweller (those of modern hippos, or reduncines among bovids, are relatively high-crowned). HEISSIG (2004:253) supposed that it preferred "dryer woodland or maybe solid soil"; this is likely but its low-crowned teeth imply that it must have fed upon some soft plants.

The coexistence of *Ceratotherium neumayri* and *Dihoplus pikermiensis* at Samos, Pikermi, Kerassia, has been discussed recently (GIAOURTSAKIS et al. 2006); Strumyani-2 shows that they may be closely associated, because they were found at the same spot. As noted by several authors (HEISSIG 1999; GIAOURTSAKIS et al. 2006) *Dihoplus pikermiensis* could be a true browser and *C. neumayri*, although not a strict grazer as *C. simum*, had adaptations to tough low-level vegetation, such as lengthened skull, lack of incisors, and greater convexity of the mandible probably related to increased pressure on the cheek-teeth. Both species probably exploited different food niches in the same “Pikermian biome” of the Balkan Turolian (SPASSOV et al. 2006, and references therein).

The presence of *Ceratotherium* in the fossil material from the middle Turolian localities of Kalimantsi and the absence of this genus in the huge collection from Hadjidimovo is an indication of the more open conditions in Kalimantsi, in agreement with other data (MERCERON et al. 2006) and possibly reflects the gradual aridification of Southwestern Bulgaria with time.

**Zoogeography**

The new data refines the western boundary of the genus *Chilotherium*, at the same time confirming its occurrence in the Balkans (Fig. 3). The discovery of *Brachypotherium* could also have important biogeographic implications, if it could be demonstrated that this genus is an African immigrant rather than a Middle Miocene survivor; at present, it is hard to chose, as there is neither any representative of this genus in the upper Miocene of Greece or Turkey, arguing against a migration, nor any report in the Late Miocene of Bulgaria, arguing against a survival.

It may be expected that ongoing rhino studies in the Aegean-Pontic region will greatly improve our understanding of the Late Miocene radiation and diversification patterns in this difficult group, where mostly uniform skull and tooth patterns, and more generally scarce apomorphies (HEISSIG 1981) and frequency of parallelism hinder phylogenetic reconstruction.

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Captions to figures

Fig. 1. Map of Bulgaria with the main rhino-bearing localities.

Fig. 2. Schematic drawing of the skull of *Dihoplus pikermiensis* from Strumyani.

Fig. 3. Map of the Balkano-Turkish region with the main occurrences of Turolian rhinoceroses.

Captions to plates.


Table 1. Measurements of upper tooth series (occlusal lengths)

<table>
<thead>
<tr>
<th>Species</th>
<th>P2-P4</th>
<th>M1-M3</th>
<th>P2-M3</th>
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<td>127</td>
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<td></td>
<td>DGUS</td>
<td>130</td>
<td>138</td>
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<td><em>C. (E.) kiliasi</em></td>
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<td>-</td>
<td>142</td>
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<td>HD-605</td>
<td>-</td>
<td>146</td>
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<td><em>Ceratotherium neumayri</em></td>
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Table 2. Measurements of mandibles

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<th>length of diastema</th>
<th>length of symphysis</th>
<th>width of symphysis</th>
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Table 3. Measurements of lower tooth series (occlusal lengths)

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Table 4. Measurements of *D. pikermiensis* skulls

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<td>210</td>
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<tr>
<td>16</td>
<td>205+</td>
<td>275</td>
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<td>210</td>
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<tr>
<td>23</td>
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<td></td>
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Table 5. Measurements of upper milk-teeth

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<td>- x 42</td>
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<td>- x 33</td>
<td>46.1 x 41**</td>
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* height of unworn metacone = 35.5

** height of unworn metacone = 37.7
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<th>Locality</th>
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<th>prox. depth</th>
<th>dist. width</th>
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Table 7. Measurements of *Ceratotherium* skulls

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