# Carnivora from the Late Miocene of Kerassiá (Northern Euboea, Greece) 


#### Abstract

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Excavations carried out in recent years in Kerassiá brought to light many fossiliferous sites. In this study, the carnivores from the three richest sites, Kerassiá 1 (K1), Kerassiá 3 (K3) and Kerassiá 4 (K4), are discussed. K1 includes Metailurus cf. parvulus and possibly Adcrocuta eximia; K3 includes Plioviverrops sp. and cf. Ictitherium pannonicum; and K4 includes Adcrocuta eximia and Machairodus giganteus. The carnivores as well as the other faunal remains from these localities indicate an early to middle Turolian age.


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## INTRODUCTION

The existence of Miocene fossil vertebrates in Euboea has been known since 1878
(Cordella 1878). Further information is provided in Woodward (1901), Déprat (1904), Mitzopoulos (1947), Melentis (1968, 1969), Jacobi (1982), and others. These authors present information on the Limni, Hagia Anna, Rhovies, Achladi, Eria, Prokopi, Palaeovrissi and Halmyropotamos localities. The presence of fossils in Kerassiá has been known to local people since 1966, when fossil bones were found during the construction of a road close to the village. In 1981, Köhler tracked down the fossil bones of Kerassiá during geological fieldwork in the area. The first excavations in Kerassiá were carried out in 1982 by Hans de Bruijn and Albert van der Meulen (University of Utrecht) and Constantin Doukas (University of Athens). The excavations were not resumed until 1992, when the University of Athens (Department of Historical Geology
and Palaeontology) started systematic excavations in the area. These excavations, financed by the University of Athens, the Municipality of Nileas and the General Secretary of Research and Technology (Project $95 \Sigma$ YN107) brought to light more than seven fossiliferous sites. The fauna of Kerassiá has been described until now by Köhler (1983), Van der Made \& Moyà-Solà (1989), Theodorou et al. (1995, 1998, this volume) and Kostopoulos et al. (2001). Köhler (1983) mentions the presence of Microstonyx sp., Bovidae, Giraffidae, two Hipparion species, two species of Proboscidea and a large carnivore. Van der Made \& Moyà-Solà (1989) briefly describe the suids and attribute these to Microstonyx major erymanthius. Moreover, they mention the presence of Deinotherium and Dorcatherium and give a middle Turolian age for Kerassiá. According to Kostopoulos et al. (2001), the Kerassiá Microstonyx major is smaller than
the Pikermi M. major erymanthius and probably represents a new subspecies.
The carnivores of Kerassiá are represented from a small number of specimens, and the material is not always in a good state of preservation. In the present study, we discuss the carnivores of the three richest sites, the sites Kerassiá 1 (K1), Kerassiá 3 (K3) and Kerassiá 4 (K4). The material of this study comes from the excavations carried out from 1992 onwards and is temporarily stored in the Athens Museum of Palaeontology and Geology. In the future, this material will be exhibited in a new museum in Kerassiá that is planned by the local prefecture.

## Methodology and abbreviations

The methodology used for the dental measurements follows Werdelin (1988a). A value inside parenthesis signifies an approximate measurement, "a" signifies a measurement taken at the alveolus and " r " signifies a measurement taken at the root. The methodology used for the postcranial material is given with each table of measurements. AMPG: Athens Museum of Palaeontology and Geology. MNHNP: Muséum National d'Histoire Naturelle, Paris. BMNH: The Natural History Museum, London.

## SYSTEMATICS

Family: Hyaenidae Gray, 1869
Subfamily: Ictitheriinae Trouessart, 1897
Genus: Plioviverrops Kretzoi, 1938
Plioviverrops sp.
Locality: Kerassiá 3 (K3)

## Material

K3/B1/4: left humerus, lacking the proximalmost end. K3/B1/15: distal epiphysis of a right femur, juvenile. K3/B1/18: distal epiphysis of a left femur, juvenile. K3.192: right calcaneum, damaged on its sustentaculum tali. K3.206: proximal epiphysis of a left humerus, juvenile. K3.161: left Mt IV. K3.195: right Mt IV. Of the specimens listed above, those with the indication B1 come
from the same ossiferous block and most probably from the same individual.

## Description and comparisons

From Tables 1-3 it is clear that the available postcranial material is similar in its dimensions to that of Plioviverrops orbignyi (GAUDRY \& LARTET, 1856) from Pikermi, mentioned by Gaudry (1862, pl. 11, figs. 6, 10) and Pilgrim (1931). The available fourth metatarsals are only slightly larger than the specimen mentioned by Pilgrim (1931). In their morphological characters, the present specimens also show great resemblance to $P$. orbignyi. The distal articular surface of the calcaneum is flat and narrow as in P. orbignyi (Pilgrim 1931). The cuboid facet of the Mt IV is not extended posteriorly more than the tuberosity for the ligament of the peroneus longus, something also observed in P. orbignyi (Pilgrim 1931). The humerus is similar to that of P. orbignyi from Pikermi, except in its smaller supratrochlear foramen. Otherwise, the humerus has an entepicondylar foramen accompanied by a strong bar, as in P. orbignyi.

Except for $P$. orbignyi, the postcranial material of the Miocene small carnivores is poorly known. "Progenetta crassa" (=Protictitherium crassum) from the Can Llobateres II in Spain (Crusafont-Pairó \& Petter 1969, fig. 5, pl. 3, fig. 7) is significantly larger from the Kerassiá 3 material. Promeles palaeatticus (Weithofer, 1888) has a smaller calcaneum and metatarsals (Pilgrim 1931;
Roussiakis 1996), and its fourth metatarsal is shorter relative to the calcaneum. The ratio height calcaneum/length Mt IV is 58.3 in $P$. palaeatticus (data from Roussiakis 1996) but 51.7-52.7 in the Kerassiá Plioviverrops. Moreover, in P. palaeatticus the cuboid facet of the calcaneum is concave and oval, with its greater diameter obliquely oriented. In the Plioviverrops sp. of Kerassiá the cuboid facet is almost flat and anteroposteriorly elongated, as in P. orbignyi.
The available postcranial material thus shows dimensions similar to $P$. orbignyi from


Figure I Logarithmic ratio diagram comparing the Kerassia 3 ictithere with various species of icitheres. H. wongii from Samos according to Werdelin (I988a), I. viverrinum from Pikermi according to Werdelin (I988b), I. pannonicum from Polgardi and the Black Sea from Table 4.

Pikermi. In most of their morphological characters, the Kerassiá specimens also agree with P. orbignyi. Nevertheless, at the present time we prefer to refer the specimens considered here to Plioviverrops sp., since the lack of more material, especially teeth, does not permit us a more precise determination. The genus Plioviverrops is considered to be the oldest known hyaenid, appearing first in MN2a of Laugnac in France, with the species Plioviverrops collectus (De Bonis, 1973) (Ginsburg 1999). Various other species of Plioviverrops are known, such as Plioviverrops gervaisi De Beaumont \& Mein, 1972, from MN7/8 of La Grive (France), Plioviverrops guerini (Villalta \& Crusafont, 1948) from MN11 of Piera and Viverro de Pinos and MN12 of Cerro de la Garita and Los Mansuetos in Spain (Ginsburg 1999), and Plioviverrops faventinus Torre, 1989
from the late Turolian (MN13) of Brisighella in Italy and MN14 of La Gloria 4 in Spain (Alcalá 1994). P. orbignyi, which shows resemblance to our findings, is reported from the early-middle Turolian (MN11/12) of Ravin des Zouaves 5, Vathylakkos 2, 3, Prochoma 1 and Perivolaki (De Bonis \& Koufos 1991, Koufos et al. 1999, Koufos 2000), and from the middle Turolian (MN12) of Pikermi and Samos (Major 1888, Major 1894, De Beaumont 1969, Bernor et al. 1996) in Greece. It is interesting to notice that in Vathylakkos 3, P. orbignyi seems to have coexisted with another form, listed as $P$. cf. guerini by De Bonis \& Koufos (1991) and Koufos (2000).

Genus: Ictitherium Roth \& Wagner, 1854 cf. Ictitherium pannonicum KretzoI, 1952 Locality: Kerassiá 3 (K3)

## Material

K3/204: right mandibular ramus with ci, p3, p 4 and m 1 . The alveoli of $\mathrm{p} 1, \mathrm{p} 2$ and m 2 are visible.

## Description

The available mandible is badly damaged and some of its characters cannot be evaluated precisely. The mandibular corpus has a mental foramen below the middle of the alveolus for p 2 . The lower border of the mandibular corpus slopes down from the symphysis to a point under m 1 . The depth of the mandible thus increases from the anterior to the posterior part and the depth in front of p 2 is smaller than that behind m 1 (Table 4). The total length of the symphysis measures about 40 mm and its posterior border lies under the middle of the alveolus for p 2 . The alveolus for p 1 is separated from the canine by a diastema of 9.9 mm . The p 3 has a posterior accessory cusp and most probably a smaller anterior one. The p4 is more elongated than p3 (Table 4) and has a robust anterior accessory cusp. The posterior part of p 4 is not very well preserved but almost certainly had an accessory cusp. The most important character of ml is the presence of a strong metaconid. The protoconid and paraconid are almost equal in height. The talonid of m 1 is large and wide, but very worn and its morphological details are not visible. The only preserved cusp is the entoconid, which is high. The m2 is not preserved, but judging from its elongated alveolus it was large.

## Comparisons

The group of the 'ictitheres' contains many species and genera that share numerous characters and sometimes have slight differences that make their identification difficult. In this study, we follow the revision of Werdelin \& Solounias (1991). The various species of the 'ictitheres' have been referred mainly to the genera Ictitherium WAGNER, 1848, Thalassictis Gervais ex Nordmann, 1850, Hyaenictitherium Kretzoi, 1938, Palinhyaena Qiu, Huang \& Guo, 1979, and Hyaenotherium

Semenov, 1989.
Ictitherium viverrinum Roth \& WAGNER, 1854 is a well-known species with a wide geographical range. Compared to I. viverrinum from Pikermi, K3.204 has clearly larger $\mathrm{p} 3, \mathrm{~m} 1$ and m 2 (Fig. 1). Thalassictis robusta Gervais ex Nordmann, 1850, from Kishinev, is also smaller than the Kerassiá ictithere (Fig. 2). Hyaenictitherium hyaenoides (ZDANSKy, 1924) has larger teeth than the Kerassiá specimen and a small m2 (Fig. 3). Moreover, the mandible of that species has a more "hyaenoid" appearance. It is robust, straight, and with a lower border that is almost parallel to the alveolar border (Qiu 1985). The mandibular corpus of K3.204, on the other hand, is deeper in its posterior part (Table 4). Palinhyaena reperta Qiu, Huang \& Guo, 1979, is known from Chinese localities only (Werdelin \& Solounias 1991). This species is also larger than the Kerassiá 3 ictithere, but with a smaller m2 (Fig. 3). The mandible of $P$. reperta is similar to that of $H$. hyaenoides, i.e., 'hyaenoid' in appearance.
The lower premolars of K3.204 are slightly more robust than those of Hyaenotherium wongii (ZDANSKY, 1924) from Samos (Fig. 3). This is especially true for p 3 , while p 4 is badly damaged and its width is only approximately given (Table 4). The most important difference concerns m 2 . This tooth is not preserved in the Kerassiá specimen, but judging from its elongated alveolus that measures about 8.9 mm , it was especially large (Table 4, Fig. 3). Such a character differentiates our specimen from $H$. wongii and makes it more similar to Ictitherium pannonicum Kretzoi, 1952, a species that was first described from Polgardi in Hungary, but it was actually ignored until Semenov $(1985,1989)$ restudied these specimens and attributed some more specimens from the Black Sea area to it. Werdelin \& Solounias (1991), however, note that we cannot be certain if the Black Sea specimens belong to the same species as the Polgardi specimens. I. pannonicum is slightly larger overall than $H$. wongii, but in some of its dimensions is within the range of $H$. won-


Figure 2 Logarithmic ratio diagram comparing the Kerassia 3 ictithere with various species of icitheres. H. wongii from Samos according to Werdelin (1988a), T. robusta from Kishinev according to Kurtén (1982), I. pannonicum from Polgardi and the Black Sea from Table 4.
gii (Fig. 3). The most important distinguishing character of I. pannonicum is the large m 2 (Figs. 2, 3). As mentioned earlier, such a character is also present in our specimen. Aside from the large m 2 , the Kerassiá 3 specimen resembles I. pannonicum in its m 1 but it has more robust p3 (Table 4, Fig. 2).
Unfortunately, the species I. pannonicum is poorly known. In addition to Polgardi (Kretzoi 1952), which is the type locality, specimens referred to that species are also known from Chobruchi in Moldova, and Novaja Emetovka 2 and Cherevichnoe in Ukraine (Semenov 1985, 1989). A specimen from Valdecebro 5 in Spain referred to Thalassictis aff. hipparionum by Adrover et al. (1986), may, according to Werdelin \& Solounias (1991), also belong to I. pannonicum. Alcalá (1994) further mentioned a fragmentary mandible from the middle Turolian (MN12) of Cerro de la Garita in Spain under the name 'Ictitherium aff. I. pannonicum'. The p 2 and p 3 of that specimen are comparable in size to those of I. pannonicum, but the
p4 is larger. Moreover, that specimen lacks any teeth posterior to p4. Another species possibly related to $I$. pannonicum, is Ictitherium intuberculatum Ozansoy, 1965, from the Yassiören in Turkey. Werdelin \& Solounias (1991) emphasize the possibility that this form might be conspecific with $I$. pannonicum, but Ozansoy (1965) does not give metric data for the lower dentition and comparison with the Kerassiá ictithere is not possible.

Of the localities mentioned, Chobruchi is considered to be early Turolian (MN11) (Semenov 1989), Valdecebro 5, Novaja Emetovka 2 and Cherevichnoe middle Turolian (MN12) (Adrover et al. 1986, Semenov 1989) and Polgardi late Turolian (MN13) (Ginsburg 1999).

Subfamily: Hyaeninae Mivart, 1882
Genus: Adcrocuta Kretzoi, 1938
Adcrocuta eximia (Roth \& WAgner, 1854)
Locality: Kerassiá 4 (K4)


Figure 3 Logarithmic ratio diagram comparing the Kerassia 3 ictithere with various species of icitheres. H. wongii from Samos, P. reperta and H. hyaenoides from China according to Werdelin (1988a), I. pannonicum from Polgardi and the Black Sea from Table 4. 4.

## Material

K4.7: skull fragment with the right P3 and P4. The P3 is badly damaged and the alveolus of M1 is visible. K4/ $\Delta 388 / 1$ : skull fragment with the left I1-P3, Cs, P1-P4 and M1, and the right I1-I3, Cs, and P3-P4. K4/ $\Delta 388 / 2$ : left mandibular ramus with the complete dentition (i1-i3, ci, p1-p4 and m1). K4/ $\Delta 388 / 3$ : right mandibular ramus with the complete dentition except p 1 . The lower mandibles K4/ $4388 / 2-3$ were found connected at their symphysis, and so come from the same individual. The specimen $K 4 / \Delta 388 / 1$ was found next to these mandibles, and it is almost certain that all these specimens belong to the same individual. This is also indicated from their ontogenetic stage, since the teeth are almost unworn and neither the upper nor the lower canines have fully erupted.

## Description of the skull and the upper dentition

The infraorbital foramen opens above the
posterior half of P3. The maxillopalatine suture is not clearly visible, but most probably its anterior limit lies opposite the posterior part of P3. The anterior palatine foramina are situated level with the middle of P 2 . The posterior palatine foramina are not visible. The I1 and I2 are arranged in an almost straight line, while I3 is situated slightly posteriorly. The I2 is only slightly larger than I1, but I3 is significant larger. The I1 and I2 have a distal cingulum composed from two cusps, with the lingual one slightly larger than the lateral one. The upper canine has a mesiolingual and a distal crest. The P2 is not well preserved, but shows a posterior accessory cusp. On its anterior part, we cannot clearly observe an accessory cusp, but only a small bulge at the lingual side of the base of the anterior crest. The P3 has a posterior accessory cusp and a small anterior one in a mesiolingual position. The greatest width is at the anterior part of the tooth. The P4 is characterized by a small protocone, a parastyle that is smaller
and lower than the paracone, and a metastyle that is slightly longer than the paracone (Table 5). The distal part of the metastyle blade is directed labially. M1 is small and its metacone projects slightly posteriorly.

## Description of the lower mandible and the lower dentition

The mandibular corpus has two mental foramina. The anterior foramen opens under the middle of p 2 and the posterior one under the contact between p 2 and p 3 . The symphysis measures about 52 mm in length and extends posteriorly to the anterior part of p 2 . The anterior border of the masseteric fossa is situated posterior to m 1 . The incisors increase in size from i1 to i3 and they have a lateral cusp, which shows an increase in size from il to i3. The lower canine has a mesiolingual and a distal crest and is slightly more curved than the upper canine. The cheek teeth are arranged in a curved line. The posterior part of p 3 slightly overlaps the anterior part of p 4 , and the posterior part of p 4 the anterior part of ml . The anterior and posterior borders of p 3 and p 4 are almost straight, giving them a rectangular outline. The p 1 is small and situated lingually relative to p 2 . The p 2 has a posterior accessory cusp but not an anterior one. The p 3 has a posterior accessory cusp and a faint anterior one at the lingual side of the base of the anterior crest. The p4 has two accessory cusps, an anterior and a posterior one. The anterior one is more robust and situated slightly lingually. Behind the posterior accessory cusp, the cingulum is elevated. The premolars lack a lingual cingulum. The m 1 lacks a metaconid, while the talonid is small and two-cusped, retaining the entoconid and the hypoconid, of which the hypoconid is the larger. There is a weakly developed cingulum at the anterior part of the paraconid, both lingually and buccally. There is no m 2 or an alveolus for it.

## Comparisons

Hyaena eximia was first described by Roth \& Wagner (1854) from the classical locality of

Pikermi. Later it has been found at many fossilliferous sites in Europe and Asia. The generic attribution of the species has changed several times, and it is now attributed in Adcrocuta Kretzoi, 1938, opinion followed also here.

The available specimens from Kerassiá have all the morphological characters of $A$. eximia. This species shows some variation in the position and the number of the mental foramina (Gaudry 1863, Werdelin \& Solounias 1991, Koufos 2000). The specimen figured by Roth \& Wagner (1854, pl. 8, fig. 6) has two mental foramina, almost in the same positions as the Kerassiá specimens. Gaudry (1863) mentioned that of three specimens available to him, two had two mental foramina, but another specimen had only one mental foramen. The same variation is also present in the Axios material. The specimen RZO-126 has two mental foramina, while the specimen RZ1-4 has one (Koufos 2000). Some variation is also observable on the accessory cusps of the upper and lower premolars. This variation concerns their presence or not, as well as their development. The P2 and P3 generally have small accessory cusps. The anterior accessory cusp of p 2 is small or absent, while that of p 3 is present but sometimes small. The ml most commonly lacks a metaconid. It has been observed, however, in some cases, as on MNHNP MAR G. 12 from Maragha figured by De Mecquenem (1925, pl. 9, fig. 7) and on RZ1- 4 from Ravin des Zouaves 1 (Koufos 2000). A small metaconid, appressed to the main body of the protoconid, has also been observed (Roussiakis 1996) on AMPG P.G. 95/1505 from Pikermi (specimen mentioned with the number 113 by Howell \& Petter 1985, tab. 6a).

The dimensions of the specimens $K 4 / \Delta 388 / 1$ and $K 4 / \Delta 388 / 2-3$ are within the range of variation of $A$. eximia from Pikermi (Tables 5-6, figs. 4,5). The observable differences are not statistically significant. The M1 of $K 4 / \Delta 388 / 3$ is slightly large relative to the premolars, but is damaged at the area between the trigonid and the talonid. The P3
and P4 of K4.7 are comparable to the largest specimens from Pikermi (Fig. 4). The metastyle of the P4 of K4.7 is large (Table 5) but that tooth is badly damaged and broken at the limit of the paracone and metastyle.
The present specimens show significant differences from Adcrocuta eximia leptoryncha Bonis \& Koufos, 1981, from the Vallesian of Ravin de la Pluie. According to Bonis \& Koufos (1981) and Koufos (2000), this subspecies differs from the typical A. eximia in its longer snout, narrower palate and more slender premolars. With regard to the lower premolars, there are no significant differences from the Pikermi $A$. eximia, something mentioned also by Howell \& Petter (1985). The p3 and p4 are slightly smaller in some of their dimensions (Fig. 5) and p4 has an index of compression (Wp4/Lp4) equal to 50.7 on RPL-15, which is slightly lower than the Pikermi range (Table 6). The indices of the upper premolars are within the range of $A$. eximia from Pikermi (Table 5, fig. 4). The skull characters of $A$. eximia leptoryncha, however, distinguish that subspecies from the Pikermi, Samos and Maragha A. eximia. The skull of $A$. eximia leptoryncha has a longer snout and relatively narrower palate, the length/width ratio of the palate being 154 (Bonis \& Koufos 1981). The corresponding ratio (estimated following the methodology of Bonis \& Koufos 1981) for K4/ $4388 / 1$ is 123 , a value very close to those given by Bonis \& Koufos (1981) for the A. eximia from Pikermi (125) and Maragha (117). On P.A. 490/91 from Pikermi, this ratio is about 127.

The species Chasmaporthetes bonisi Koufos, 1987 is known from the early Turolian locality of Ravin des Zouaves 5 and the late Turolian locality of Dytiko 1. This species differs from A. eximia mainly in the upper dentition. The M1 is larger than that of A. eximia (Fig. 4) and the protocone of P 4 is more developed and directed slightly anteriorly. The P2 is also slightly less robust than that of $A$. eximia from Pikermi, but not significantly so. In C. bonisi the robusticity of the
lower premolars, lies close the minimum values of the A. eximia from Pikermi. The robusticity index for the p 2 of DTK-126 (sin.) is also similar to that of the A. eximia specimen RZ1-4 (sin.) (Fig. 5). The morphology of the lower premolars of C. bonisi, however, differs from those of $A$. eximia. The lower premolars of $C$. bonisi have a more elliptical outline, especially anteriorly, while the accessory cusps lie almost on the axis of the teeth. In A. eximia the anterior cusps of the lower premolars are in a more lingual position, and the anterior and posterior borders are almost straight, giving an almost rectangular shape to the teeth.
Most probably A. eximia is also present at the Kerassiá $1(\mathrm{~K} 1)$ site. This is indicated by the specimen $\mathrm{K} 1 / \Delta 104$ (a left P4). Unfortunately, this specimen is so fragmentary that no measurements can be taken.
The species $A$. eximia is known mainly from the Turolian (MN11-13) of Europe and Asia as well as from northern Africa, with its maximum in the middle Turolian (MN12; Howell \& Petter 1985, Koufos 2000). Its existence in the Vallesian of Hostalets de Pierola in Spain and Soblay in France, has been shown to be unconfirmed (Howell \& Petter 1985). Ravin de la Pluie, Ravin des Zouaves 1 and Xirochori 1 are the only known late Vallesian localities with Adcrocuta eximia remains (De Bonis et al. 1988; Koufos 2000). As shown earlier, the Ravin de la Pluie subspecies A. e. leptoryncha, is clearly different from the Kerassiá $A$. eximia. The Ravin des Zouaves 1 A. eximia is also different from the Kerassiá specimens. According to Koufos (2000), the late Vallesian A. eximia from Ravin des Zouaves 1 has strong lingual cingulum on the lower premolars, a character considered to be primitive by Koufos (2000). Such a cingulum is absent on the Kerassiá specimens, indicating that the Kerassiá $A$. eximia is more similar to the typical Turolian forms.


Figure 4 Logarithmic ratio diagram comparing the upper dentition of the Kerassia 4 Adcrocuta with A. eximia and C. bonisi from various localities. H. hyaena (standard) according to Howell \& Petter (I980). Data for the Axios, Ravin des Zouaves 5 and Ravin de la Pluie hyaenids according to Koufos (2000). The rest from Table 5.

Family: Felidae Gray, 1821
Subfamily: Machairodontinae Gill, 1872
Genus: Machairodus Kaup, 1833
Machairodus giganteus (WAGNER, 1848)
Locality: Kerassiá 4 (K4)

## Material

K4.14: right upper canine. $\mathrm{K} 4 / \Delta 69 / 1$ : mandible, with both left and right mandibular rami. The condyles are only partly preserved, while the mandibular angles and the coronoid processes are broken. In the right ramus the $\mathrm{i} 1-\mathrm{i} 3, \mathrm{p} 3-\mathrm{p} 4$ and m 1 are preserved, but the tips of i2 and i3 are broken and the posterior part of p4 is badly damaged. Only the root of the right ci is preserved. In the left ramus the complete dentition (i1-i3, ci, p3-p4 and m1) is preserved but the main cusp of p 3 , the anterior half of p 4 and the posterior half of m 1 are broken. $\mathrm{K} 4 / \Delta 110 / 16$ : Left radius, complete. K4/د82: Right Mt IV, badly damaged.

## Description

The upper C (K4.14) is laterally compressed, has an anterior keel in a slightly lingual position, as well as a posterior one. Both keels are poorly preserved, but show crenulations at least near the tip of the canine. The length at the base of the canine is about 30.0 mm and the width about 11.6 mm . The resulting index ( $\mathrm{W} \times 100 / \mathrm{L}$ ) Cs is 38.7 , showing the degree of compression of the canine.

The mandible has a strong, downwardly expanded and angular mental crest. The height of the symphysis measured anteriorly is about 75 mm . The area of the symphysis just below the incisors and anteriorly is deeply grooved, while its lower part is smoother. The upper part of the symphyseal area is situated in front of the mental crests. There are two mental foramina. These foramina are on the same level and closer to the lower margin of the mandible than to the upper one. The anterior mental foramen opens under the middle of the postcanine diastema and the poste-


Figure 5 Logarithmic ratio diagram comparing the lower dentition of the Kerassia 4 Adcrocuta with A. eximia and C. bonisi from various localities. H. hyaena (standard) according to Howell \& Petter (1980). Data for the Ravin des Zouaves I \& 5, Dytiko I and Ravin de la Pluie hyaenids according to Koufos (2000). A. eximia from Halmyropotamos according to Melentis (I968). The rest from Table 6.
rior one just in front of p 3 . The masseteric fossa is deep and well defined by an acute crest on its upper part. It extends anteriorly to the posterior limit of ml on the right ramus, slightly more anteriorly on the left. The condyle is situated 1.5 cm lower than the level of the cheek teeth, while the incisors and the canine are clearly at a higher level. The mandibular foramen opens close to the lower margin of the mandible and about 65 mm in front of the condyle.

The lower incisors are arranged in an almost straight line, are conical in shape, increasing in size from i1 to i3 and situated clearly in front of the lower canines. Their tips are directed upwards. The total width of the incisors (i3-i3) measures about 43 mm . The lower canine is not significantly laterally compressed (Table 7). A large diastema separates the lower canine from p3 (Table 7) and a small diastema of 2.5 mm separates p3
from p 4 . The mandibular corpus, at the region of the diastema and on its upper part, is laterally concave and an acute crest is formed dorsally. There is no trace of a p2. The p 3 is elongated (Table 7). The right p 3 has a faint anterior accessory cusp and the left one a larger one. There is also a posterior accessory cusp that is larger than the anterior one. The greatest width of the tooth is in its posterior part. In p 4 , both the anterior and the posterior accessory cusps are strong, but the anterior one is thicker. Behind the posterior accessory cusp there is also a small supplementary cuspid. Both the posterior accessory cusp and the supplementary cuspid are directed slightly backwards. The greatest width of the tooth lies posteriorly. In m 1 , the paraconid is robust while the protoconid is more slender and slightly longer. There is a small metaconid. The greatest width of the tooth lies in the posterior part of the paraconid.

## Comparisons

Various species of Machairodus have been described from the Late Miocene of Europe and Asia, but De Beaumont (1975) retained only two species, the Vallesian Machairodus aphanistus (KaUP, 1833) and the Turolian Machairodus giganteus (WAGNER, 1848). These two species are of comparable dimensions, but differ in many morphological characters. De Beaumont (1975) considered the species Machairodus leoninus (Rотн \& Wagner, 1854), Machairodus tarakliensis Riabinin, 1929, Machairodus palanderi Zdansky, 1924, and possibly Machairodus tingii Zdansky, 1924, to be synonyms of $M$. giganteus. The available Kerassiá 4 specimens are compared to bibliographic data as well as to three undescribed specimens of $M$. giganteus from Pikermi stored in the AMPG (Table 7).
As is indicated by De Beaumont (1975) and Koufos (2000) the upper canine of M. aphanistus is less compressed than that of $M$. giganteus. According to Koufos (2000), the index of compression (Wx100/L) of the upper canine in M. aphanistus ranges from 41.5-52 and in M. giganteus from 38-41. The canine from Kerassiá 4 has an index of compression equal to 38.7 , well within the range of $M$. giganteus. An upper canine from Pikermi (AMPG no 1967/7) has an index of compression 38.8. This specimen is attributed by Melentis (1968) to M. aphanistus, but in our opinion must be attributed in M. giganteus. For the Machairodus skull from Halmyropotamos (Melentis 1968), the compression index of the upper canine is about 40.6. This specimen has been referred to M. aphanistus by Melentis (1968), but was considered to belong to M. giganteus by Petter \& Howell (1987) and Koufos (2000).

In lateral view, the lower incisors of $\mathrm{K} 4 / \Delta 69 / 1$ are clearly in front of the canines. They are arranged in an almost straight line, while in P.G. 01/100 from Pikermi the incisors are arranged in an arc. De Beaumont (1975) observed that in M. giganteus the lower incisors are arranged in an arc. It is not
known, however, on how many specimens this has been observed and how much variability is involved. Judging from the drawings of some mandibles in lateral view, the arrangement of the lower incisors varies in $M$. giganteus. On the mandible from Pikermi figured by Roth \& Wagner (1854, pl. 9, fig. 1 ), for example, i2 is only just in front of i3. On another mandible (Wagner 1857, pl. 5, fig. 11), i2 is significantly in front of i3. These observations show that the disposition of the lower incisors varies. The width of the lower incisor row (i3-i3) of K4/ $\Delta 69 / 1$ is 43 mm , only slightly different from P.G. 01/100 (estimated at 47 mm ). The height of the symphysis of P.G. $01 / 100$ is about 67 mm , slightly smaller than that of the Kerassiá mandible. De Beaumont (1975) mentioned that the postcanine diastema is generally large in $M$. giganteus (as also on the Machairodus from Kerassiá 4) and shorter in M. aphanistus, but he also mentioned that there is considerable variation in this character. On the type specimen of M. aphanistus from Eppelsheim, this diastema is $30-35 \mathrm{~mm}$ and in another specimen from the same locality, it is 35 mm (De Beaumont 1975). The length of this diastema varies significantly in $M$. giganteus from Pikermi. It is 55 mm in a specimen from Pikermi described by Roth \& Wagner (1854, pl. 9, fig. 1) and only 24 mm in another specimen from the same locality described by Wagner (1857, pl. 5, fig. 11). On P.G. 01/100 from Pikermi, this diastema is 48.8 mm , which is slightly less than that of $\mathrm{K} 4 / \Delta 69 / 1$ (Table 7). In P.G. 01/100, there is a minute alveolus for p 2 at the middle of the diastema. De Beaumont (1975) mentioned that in $M$. giganteus there is no trace of p 2 . In our opinion, however, such a character is not very important since, as we mention also later, it is sometimes present in Machairodus alberdiae Ginsburg, Morales \& Soria, 1981, as well as in species such as Metailurus parvulus (Hensel, 1862). In the Kerassiá 4 Machairodus, the width of the lower canines is much less than the distance that separates them from the symphysis. Such a character is
always found in M. giganteus while in $M$. aphanistus the opposite is observed (De Beaumont 1975). According to the same author, the lower canine of M. giganteus is less compressed than that of M. aphanistus. In a specimen of M. giganteus from Samos, the lower canine has dimensions $(\mathrm{L} \times \mathrm{W})$ at the base of the crown of $18.0 \times 14.0$ ( De Beaumont 1975), giving it an index of compression of 77.8. According to the measurements given by De Beaumont (1975), the lower canine of $M$. aphanistus from Eppelsheim described by Kaup (1833, pl. I, fig. 3) under the name "Agnotherium antiquит" has an index of 59.6, while in another specimen of M. aphanistus from Eppelsheim (BMNH 4996a) the corresponding index is 58.1. The index of compression of the lower canine of $\mathrm{K} 4 / \Delta 69 / 1$ is 74.7 , a value significantly greater than in M. aphanistus and closer to M. giganteus. In P.G. 01/100 from Pikermi, the lower incisor has an index of compression of 66.9. This value, however, is taken at the root of the tooth. Higher on the crown, we would expect a greater value, since the anteroposterior diameter (L) of the canine decreases more than the transverse diameter (W).

The lower premolars and the lower carnassial of M. giganteus show no significant differences from those of M. aphanistus. As shown in Figure 6, the cheek teeth vary significantly in size, something also mentioned by De Beaumont (1975). Aside from the size variation, De Beaumont (1975) also emphasized the morphological variation of M. giganteus, a variation that concerns the postcanine diastema that we mentioned above, the development of the anterior accessory cusp of p3, and the development of the metaconid/talonid complex of m 1 . As shown in Figure 6, the p3 and p 4 of $M$. giganteus are generally more elongated than they are in M. aphanistus. There are, however, specimens referred to $M$. aphanistus with p 3 and p 4 equally or almost equally elongated as those of M. giganteus, as for example the M. aphanistus from Charmoille and the type specimen from

Eppelsheim (Fig. 6). The p3 of P.G. 01/100 has a larger anterior accessory cusp than K4/ $669 / 1$, but De Beaumont (1975) mentioned that in M. giganteus the anterior cusp varies from faint to well developed. The accessory cusps of p 4 in $\mathrm{K} 4 / \Delta 69 / 1$ are not different from those of P.G. 01/100 and P.G. 01/101 from Pikermi, the anterior one being slightly more robust than the posterior one. In $\mathrm{K} 4 / \Delta 69 / 1$, the metaconid of M1 is slightly more developed than in P.G. 01/100, P.G. $01 / 101$ and P.G. $01 / 102$, in which feature it is more like two specimens of M. giganteus figured by De Beaumont (1975, figs. 6e, 6f). In M. giganteus, however, the m 1 is, in general, more slender than in M. aphanistus (Fig. 6).

The species M. pseudailuroides SchmidtKittler, 1976, from the localities of Akçakoy and Eskihisar in Turkey, differs from the Machairodus from Kerassiá in its shorter p4 and m 1 . The upper canine of this species is more slender than that of M. giganteus, with an index of compression of 35.9.
M. alberdiae Ginsburg, Morales \& Soria, 1981, from the Vallesian locality of Los Valles de Fuentidueña has been considered as a possible ancestor of M. giganteus (Ginsburg 1999). This species has smaller dimensions than the specimens described here and sometimes preserves a p2. Its upper canine is significantly less compressed ( $\mathrm{W} \times 100 / \mathrm{L}=50.0-53.9, \mathrm{n}=2$ ) than that of the Kerassiá 4 machairodont, but its lower canine is slightly more compressed ( $\mathrm{W} \times 100 / \mathrm{L}=63.4$ $71.4, \mathrm{n}=4$ ). Most characteristic of $M$. alberdiae is the large p 3 relative to p 4 and m 1 (Fig. 7).
Two machairodonts have been described from the Late Miocene of China, M. tingii and M. palanderi (Zdansky 1924). As we mentioned earlier, M. palanderi has been considered conspecific with M. gignateus by De Beaumont (1975), and the same may be true also of M. tingii. As we can see (Fig. 7) M. tingii is very similar in dimensions to K4/ $\times 69 / 1$, but $M$. palanderi has shorter p3 relative to p 4 and m 1 .


Figure 6 Logarithmic ratio diagram comparing the Kerassia 4 Machairodus with M. giganteus and M. aphanistus from various localities. M. aphanistus from Eppelsheim, Soblay and Charmoille according to Beaumont (1975), M. giganteus from Samos and M. ex gr. giganteus from Baccinello according to Rook et al. (I99I), M. giganteus from Ravin X according to Koufos (2000), M. giganteus from Pikermi according to the authors.

Sotnikova (1992) described a new species, M. kurteni, from the Late Miocene locality of Kalmakpai in Kazakhstan. According to Sotnikova (1992), this species has a proportionally narrower P4 than other species of Machairodus, retains a small P2, the tips of the lower incisors are directed upwards, the length of the lower premolars is small compared with that of m 1 , and has completely lost the metaconid/talonid complex on $\mathrm{M}_{1}$. Sotnikova (1992) did not support her statements with adequate comparisons, but actually compared her specimens to ' $M$. tarakliensis' from Taraklia, a form that has been considered a synonym of $M$. giganteus by De Beaumont (1975) and a subspecies of $M$. giganteus by Sotnikova (1992). Most of the above mentioned characters of M. kurteni are also found in M. giganteus. Concerning for example the upper teeth of the Kalmakpai
form, a compression index of P4 of 38.7 signifies that P 4 is not narrower than that of $M$. giganteus. The M. giganteus from Halmyropotamos, for example, has a compression index equal to 34.3 , which indicates a narrower P4 compared to the Kalmakpai form. The same is true for the Ravin X skull portion of M. giganteus, which has a compression index of P4 equal to 36.4 (Koufos 2000). A similar comparison shows that the P3 of the Kalmakpai form is not shortened relative to P 4 , more than it is in M. giganteus. Concerning the lower teeth, the index $\mathrm{Lm} 1 \times 100 / \mathrm{Lp} 3-4$, which shows the relative development of the premolars, varies in the Kalmakpai form from 67.7 to $74.5(\mathrm{n}=3)$. These values are not different from the Pikermi specimens P.G. 01/100 (73.2) and P.G. 01/101 (70.4). According to Sotnikova (1992), the tips of the lower incisors of the Kalmakpai form are
directed upwards, contrary to the Taraklia M. giganteus, where they are directed forwards. In P.G. 01/100, however, as well as on the mandible figured by Roth \& Wagner (1854, pl. 9, fig. 1), the tips of the lower incisors are also directed upwards. Other characters, such as the presence of P 2 and p 2 are not reliable alone, but only in combination with other characters, since they seem variable. Sotnikova (1992) for example figured a mandible of M. laskarevi from Kalfa, where there is a p2 on the right ramus but not on the left. The same tooth varies in development in $M$. alberdiae, being absent or rudimentary. The P2 is in general absent in M. giganteus, but present in M. palanderi and M. tingii. M. kurteni has completely lost its metaconid/talonid complex on m 1 . This character varies significantly in M. giganteus and can be almost absent (De Beaumont 1975). From the above observations is seen that, except for the last character, M. kurteni is not clearly different from M. giganteus and further comparison has to be made.
The available radius is comparable in size to that described by Gaudry (1863, pl. 16, figs. 2, 3). The radial tuberosity is not very prominent. The radius has two wide grooves in its anterodistal part, a medial one for the extensors carpi radialis longior and brevior indicis and a lateral one for the extensor communis digitorum. These two grooves are separated by a blunt crest. More medially lies the groove for the extensor ossis metacarpi pollicis and more laterally the triangular articular surface for the ulna. A proximal part of a radius from Halmyropotamos is also mentioned by Melentis (1969) under the name Machairodus aphanistus. In our opinion it is not certain that this specimen represents a Machairodus, since it has smaller dimensions. The maximum diameter of the head of that specimen is 23 mm (Melentis 1969), significantly smaller than in the radius of Kerassiá 4. The present Mt IV has a badly damaged shaft and its distal epiphysis is slightly broken. Its total length is 128 mm , and must be expected to have been slightly
greater. The anteroposterior diameter of the proximal part is 29 mm .
The species M. giganteus is known from the Turolian of Halmyropotamos (possibly MN11 or MN12), the early-middle Turolian (MN11-12) of Ravin X in Greece and from the middle Turolian (MN12) of Mont Leberon (France), Kalimanchi (Bulgaria), Samos, Pikermi (Greece) and Taraklia (Moldavia) (Ginsburg 1999, Koufos 2000). It is reported also under the name Amphimachairodus giganteus from various localities in Spain, ranging from MN11 to MN13 (Fraile et al. 1997). Rook et al. (1991) also mentioned some specimens under the name Machairodus ex gr. giganteus from the MN12 locality of Baccinello-V3 in Italy.

Subfamily: Felinae Trouessart, 1885
Genus: Metailurus Zdansky, 1924
Metailurus cf. parvulus (Hensel, 1862)
Locality: Kerassiá 1 (K1)

## Material

$\mathrm{K} 1 / \Delta 18$ : left mandible with ci, p3 and p4. The m 1 has fallen out and the tip of the canine is broken.

## Description

The mandibular corpus has a large mental foramen situated under the middle of the postcanine diastema and a very small one under the front end of p 4 . The lower canine is in a poor state of preservation, but shows an antero-lingual keel and most possibly a posterior one. The p 3 has only a posterior accessory cusp and its greatest width is in its posterior part. The p 4 has an anterior and a posterior accessory cusp, the posterior one being slightly stronger. As in p3, the greatest width occurs in the posterior part of the tooth. The alveolus for ml is especially large and its posterior limit lies well up on the ascending ramus of the mandible. Moreover, the mandibular corpus shows an abrupt thickening at the region of the m 1 . The above characters may indicate that our specimen is a pathological one.


Figure 7 Logarithmic ratio diagram comparing the Kerassia 4 Machairodus with other species. M. aphanistus from Eppelsheim (standard) according to Beaumont (I975), M. tingii and M. palanderi according to Zdansky (I924), M. alberdiae according to Ginsburg et al. (I98I), M. pseudailuroides according to Schmidt-Kittler (1976) and M. Kurteni according to Sotnikova (I992).

## Discussion and comparisons

The species M. parvulus (Hensel, 1862) has a wide range from Spain to China. Following Thenius (1951) and De Beaumont (1961), we include in this species the "Felidae indet., 3rd species" (Gaudry 1863), Felis leiodon Weithofer, 1888, Metailurus minor Zdansky, 1924, and other remains referred to under various specific or generic names.
The available specimen from Kerassiá 1 has been compared with specimens of $M$. parvulus from Pikermi, Chomateri and China. We have included also in our comparison an undescribed until now specimen from Pikermi, with the catalogue number AMPG P.G. 01/103 (Table 9). The mandible of $\mathrm{K} 1 / \Delta 18$ has the typical form of the Felinae,
without a mental crest. The height of the mandibular corpus in front of P3 is only slightly smaller than that of the Chomateri specimen (Symeonidis 1978) or P.G. 01/103 (Table 9). There is no alveolus for p 2 between the lower canine and the p3. Such an alveolus, with a single, very small root, exists on the specimen from Chomateri, on the specimen P.G. 01/103 from Pikermi, as well as in one specimen from China (Zdansky 1924, ex. 3).

Unfortunately, the teeth are not very well preserved and some of their measurements could not be taken with the appropriate precision. The p 3 is slightly smaller than other specimens of $M$. parvulus, but the indices of the teeth are within the range of M. parvulus


Figure 8 Logarithmic ratio diagram comparing the Kerassia I Metailurus with M. parvulus from Pikermi, Chomateri and China. Data from Table 9.
from China (Table 9). As we can see (Fig. 8), the Greek specimens show a tendency to smaller size than the Chinese specimens. Some of these specimens also show a narrower p 4 . The Kerassiá one specimen is slightly smaller in most of its dimensions than other specimens referred to $M$. parvulus, but the observed differences are of little value, as the statistical sample is not adequate. The present specimen, however, lacks the ml and it is preferable to refer it as Metailurus cf. parvulus.
Paramachairodus ogygius (Kaup, 1832) has teeth comparable is size with the largest specimens of M. parvulus from China. The Kerassiá one specimen, however, differs from P. ogygius. The height of the mandibular corpus in front of p 3 measures about 26 mm (measurement from the figure) on the specimen figured by Kaup (1833, pl. 2, fig. 3), as opposed to 17.5 mm on $\mathrm{K} 1 / \Delta 18$. Moreover,
in the type specimen of $P$. ogygius the postcanine diastema measures about 17 mm (De Beaumont 1975), as opposed to $8,5 \mathrm{~mm}$ in the Kerassiá one specimen.
M. parvulus is known from the Turolian of Halmyropotamos (possibly MN11 or MN12) (Melentis 1968), the middle Turolian (MN12) of Pikermi (Hensel 1862, Thenius 1951) and Chomateri (Symeonidis 1978) in Greece, Los Mansuetos in Spain (Morales \& Soria 1979, Fraile et al. 1997) and the late Turolian of El Arquillo in Spain (Morales \& Soria 1979, Fraile et al. 1997).

## GENERAL DISCUSSION AND CONCLUSIONS

The carnivores of K1, K3 and K4 are consistent with a Turolian age for these sites. $A$. eximia is very common in the middle Turolian, but rarer in the early or late Turolian (Howell \& Petter 1985). In Greece,
A. eximia seems absent from the late Turolian localities, such as those of Dytiko (De Bonis et al. 1992, Koufos 2000). The same is true for the rare M. giganteus, which is absent from Dytiko. Both species are present in the Main Bone Beds of Samos (Bernor et al. 1996), dated to 7.1 Ma (Swisher 1996), while $A$. eximia is also present in the older White Beds of Samos (Bernor et al. 1996; Swisher 1996). According to the recent calibration of the MN zones (Steininger 1999), the Main Bone Beds of Samos can be placed in MN12. In addition to the carnivores discussed here, the K1 faunal list (Theodorou et al. this volume) also includes a Tragoportax, provisionally referred to Tragoportax cf. amalthea. T. amalthea (Roth \& Wagner 1854) is a boselaphine that is well known from many Late Miocene localities, with a stratigraphic range from the early to the middle Turolian (Gentry et al. 1999).

The suids referred to M. major erymanthius by Van der Made \& Moyà-Solà (1989) most probably come from K1 and according to these authors indicate a middle Turolian age. Kostopoulos et al. (2001), however, have shown that the Kerassiá M. major is smaller than the Pikermi M. major erymanthius, and probably represents a new subspecies. The geological prospecting of the Kerassiá sites has shown that K1 is situated at a higher level than K3 and K4, so K1 must be slightly younger. K3 and K4 possibly belong to the same stratigraphic level, but no suids have been found in these sites until now (Theodorou et al. this volume).
A biostratigraphic distinction between K 1 , K 3 and K 4 is not possible at the moment, but judging from the above data, an early to middle Turolian age is plausible for the fauna of Kerassiá. In the future, we plan to continue the fieldwork, while sedimentological and taphonomic study of the numerous fossiliferous sites at Kerassiá is already in progress.

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PLATE I Plioviverrops sp., Kerassia 3. Fig. I: Left humerus (K3/BI/4), anterior view. Fig. 2: Head of a left humerus (K3.206), juvenile, proximal view. Fig. 3: Distal epiphysis of a right femur (K3/BI/I5), juvenile, distal view. Fig. 4: Left Mt IV (K3.I9I), lateral view. Fig. 5: Right calcaneum (K3.192) medial view. × I. cf. Ictitherium pannonicum, Kerassia 3. Fig. 6: Right mandibular ramus (K3/204), labial view. Fig. 7: idem, occlusal view. Fig. 8: idem, lingual view. $\times 3 / 4$.



PLATE 3 Adcrocuta eximia, Kerassia 4. Fig. I: Left mandibular ramus (K4/ $\Delta 388 / 2$ ), labial view. Fig. 2: idem, occlusal view. Fig. 3: idem, lingual view. $\times 3 / 4$.



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PLATE 5 Machairodus giganteus, Kerassia 4. Fig. I: Right upper canine (K4.I4), labial view ( $\times 1 / 2$ ). Fig. 3: Left radius ( $\mathrm{K} 4 / \Delta \mid \mathrm{I} / \mathrm{I} / \mathrm{I}$ ) anterior view ( $\times$ I/3). Machairodus giganteus, Pikermi. Fig. 2: Left upper canine (No 1967/7), lingual view ( $\times$ I/2). Fig. 4: Left mandibular ramus (P.G. $01 / \mathrm{IOO}$ ), labial view. Fig. $5:$ idem, occlusal view. $\times \mathrm{I} / 2$.


PLATE 6 Metailurus cf. parvulus, Kerassia I. Fig. I: Left mandibular ramus (KI/DI8), labial view. Fig. 2: idem, occlusal view. Fig. 3: idem, lingual view. x I. Metailurus parvulus, Pikermi. Fig. 4: Left mandibular ramus (P.G. 0I/IO3), labial view. Fig. 5: idem, occlusal view. $\times 1$.


Table I Measurements of the humerus. Lmax: maximum length; Lf: maximum functional length; DTpr: maximum transverse diameter of the proximal epiphysis; DAPpr: maximum anteroposterior diameter of the proximal epiphysis; DTdist: maximum transverse diameter of the distal epiphysis; DAPdist: maximum anteroposterior diameter of the distal epiphysis.

|  | Lmax | Lf | DTpr | DAPpr | DTdist | DAPdist |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: |
| Plioviverrops sp., K3.206, Kerassia 3 | - | - | 15.5 | 19.9 | - | - |
| Plioviverrops sp., K3/B1/4, Kerassia 3 | - | - | - | - | 15.3 | 9.7 |
| P. orbignyi, MNHNP PIK.3107, Pikermi | 93.8 | 90.5 | $(13.8)$ | 19.5 | 15.2 | $(10.5)$ |
| P. orbignyi, (Pilgrim 1931), Pikermi | $(81)$ | - | - | - | 14 | - |

Table 2 Measurements of the calcaneum. Hmax: maximum height; DTpr: maximum transverse diameter of the head; DTcol: transverse diameter at the middle of the neck; DTmax: maximum transverse diameter: DAPpr: maximum anteroposterior diameter of the head; DAPcol: anteroposterior diameter at the middle of the neck; DAPmax: maximum anteroposterior diameter.

|  | Hmax | DTpr | DTcol | DTmax | DAPpr | DAPcol | DAPmax |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Plioviverrops sp., K3.192, Kerassia 3 | 25.6 | 6.9 | 4.5 | +8.3 | 8.9 | 8.7 | 11.3 |
| P. orbignyi, MNHNP PIK. 3111, Pikermi | 24.2 | 6.5 | 4.5 | 10.1 | 8.8 | 9.0 | 10.0 |

Table 3 Measurements of the fourth metatarsal. Lmax: maximum length; DAPpr: maximum anteroposterior diameter of the proximal end; DAPdistart: anteroposterior diameter of the distal articular surface; DTpr: maximum transverse diameter of the proximal end; DTdistart: transverse diameter of the distal articular surface; DTdistmax: maximum transverse diameter of the distal end.

|  | Lmax | DAPpr | DAPdistart | DTpr | DTdistart | DTdistmax |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: |
| Plioviverrops sp., K3.161, Kerassia 3 | 48.5 | $(6.8)$ | $(4.5)$ | $(3.2)$ | 3.9 | 4.4 |
| Plioviverrops sp., K3.195, Kerassia 3 | 49.3 | 7.6 | 4.8 | $(3.5)$ | 3.8 | 4.3 |
| P. orbignyi, (Pilgrim 1931), Pikermi | 44 | - | - | - | - | - |

Table 4 Measurements of cf. I. pannonicum from Kerassia 3 and I. pannonicum from Polgardi and the Black Sea.

|  | cf. I. pannonicum K3.204, Kerassia 3 | I. pannonicum, Polgardi (Semenov 1985) |  |  | I. pannonicum, Black Sea (Semenov 1985, 1989) |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | n | mean | min-max | n | mean | min-max |
| $\mathrm{LC}_{\mathrm{i}}$ | (10.6) | - | - | - | 2 | 12.05 | 11.5-12.6 |
| $\mathrm{WC}_{i}$ | (8.1) | - | - | - | 3 | 7.83 | 7.5-8.3 |
| $\mathrm{LP}_{2}$ | 11.3 a | 3 | 12.90 | 12.7-13.0 | 2 | 13.40 | 13.3-13.5 |
| $\mathrm{WP}_{2}$ | 6.2 a | 1 | 6.2 | - | 2 | 7.00 | 6.9-7.1 |
| $\mathrm{LP}_{3}$ | 14.8 | 5 | 15.60 | 15.3-16.0 | 4 | 16.58 | 15.9-17.2 |
| $\mathrm{WP}_{3}$ | 8.1 | 1 | 7.5 | - | 4 | 7.72 | 7.0-8.3 |
| $\mathrm{LP}_{4}$ | 16.6 | 3 | 16.67 | 16.0-17.8 | 4 | 16.95 | 16.4-17.6 |
| $\mathrm{WP}_{4}$ | (7.7) | 2 | 8.40 | 8.2-8.6 | 4 | 9.05 | 8.3-9.8 |
| $\mathrm{LpP}_{4}$ | 8.2 | - | - | - | - | - | - |
| $\mathrm{LM}_{1}$ | 20.5 | 2 | 19.58 | 19.3-20.4 | 5 | 20.18 | 19.3-21.0 |
| $\mathrm{WM}_{1}$ | 9.0 | 1 | 9.7 | - | 5 | 9.16 | 8.4-10.0 |
| LtM ${ }_{1}$ | (14.9) | - | - | - | 5 | 14.66 | 13.9-15.4 |
| $\mathrm{LM}_{2}$ | 8.9 a | 1 | 9.6 | - | 3 | 9.63 | 9.4-9.8 |
| $\mathrm{WM}_{2}$ | - | 1 | 8.3 | - | 3 | 8.10 | 7.2-8.7 |
| $\left(\mathrm{WP}_{3} / \mathrm{LP}_{3}\right) \times 100$ | 54.7 | - | - | - | 4 | 46.55 | 44.1-48.2 |
| $\left(\mathrm{WM}_{1} / \mathrm{LM}_{1}\right) \times 100$ | 43.9 | - | - | - | 5 | 45.56 | 43.9-47.8 |
| $\left(\mathrm{LM}_{2} / \mathrm{LM}_{1}\right) \times 100$ | (43.4) | - | - | - | 3 | 46.8 | 46.2-47.3 |
| $\mathrm{LP}_{2}-\mathrm{M}_{1}$ | (64.0) | - | - | - | 3 | - | 64-68 |
| $\mathrm{LP}_{1}-\mathrm{M}_{2}$ | (79.0) | - | - | - | 1 | 77 | - |
| Hdia | 24.3 | - | - | - | - | - | - |
| $\mathrm{HbehM}_{1}$ | 28.7 | - | - | - | - | - | - |

Table 5 Measurements of A. eximia upper teeth from Kerassia and Pikermi. The data concerning the A. eximia from Pikermi are from Roussiakis (1996). These are based on the specimens mentioned by Howell \& Petter (I985, tab. 6a) as well as on AMPG P.A. 490/9I described by Roussiakis (1996) and original measurements of MNHNP PIK. 3000 and AMPG P.G. 95/I 507. The last specimen is mentioned as unnumbered by Howell \& Petter (1985, tab. 6a). The statistical data have been recalculated, since there are some statistical inaccuracies in Howell \& Petter (1985, tab. 7).

|  | A. eximia, Kerassia 4 |  |  | A. eximia, Pikermi |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  | K4.7 | n | mean | min-max |
|  | sin. | dext. |  |  |  |  |
| LP ${ }^{1}$ | 6.7 | - | - | 3 | 7.7 | 7.6-8.0 |
| WP ${ }^{1}$ | 6.4 | - | - | 3 | 7.4 | 7.0-8.0 |
| $\mathrm{LP}^{2}$ | (19.3) | - | - | 11 | 11.9 | 15.7-19.1 |
| $W^{2}{ }^{2}$ | (13.3) | - | - | 11 | 12.7 | 11.4-15.0 |
| $\mathrm{LP}^{3}$ | 22.5 | 22.1 | (24.6) | 12 | 22.8 | 20.8-25.0 |
| WP ${ }^{3}$ | 16.4 | 15.9 | - | 12 | 15.4 | 14.4-16.8 |
| LP ${ }^{4}$ | 36.2 | 36.1 | (41.1) | 11 | 37.4 | 34.6-40.0 |
| WP ${ }^{4}$ | 19.2 | 19.3 | (20.7) | 12 | 18.0 | 14.4-21.0 |
| $\mathrm{LpP}^{4}$ | 13.2 | 12.9 | (13.3) | 3 | 13.2 | 13.0-13.5 |
| $\mathrm{LmP}^{4}$ | 15.5 | 15.5 | (17.8) | 4 | 15.5 | 14.5-16.4 |
| $L^{1}{ }^{1}$ | 6.2 | - | - | 4 | 6.6 | 6.0-7.0 |
| WM ${ }^{1}$ | 12.9 | - | - | 4 | 13.4 | 13.0-14.0 |
| $\left(W^{2} / L^{2}\right) \times 100$ | (68.9) | - | - | 11 | 70.8 | 63.5-88.2 |
| $\left(W P^{3} / L^{3}\right) \times 100$ | 72.9 | 71.9 | - | 12 | 67.8 | 60.0-72.7 |
| $\left(\mathrm{WP}^{4} / \mathrm{LP}^{4}\right) \times 100$ | 53.0 | 53.5 | 50.4 | 11 | 48.3 | 41.6-52.5 |
| $\left(W P^{3} / W P^{4}\right) \times 100$ | 85.4 | 82.4 | - | 11 | 85.6 | 76.2-108.3 |
| $\left(\mathrm{LmP}^{4} / \mathrm{LP}^{4}\right) \times 100$ | 42.8 | 42.9 | (43.3) | 4 | 41.4 | 40.0-43.0 |
| $\left(\mathrm{LP}^{4} / \mathrm{LP}^{2+3}\right) \times 100$ | 86.2 | - | ) | 9 | 92.6 | 84.1-102.6 |
| C-C | 62.8 |  | - | - | - | - |
| P-P | 99.4 |  | - | - | - | - |

Table 6 Measurements of A. eximia lower teeth from Kerassia and Pikermi. The data concerning the A. eximia from Pikermi are from Roussiakis (1996).These are based on the specimens mentioned by Howell \& Petter (1985, tab. 6a) as well as on AMPG P.A. 57/9I, P.A. 445/9I and P.A. I296/9I described by Roussiakis (1996) and original measurements of MNHNP PIK. 300I, AMPG P.G. 95/I505 and P.G. 95/I506. The last two specimens are mentioned as No 113 and unnumbered respectively, by Howell \& Petter ( 1985, tab. 6a). The statistical data have been recalculated since there are some statistical inaccuracies in Howell \& Petter (I985, tab. 7).

|  | A. eximia, Kerassia 4 |  | A. eximia, Pikermi |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: |
|  | K4/ $\Delta 388 / 2$ | K4/ $\Delta 388 / 3$ | n | mean | min-max |
| $\mathrm{LP}_{1}$ | 5.0 | - | 6 | 5.7 | 5.5-9.1 |
| $\mathrm{WP}_{1}$ | 5.4 | - | 6 | 5.8 | 5.0-7.0 |
| $\mathrm{LP}_{2}$ | 16.1 | 16.0 | 13 | 16.6 | 15.0-17.4 |
| $\mathrm{WP}_{2}$ | 11.1 | 11.2 | 12 | 11.5 | 10.3-12.8 |
| $\mathrm{LP}_{3}$ | 19.4 | 19.5 | 12 | 19.8 | 18.6-22.0 |
| $\mathrm{WP}_{3}$ | 13.7 | 13.1 | 12 | 13.7 | 12.6-15.1 |
| $\mathrm{LP}_{4}$ | 21.4 | 21.5 | 11 | 22.3 | 20.9-24.0 |
| $\mathrm{WP}_{4}$ | 12.9 | 12.9 | 10 | 13.3 | 12.3-14.0 |
| $\mathrm{LM}_{1}$ | 27.7 | (29.0) | 10 | 27.4 | 25.9-29.7 |
| $\mathrm{WM}_{1}$ | 13.1 | 13.0 | 10 | 12.7 | 11.5-13.7 |
| $\mathrm{LtM}_{1}$ | 22.8 | 24.0 | 7 | 22.8 | 21.8-24.0 |
| $\left(\mathrm{WP}_{2} / \mathrm{LP}_{2}\right) \times 100$ | 68.9 | 70.0 | 12 | 69.7 | 64.7-75.7 |
| $\left(\mathrm{WP}_{3} / \mathrm{LP}_{3}\right) \times 100$ | 70.6 | 67.2 | 12 | 69.4 | 63.6-76.3 |
| $\left(\mathrm{WP}_{4} / \mathrm{LP}_{4}\right) \times 100$ | 60.3 | 60.0 | 10 | 60.2 | 53.7-65.4 |
| $\left(\mathrm{WP}_{3} / \mathrm{WP}_{4}\right) \times 100$ | 106.2 | 101.6 | 10 | 104.4 | 100.0-110.6 |
| $\left(\mathrm{LtM}_{1} / \mathrm{LM}_{1}\right) \times 100$ | 82.3 | 82.7 | 7 | 82.6 | 80.8-84.2 |
| $\left(\mathrm{LM}_{1} / \mathrm{LP}_{4}\right) \times 100$ | 129.4 | (134.9) | 8 | 126.9 | 117.7-133.3 |
| $\left(\mathrm{LM}_{1} / \mathrm{LP}_{2+3+4}\right) \times 100$ | 48.7 | 50.9 | 8 | 47.96 | 44.7-50.9 |
| $\mathrm{LP}_{2}-\mathrm{M}_{1}$ | 81.2 | (84.0) | - | - | - |
| $\mathrm{HbehM}_{1}$ | 43.3 | 44.1 | - | - | - |
| Hdia | 41.2 | 41.2 | - | - | - |

Table 7 Measurements of M. giganteus from Kerassia 4 and Pikermi

|  | M. giganteus, K4/ <br> sin. | $c$ <br> dext. | AMPG P.G.01/100 | M. giganteus, Pikermi |  |
| :--- | :---: | :---: | :---: | :---: | :---: |
|  | AMPG P.G.01/101 | AMPG P.G.01/102 |  |  |  |
| $(\mathrm{L} \times \mathrm{W}) \mathrm{C}_{\mathrm{i}}$ | $15.0 \times 11.2$ | $-\times 10.9$ | $(17.3 \times 11.6) \mathrm{r}$ | - | - |
| $(\mathrm{W} \times 100 / \mathrm{L}) \mathrm{C}_{\mathrm{i}}$ | 74.6 | - | $(66.9) \mathrm{r}$ | - | - |
| $(\mathrm{L} \times \mathrm{W}) \mathrm{P}_{3}$ | $(18.7 \times 9.2)$ | $18.8 \times 9.0$ | $17.8 \times 7.7$ | $(16.4) \times 7.9$ | - |
| $(\mathrm{L} \times \mathrm{W}) \mathrm{P}_{4}$ | $-\times 12.8$ | $28.3 \times 12.9$ | $(27.0) \times 11.2$ | $28.8 \times 11.6$ | $-\times 11.3$ |
| $(\mathrm{~L} \times \mathrm{W}) \mathrm{M}_{1}$ | $-\times(14.2)$ | $35.6 \times 15.1$ | $32.5 \times 12.7$ | $31.4 \times 12.1$ | $32.0 \times 13.9$ |
| $\mathrm{LP}_{3}-\mathrm{M}_{1}$ | $(77.2)$ | 81.4 | 75.7 | $(73.2)$ | - |
| $\mathrm{LP}_{3-4}$ | 47.0 | 48.4 | 44.4 | 44.6 | - |
| ${\text { diastema } \mathrm{C}_{\mathrm{i}}-\mathrm{P}_{3}}^{\mathrm{C}_{\mathrm{i}}-\text { cond }}$ | 60.2 | 58.0 | 48.8 | - | - |
| Hdia | 240.6 | 244.0 | - | - | - |
| HbehM | 42.3 | 43.3 | 39.9 | 40.1 | - |

Table 8 Measurements of the radius. Lmax: maximum length; DTpr: maximum transverse diameter of the proximal epiphysis; DTcol: transverse diameter of the neck; DAPpr: maximum anteroposterior diameter of the proximal epiphysis; DTdia: transverse diameter at the middle of the diaphysis; DAPdia: anteroposterior diameter at the middle of the diaphysis; DTdist: maximum transverse diameter of the distal epiphysis; DAPdist: maximum anteroposterior diameter of the distal epiphysis.

|  | Lmax | DTpr | DTcol | DAPpr | DTdia | DAPdia | DTdist | DAPdist |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| K4/D110/16, Kerassia 4 | 306.1 | 38.1 | 26.2 | 31.5 | 35.8 | 20.8 | 59.9 | 40.6 |
| MNHNP PIK.3236, Pikermi | - | 37.6 | 24.6 | 29.0 | - | - | - | - |

Table 9 Measurements of M. cf. parvulus from Kerassia I and M. parvulus from Pikermi, Chomateri and China.

|  | M. cf. parvulus K1/ $\Delta 18$ Kerassia 1 | M. parvulus |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | Pikermi | Pikermi | P.G. 01/103 | Chomateri | China |
|  |  | (Weithofer 1888) | (Thenius 1951) | Pikermi |  | (Zdansky 1924) |
| $\mathrm{LC}_{\mathrm{i}}$ | (7.8) | 9.1 | - | 8.1 | 8.3 | 8.8-9.6 (n=5) |
| $\mathrm{WC}_{i}$ | (5.8) | 6.6 | - | 5.7 | 6.0 | $6.3-(7.4)(\mathrm{n}=5)$ |
| diastema $\mathrm{C}_{\mathrm{i}}-\mathrm{P}_{3}$ | 8.5 | 7.5 | 9.2 | 8.9 | 8.6 | 6.8-(11.0) ( $\mathrm{n}=5$ ) |
| $\mathrm{LP}_{3}$ | (9.4) | - | 10.3 | 9.7 | 9.9 | 9.9-10.7 ( $\mathrm{n}=3$ ) |
| $\mathrm{WP}_{3}$ | (4.6) | - | 5.1 | 4.9 | 5.1 | 5.1-5.3 ( $\mathrm{n}=3$ ) |
| $\mathrm{LP}_{4}$ | (13.3) | 15.0 | 14.4 | 14.2 | 13.9 | $14.5-15.4(\mathrm{n}=4)$ |
| $\mathrm{WP}_{4}$ | 6.0 | 6.5 | 5.9 | 5.7 | 5.8 | 6.1-6.9 ( $\mathrm{n}=4$ ) |
| $\mathrm{LM}_{1}$ | - | 17.8 | 17.2 | 17.3 | 17.1 | 16.7-18.1 ( $\mathrm{n}=4$ ) |
| $\mathrm{WM}_{1}$ | - | 7.0 | 6.9 | 6.9 | 7.0 | 6.9-7.8 ( $\mathrm{n}=4$ ) |
| $\mathrm{C}_{\mathrm{i}}$-cond | - | - | - | - | 101.2 | - |
| Hdia | (17.5) | - | - | 18.8 | 18.9 | - |
| $\mathrm{HbehM}_{1}$ | - | - | - | 20.1 | 19.4 | - |
| $\left(\mathrm{WC}_{i} / \mathrm{LC}_{\mathrm{i}}\right) \times 100$ | (74.4) | 72.5 | - | 70.4 | 72.3 | 69.2-(77.1) $(\mathrm{n}=5)$ |
| $\left(\mathrm{WP}_{3} / \mathrm{LP}_{3}\right) \times 100$ | (48.9) | - | 49.5 | 50.5 | 51.5 | 48.6-53. 5 ( $\mathrm{n}=3$ ) |
| $\left(\mathrm{WP}_{4} / \mathrm{LP}_{4}\right) \times 100$ | 45.1 | 43.3 | 41.0 | 40.1 | 41.7 | 42.1-45.4 $(\mathrm{n}=4)$ |
| $\left(\mathrm{WM}_{1} / \mathrm{LM}_{1}\right) \times 100$ | - | 39.3 | 40.1 | 39.9 | 40.9 | 39.8-45.1 ( $\mathrm{n}=4$ ) |
| $\left(\mathrm{LP}_{3} / \mathrm{LP}_{4}\right) \times 100$ | (70.7) | - | 71.5 | 68.3 | 71.2 | 68.3-70.3 ( $\mathrm{n}=3$ ) |
| $\left(\mathrm{LP}_{4} / \mathrm{LM}_{1}\right) \times 100$ | - | 84.3 | 83.7 | 82.1 | 81.3 | 80.1-86.8 ( $\mathrm{n}=3$ ) |
| $\underline{\left(\mathrm{LM}_{1} / \mathrm{LP}_{3+4}\right) \times 100}$ | - | - | 69.6 | 72.4 | 71.8 | 67.6-74.2 ( $\mathrm{n}=3$ ) |

