

The Vallesian large *Palaeotragus* Gaudry, 1861 (Mammalia: Giraffidae) from Northern Greece

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The Vallesian large *Palaeotragus* Gaudry, 1861 (Mammalia: Giraffidae) from Northern Greece

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ABSTRACT

We herein re-discuss the systematics of the Late Miocene representatives of the most common but poorly documented Eurasian giraffid genus *Palaeotragus* on the occasion of the review and description of new samples from the Vallesian faunas of Northern Greece. Our results detect five Late Miocene *Palaeotragus* morphotypes, recognizing at least four species. The so called ‘small-sized palaeotrages’ are represented by the type species *P. rouenii* and the Chinese *P. microdon*, whereas the validity of *P. pavlowae* from Grebeniki (Ukraine) is doubted. ‘Large-sized palaeotrages’ are mainly represented by *P. coelophrys* (synonym of *P. expectans*, *P. borissiaki*, *P. hoffstetteri*, *P. quadricornis*, and probably *P. moldavicus*), a species that thrived in the peri- Black Sea territories during Vallesian and survived during Turolian in the Irano-Anatolian domain, likely by adopting a more robust appearance. Large palaeotrages from the Vallesian faunas of Pentalophos and Ravin de la Pluie (Axios Valley, Greece) are identified as *P. coelophrys* with certain confidence. The Vallesian *P. berislavicus* from Berislav (Ukraine) has intermediate morphometric features between *P. rouenii* and *P. coelophrys* and it is, therefore, recognized as a most probably valid species. The latest Vallesian Nikiti-1 (Chalkidiki peninsula) large palaeotrage shares many morphometric features with *P. berislavicus*, suggesting that the species may have invaded Balkans by the end of Vallesian and possibly survived there until middle Turolian. The Late Miocene *Palaeotragus asiaticus* from Central Asia is a quite problematic species; it appears closely related to the Turolian equivalent *P. cf. coelophrys* from China and both may be linked to the older Berislav taxon.

KEY WORDS

Late Miocene,
Palaeotragiinae,
giraffes,
Europe.

RÉSUMÉ

Les Palaeotragus Gaudry, 1861 (Mammalia: Giraffidae) du Vallésien de Grèce du Nord.

Nous re-discutons ici de la systématique des représentants du Miocène supérieur du genre de girafide eurasien le plus commun mais mal documenté, *Palaeotragus* à l'occasion de révision et de la description originale de matériel vallésien du nord de la Grèce. Nos résultats permettent de détecter cinq morphotypes de *Palaeotragus* du Miocène supérieur, reconnaissant au moins quatre espèces. Les « paléotrages de petite taille » sont représentés par l'espèce type *P. rouenii* et *P. microdon* chinoise, alors que la validité de *P. pavlowae* de Grebeniki, Ukraine est mise en doute. Les « paléotrages de grande taille » sont principalement représentés par *P. coelophrys* (synonyme de *P. expectans*, *P. borissiakii*, *P. hoffstetteri*, *P. quadricornis*, et probablement *P. moldavicus*), une espèce qui a prospéré dans les territoires péri-Mer Noire pendant Vallesian et a survécu au Turolien dans le domaine irano-anatolien, probablement en adoptant une apparence plus robuste. De grands paléotrages des faunes vallésiennes du Pentalophos et du Ravin de la Pluie (vallée d'Axios, Grèce) sont attribués à *P. coelophrys* avec une certaine confiance. Le vallésien paléotrage de Berislav (*P. berislavicus*) d'Ukraine présente des caractéristiques morphométriques intermédiaires entre *P. rouenii* et *P. coelophrys* et il est donc reconnu comme une espèce très probablement valide. Le grand paléotrage de Nikiti-1 (péninsule de Chalcidique) présente des ressemblances importantes avec *P. berislavicus*, ce qui suggère que l'espèce pourrait envahir les Balkans à la fin du Vallésien et y survivre jusqu'au Turolien moyen. *Palaeotragus asiaticus* d'Asie centrale est une espèce assez problématique; il apparaît étroitement lié à *P. cf. coelophrys* de Chine et les deux peuvent être liés à l'ancien taxon de Berislav.

MOTS CLÉS
Miocène supérieur,
Palaeotragiinae,
girafes,
Europe.

INTRODUCTION

Palaeotragiinae is probably a polyphyletic or paraphyletic subfamily including species of the genus *Palaeotragus*, and according to some authors members of *Samotherium* too (Geraads 1986; Godina 2002; Hou *et al.* 2014; Danowitz *et al.* 2015). A phylogenetic parsimony analysis by Ríos *et al.* (2017: 31) restricted on the most well-known species supports, however, the genus monophyly. *Palaeotragus* was the most common giraffid genus of Eurasia during the Late Miocene, expanded from North Africa to the Black Sea and from the Balkans to China. It achieved maximal dispersion and diversity in the Turolian mammal communities of Eastern Mediterranean.

Numerous species have been ascribed to *Palaeotragus* from the Miocene of the Old World, most of them from the Late Miocene of W. Eurasia. The type species is *Palaeotragus rouenii* Gaudry, 1861, originally from Pikermi (Greece), archetype of the so-called 'small-sized palaeotrages'. Another widely accepted taxon is *Palaeotragus coelophrys* (Rodler & Weithofer, 1890) originally from Maragheh (Iran), usually regarded as the basic model of a 'large-sized palaeotrage'. Other Eurasian Late Miocene palaeotragine taxa include: *Palaeotragus microdon* (Koken, 1885) from Shansi (China); *Palaeotragus asiaticus* Godina, 1975 from Ortok (Kyrgyzstan); *Palaeotragus borissiakii* (Alexeev, 1930) from Eldari (Georgia); *Palaeotragus expectans* (Borissiak, 1914) from Sevastopol, *Palaeotragus pavlowae* (Pavlow, 1913) from Grebeniki, *Palaeotragus berislavicus* Korotkevitch, 1957 from Berislav (Ukraine); *Palaeotragus moldavicus* Godina, 1979 from Starýe Bogeny (Moldova); *Palaeotragus hoffstetteri* Ozansoy, 1965 from Sinap (Turkey); and *Palaeotragus quadricornis*

Bohlin, 1926 from Samos (Greece). The validity, generic affiliations and supra-generic relationships of most of these species are, however, strongly debated (e.g., Geraads 1974, 1986; Godina 1975, 1979; Ríos *et al.* 2017).

In Greece, *Palaeotragus* oldest record is from the Vallesian faunas of Axios Valley (Koufos 2006), whereas its last occurrence is from the Early Pleistocene faunas of continental Greece and Lesvos Island (Steensma 1988; Kostopoulos & Koufos 1994; de Vos *et al.* 2002; Athanassiou 2014). During the Turolian, the genus was mainly represented in the local mammal assemblages by the small and slender *P. rouenii* (Iliopoulos 2003; Koufos 2006; Kostopoulos 2009; Koufos *et al.* 2016; Lazaridis 2015; Xafis *et al.* 2019). Late Miocene large sized palaeotrages are rather scarce in the Greek record (Fig. 1), including sparse findings from the Vallesian faunas of Pentalophos, Xirochori, Ravin de la Pluie (Axios valley) and Nikiti-1 (Chalkidiki peninsula), as well as from the Turolian faunas of Kerassia, Samos and Thermopigi (Geraads 1979; Iliopoulos 2003; Kostopoulos 2009; Xafis *et al.* 2019).

The main aim of this study is the taxonomic review of the Vallesian large sized *Palaeotragus* from Northern Greece. The studied material comes from the Upper Miocene fossil sites of Nikiti-1 (NKT) in the Chalkidiki peninsula, and Pentalophos (PNT), Ravin de la Pluie (RPL) and Xirochori (XIR) at the Axios Valley (Fig. 1). The geological, stratigraphic and chronological settings of both fossiliferous areas are provided by Koufos (1990, 2016), Koufos *et al.* (1991), and Sen *et al.* (2000). The PNT material was previously referred to *Palaeotragus coelophrys* (Koufos 2006; Konidaris 2013) but never described; the associated fauna is suggested as of early late Vallesian, (latest MN9; Koufos 2006; Konidaris

2013). The large-sized *Palaeotragus* material from RPI was attributed to *Palaeotragus* cf. *coelophrys* by Geraads (1978), whereas the poor XIR material remained unpublished until now; faunal assemblages of both RPI and XIR are suggested as late Vallesian (MN10; Koufos 2006; Konidaris 2013). The NKT material was originally described as *Palaeotragus* cf. *rouenii* (Kostopoulos *et al.* 1996) and later considered as *Palaeotragus* sp. (Koufos *et al.* 2016); the NKT fauna is suggested as of latest Vallesian age (latest MN10; Koufos *et al.* 2016). The revision of the Greek Vallesian large-sized *Palaeotragus* leads to the necessary reassessment of the systematics of Late Miocene Eurasian large sized *Palaeotragus*.

MATERIAL AND METHODS

We follow the nomenclature proposed by Gentry *et al.* (1999) for the adult dental material; Geraads *et al.* (2013) for the deciduous lower premolars; Ríos *et al.* (2016) for the metapodials; Solounias & Danowitz (2016) for the astragali; and Schmid (1972) for the other postcranials. Postcranial measurements are according to the system proposed by von der Driesch (1976).

In order to test possible groupings within Late Miocene *Palaeotragus* species we performed Principal Coordinates Analysis (PCO) with Gower similarity, and Principal Components Analysis (PCA) based on 12 dental metrical features and indices, 15 postcranial measures and indices and 4 qualitative characters of P₃ and P₄ morphology (see Appendix 1). As available material appears fragmentary in several cases, the original dataset suffers from a rather high number of missing values. They are treated by iterative imputation option in PCA, whereas a series of subsequent analyses of both PCA and PCO were run by omitting taxa and/or parameters to test consistency of the results. A set of scatter diagrams based on selected and better documented variables associates multivariate analyses.

The PAST software (Hammer *et al.* 2001) was used for the statistical analyses. The program Inkscape was used for the illustrations provided. The program Adobe Photoshop was used for the processing of the studied material from NKT, PNT and RPI. All studied material is housed in the Museum of Geology-Palaeontology-Palaeoanthropology of the Aristotle University of Thessaloniki (LGPOT).

ABBREVIATIONS

AMNH	American Museum of Natural History, New York;
LGPOT	Museum of Geology – Palaeontology – Palaeoanthropology, Aristotle University of Thessaloniki;
NHM	Natural History Museum, Izmir;
MNHN	Muséum national d'Histoire naturelle, Paris;
OSU	Odessa State University, Odessa;
PIN	Palaeontological Institute, Moscow;
RSGU	Russian State Geological Prospecting University, Moscow;
SIZK	Schmalhausen Institute of Zoology of National Academy of Sciences of Ukraine, Kyiv;
SIT	Thermopigi Local Palaeontological Collection, Serres Prefecture, Greece.



FIG. 1. — Occurrences of large sized *Palaeotragus* Gaudry, 1861 in Greece: 1, Thermopigi; 2, Nikiti-1; 3, Pentalophos; 4, Xirochori; 5, Ravin de la Pluie; 6, Kerassia; 7, Mytilinii, Samos. Red circles, Vallesian occurrences; green circles, Turolian occurrences.

SYSTEMATIC PALAEONTOLOGY

Order ARTIODACTYLA, Owen 1848

Family GIRAFFIDAE Gray, 1821

Subfamily PALAEOTRAGIINAE Pilgrim, 1911

Genus *Palaeotragus* Gaudry, 1861

TYPE SPECIES. — *Palaeotragus rouenii* Gaudry, 1861 by original designation.

TYPE LOCALITY. — Pikerimi, Greece.

AGE. — Late Miocene to Early Pleistocene.

GENERIC DIAGNOSIS (based on Geraads 1974; Churcher 1978). — Giraffids of small to medium size. Skull wide with long postorbital cranial part. Ossicones simple, straight to slightly curved with smooth surface, single paired, pointed, uprightly inserted on frontals in supraorbital position, widely spaced on the cranial roof, and subparallel to each other. Frontals concave between the ossicones. Diastema long; dentition brachydont, premolar section proportionally long. Metapodials elongate, slender with moderately to strongly deep central palmar/plantar through.

DESCRIPTION

Pentalophos sample (end MN9)

Studied material (LGPOT, *Palaeotragus coelophrys*). Right maxilla with P₂-M₃ (PNT-113); part of left maxilla with M₂-M₃ (PNT-165); upper right P₂ (PNT-161); upper right P₂-P₃ (PNT-162); upper right P₂-P₃ (PNT-163); upper left P₃ (PNT-164); part of right mandible with M₁-M₃ (PNT-328F); part of right mandible

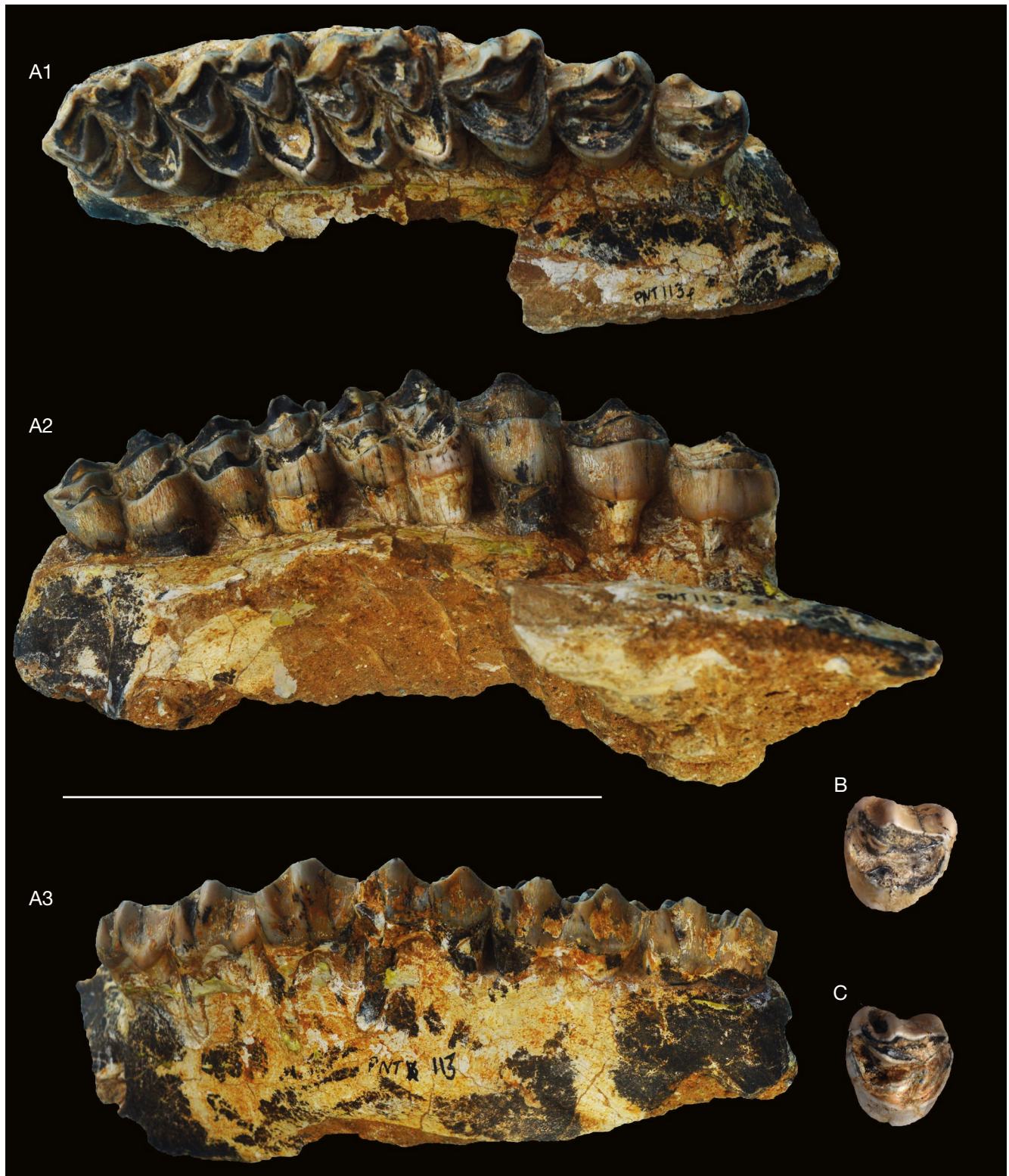


FIG. 2. — *Palaeotragus coelophrys* (Rodler & Weithofer, 1890) from Pentalophos, Axios Valley, Greece (late early Vallesian): **A**, upper tooththrow LGPUT PNT-113F: **1**, occlusal; **2**, lingual; **3**, labial views; **B**, upper P² (LGPNT PNT-161) in occlusal view; **C**, upper P³ (LGPNT PNT-164) in occlusal view. Scale bar: 10 cm.

with dP₂-M₁ (PNT-121F); distal part of left humerus (PNT-166); proximal parts of a right and a left metatarsal (PNT-114F, PNT-119F). The taxon is represented by at least four adult and a juvenile individual. Measurements are provided in Appendix 2: Tables 1-4.

Upper dentition. All the labial ribs and styles of the premolars (parastyle, paracone, metastyle) are well developed (Fig. 2A-C; Table 1). The hypocone is weak, but distinct. The enamel is finely rugose especially on the lingual side

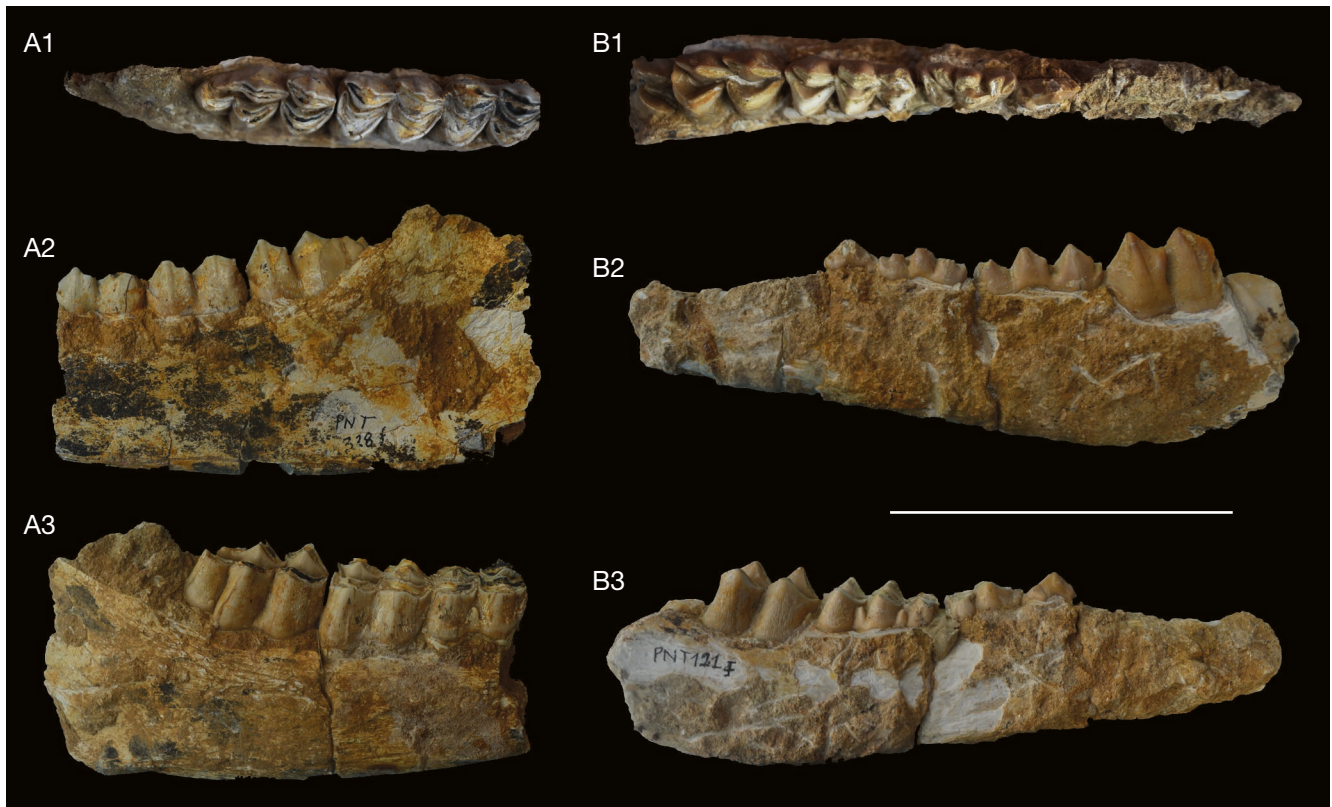


FIG. 3. — *Palaeotragus coelophrys* (Rodler & Weithofer, 1890) from Pentalophos, Axios Valley, Greece (late early Vallesian): **A**, part of mandible with M_1 - M_3 , LGPUT PNT-328F; **B**, part of deciduous mandible with dP_2 - M_1 , LGPUT PNT-121F; **1**, occlusal; **2**, lingual; **3**, labial views. Scale bar: 10 cm.

of the premolars. In some of the premolars there is a well-developed lingual cingulum (PNT-161, PNT-164; Fig. 2B, C). In the preserved P^2 and P^3 (Fig. 2A-C) the parastyle and the paracone rib are close to each other and the enamel is folded in the area of the parastyle towards the paracone. Moreover, these teeth are inflated at the basis, lingually rounded and they delineate occlusally an almost half circle. The P^2 of the PNT-113 tooththrow (Fig. 2A) has a prominent disto-labial style that reaches almost $\frac{1}{3}$ of the crown's height. In the single available P^4 , the paracone is more centrally placed, still slightly folded towards the parastyle, and the tooth has a triangular occlusal shape. The fossettes of the P^2 are distorted, but they look wide. The fossettes of the P^3 and P^4 are also wide and bear a hypoconal fold distally.

The upper molars are similar to those of other giraffids (Fig. 2A). The most prominent labial features are the parastyle, the paracone rib and the mesostyle. The metastyle is barely developed in the M^1 - M^2 , but similarly strong to the parastyle in the M^3 . Lingually, the protocone is more prominent than the hypocone, especially in the M^3 ; hence the mesial lobe is wider and square shaped whereas the distal lobe is narrower and rounder. The fossettes converge towards the center of the tooth but they do not fuse. The hypocone flange reaches almost at the labial side of the molar, stopping just before the mesostyle mesially and the metastyle distally. The protocone flange almost reaches the parastyle mesially. Distally, it reaches almost at the

center of the tooth. The enamel is finely rugose in all the molars, which also have a fine cement cover, mainly labially. Although in a less advanced wear stage, PNT-165 upper molars share the same basic morphological features with those of PNT-113. A lingual cingulum, occurs, especially on the mesial lobe.

Lower dentition. The right mandible PNT-328F preserves only the molars (Fig. 3A; Table 2). The mandibular height is 48.9 mm at the M_1 - M_2 level, and 53.6 mm below the distal lobe of M_3 . All the lower molars preserve a fine cement cover. The most prominent lingual feature is the metastylid, and then the metaconid; the entoconid is traceable but less strong. In the disto-lingual side of the M_1 and M_2 the entoconulid is present but not prominent. The entoconulid of the M_2 is completely separated from the hypocone region in the upper part of the crown. The same applies probably for the entoconulid of M_1 . The praentocristid penetrates in the mesial fossette, separating the labial and lingual side of the mesial lobe. The protoconid and the hypoconid are almost equally developed. An ectostylid is present on M_1 , less developed on M_2 , and absent on M_3 . Finally, the hypoconulid is pointed labially and parallel to the protoconid and hypoconid. Lingually, the third lobe of the M_3 is separated from the second one by a shallow furrow. The cingulum appears strong on M_1 , but weak on M_2 and absent on M_3 .

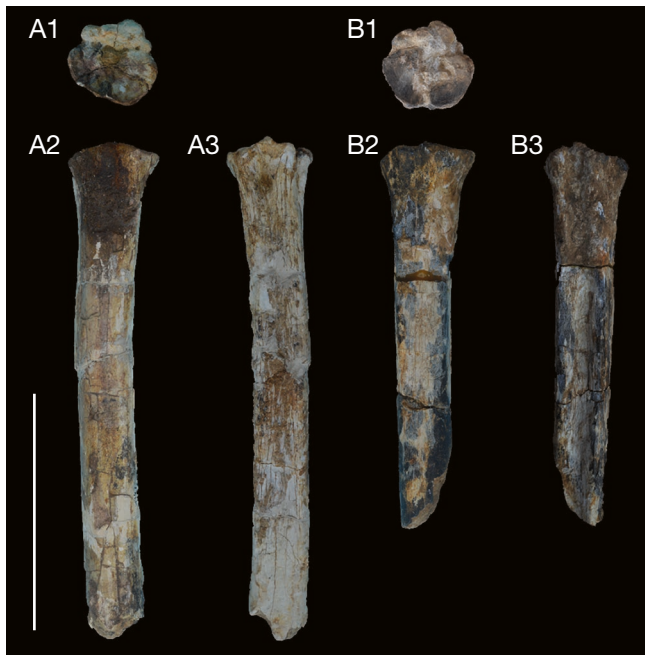


FIG. 4. — Partial metatarsals of *Palaeotragus coelophrys* (Rodler & Weithofer, 1890) from Pentalophos, Axios Valley, Greece (late early Vallesian): **A**, LGPUT PNT-114F; **B**, LGPUT PNT-119F; **1**, proximal; **2**, dorsal; **3**, plantar views. Scale bar: 15 cm.

Lower deciduous dentition. The right mandible PNT-121F (Fig. 3B; Table 3) consists of the series of the deciduous premolars and the M_1 . The dp_2 has a primitive morphology. The paraconid and anteparaconid are barely distinguished from each other. The protoconid is placed medially and it is the more developed conid. Distally, the entoconid and the hypoconulid are distinguished in the upper part of the crown, whereas they fuse in the lower. The hypoconid is prominent, placed more labially.

On the dp_3 the paraconid and anteparaconid are pointed lingually and they are well-distinguished in the upper half of the crown, whereas they fuse each other towards the base. The metaconid and the protoconid are the most prominent features (especially the protoconid); they fuse distally and they are folded and oriented mesially. The mesosinusid is lingually open. The distal and mesial part of the dp_3 are clearly separated in the area of the metaconid and protoconid fusion. The hypoconulid is as developed as the protoconid. The hypoconulid and the entocristid are distinguished only in the upper part of the crown. The metasinusid is clear but weaker than the mesosinusid.

The dp_4 has two prominent ectostylids. The mesial lobe is distorted, although it seems that its ribs and stylids are weaker. In the two distal lobes the labial cones are equally developed. The lingual ribs and stylids are weaker too, as in the mesial lobe. The distal flange of the third labial cone reaches the lingual side of the tooth. The distal flange of the second labial cone is also well-developed and it is placed next to the metastylid; it is the feature that separates the second from the third lobe.

The M_1 morphology agrees with the morphology of the M_1 from the adult tooththrow PNT-328F: the metaconid, metastylid, entoconid and entoconulid ribs are quite well-developed in the upper half of the crown.

Postcranials. The distal part of the left humerus PNT-166 is badly preserved prohibiting a detailed morphological description. The olecranon fossa is rather long, narrow, and deep. The lateral supracondyloid crest is well developed and raises quite high on the distal diaphysis. Both the radial and coronoid fossae are shallow and weakly separated by each other (but also probably due to the bad preservation). The distal trochlea is asymmetric and its keel wide and rather blunt. In lateral view, the lateral epicondyle slopes weakly downwards, whereas in medial view the medial epicondyle is perpendicular compared to the diaphysis longitudinal axis and not significantly protruding.

The preserved proximal parts of the metatarsals PNT-114F and PNT-119F (Fig. 4A, B; Table 4) indicate rather long but relatively robust metapodials. On the proximal articular surface, the plantar and dorsal heads of the medial epicondyle are separated by a bone protrusion, whereas a groove separates the two heads on the lateral side. The plantar head does not tilt medially or laterally. The lateral dorsal head is placed parallel to the proximal articular surface's axis. The pygmaios is present but rather small and not prominent.

Ravin de la Pluie sample (MN10)

Studied material (LGPUT, *Palaeotragus* cf. *coelophrys*). A 'hornless' skull with a highly worn tooththrow (RPI-91B); part of right mandible, with highly worn M_1 - M_3 (RPI-104F); isolated upper molar (M^1 or M^2) (RPI-315n). The taxon is represented by at least an adult and an old individual. Dental measurements are provided in Appendix 2: Table 5.

Cranium (RPI-91B). The skull is strongly deformed (Fig. 5), and hence a thorough description is prevented. Geraads (1978) provided the basic recognizable cranial and dental features; we repeat here the most important ones and add a few more. The cranium lacks ossicones, and has a flattened frontal region (Fig. 5B) suggesting a female individual. The parietal region is highly lateromedially compressed. The postorbital region is elongated. The mesial border of the orbit reaches almost at the level of the center of the M^3 (Fig. 5A). The length from the mesial margin of the orbit to the mesial root of P^2 is 153.8 mm. The height of the orbit is 58.9 mm, while its horizontal (caudo-rostral) diameter is 59.2 mm. The width of the region of the frontal bone behind the orbits is 95.4 mm.

Upper dentition. The tooththrow of the cranium RPI-91B is strongly worn (Fig. 5C). The premolars' width is similar to that of the molars, especially the width of the P^4 . The P^2 and P^3 are lingually rounded. Labially, the parastyle, paracone rib, and metastyle are all well-developed, with the paracone rib set mesially and close to the parastyle. The premolar fossettes demonstrate a slight hypoconal fold. P^4 is labially

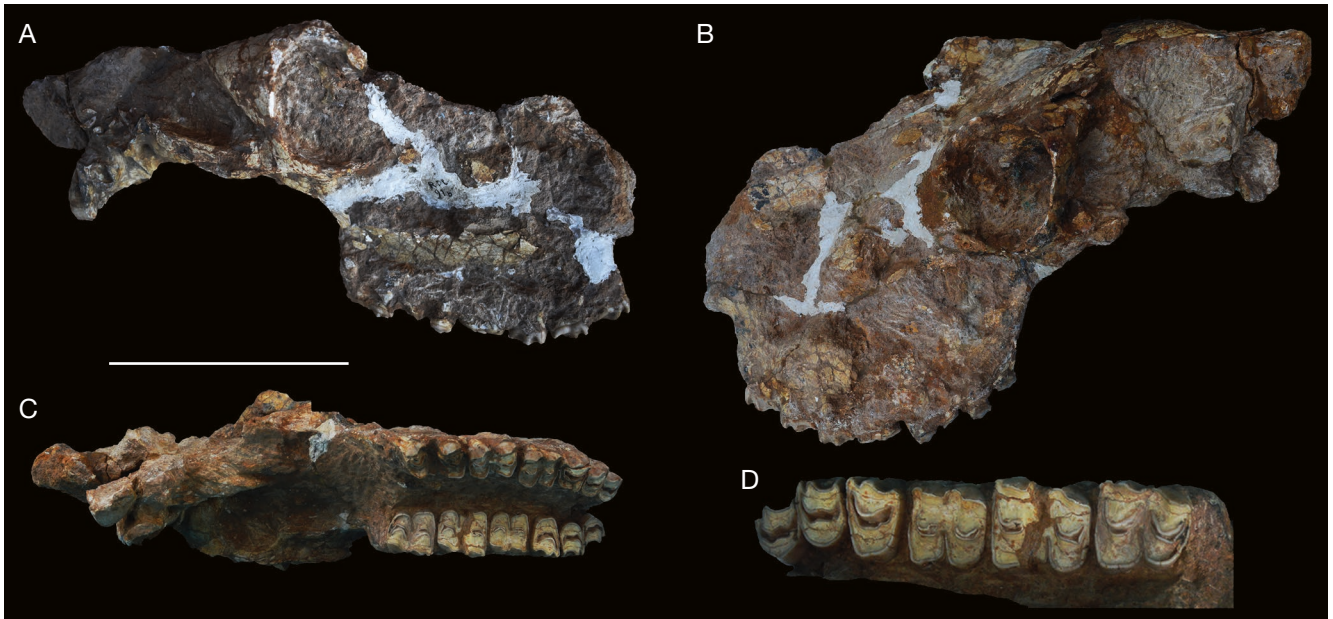


FIG. 5. — Partial cranium, LGPUT RPI 91-B of *Palaeotragus* cf. *coelophrys* from Ravin de la Pluie, Axios Valley, Greece (late Vallesian): **A**, right-lateral; **B**, left-lateral; **C**, ventral views. Scale bar: 15 cm.

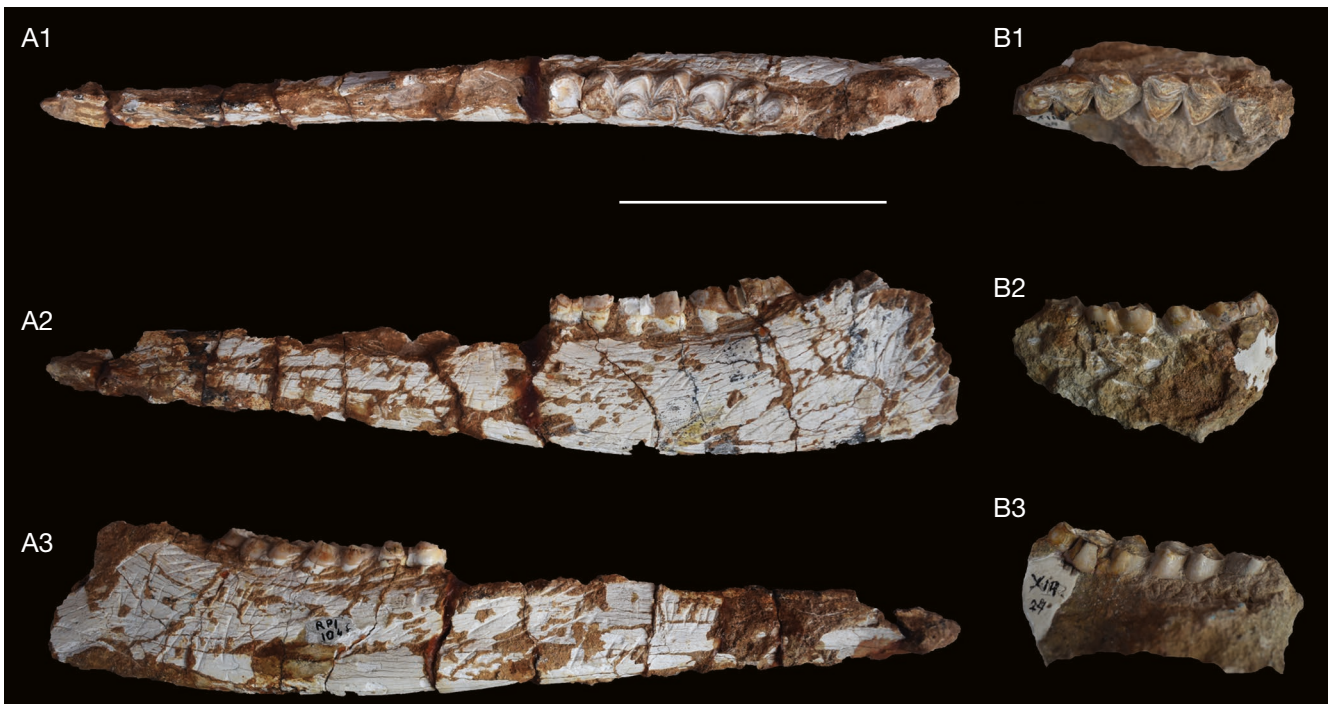


FIG. 6. — **A**, Part of mandible, LGPUT RPI-104F with M_1 - M_3 of *Palaeotragus* cf. *coelophrys* from Ravin de la Pluie, Axios Valley, Greece (late Vallesian) in: **1**, occlusal; **2**, lingual; **3**, labial views; **B**, part of mandible, LGPUT XIR-24 with M_2 - M_3 of *Palaeotragus* sp. from Xirochori, Axios Valley, Greece (late Vallesian) in: **1**, occlusal; **2**, lingual; **3**, labial views. Scale bar: 10 cm.

more flattened and more pointed lingually than the P^2 and P^3 ; a weak, shallow lingual furrow unequally divides the lingual cone. P^4 also has a strong parastyle, paracone rib and metastyle. The upper molars have a thin cement cover labially. The protocone widens significantly at the basal part of the crown; it is slightly constricted lingually on M^1 and M^2 , and more angular on M^3 . The labial ribs and styles are

extremely worn but apart from the parastyle they do not seem well-developed.

Lower dentition. The mandible RPI-104F preserves only the molars (Fig. 6A; Table 6). The mandibular height is 48.9 mm at the level of M_1 - M_2 and 53.6 mm at level of the distal lobe of the M_3 . The lingual ribs seem very weak though the dentition

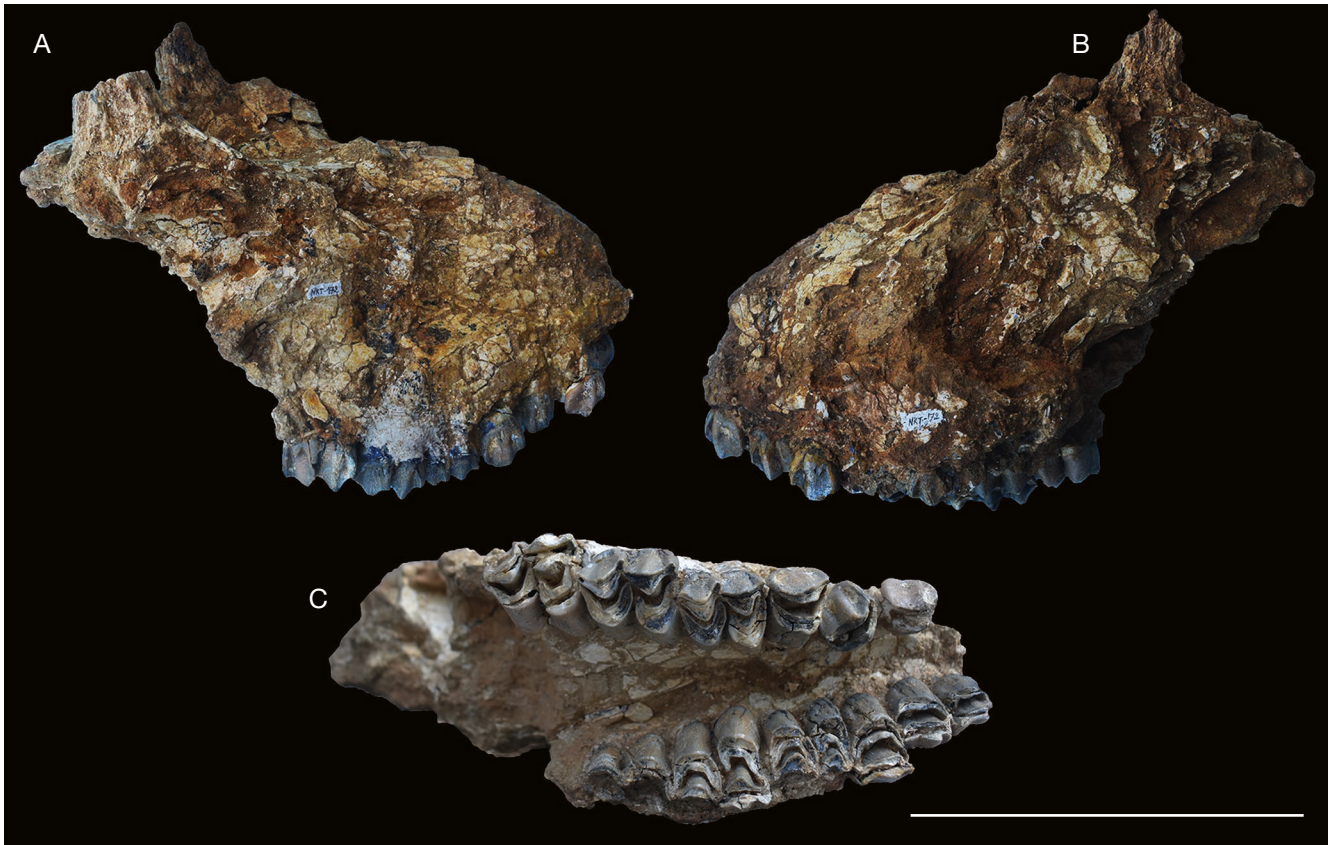


FIG. 7. — Partial cranium, LGPUT NKT-172 of *Palaeotragus* aff. *berislavicus* from Nikiti-1, Chalkidiki peninsula, Greece (latest Vallesian) in: **A**, right-lateral; **B**, left-lateral; **C**, palatal views. Scale bar: 15 cm.

is in a very advanced wear stage. Labially, the lobes of the M_1 , and but less those of M_2 are pointed and centrally oriented, while those of M_3 are directed more distally. The distal (3rd) lobe of M_3 is quite large and disto-labially oriented, separated lingually from the rest of the tooth by a slight groove. There is no evidence of cingulum or cement on the molars.

Xirochori sample (MN10)

Studied material (LGPUT, *Palaeotragus* sp.). Right mandible fragment with M_2 – M_3 (XIR-24). The taxon is represented by a single adult individual. Measurements are provided in Appendix 2: Table 7.

The lingual ribs and stylids are weak on the preserved M_2 and M_3 of XIR-24 (Fig. 6B) but the metaconid and the metastylid are quite prominent. The second and third lobes of the M_3 are separated by a very shallow groove. Labially, the protoconid and the hypoconid direct distally and they are both fairly more pointed in the M_2 than in the M_3 . A weak entoconid is observed on the M_2 . A strong ectoconid is observed between the first and the second lobe of M_3 , while there is a weaker one between the second and the third lobe. Neither cingulum nor cement are observed.

Nikiti-1 sample (end MN10)

Studied material (LGPUT, *Palaeotragus* aff. *berislavicus*). A partial cranium preserving both tooththrows and the very

proximal part of the ossicones (NKT-172); a left humerus distal articulation (NKT-161); three left radii (NKT-156, NKT-159, NKT-169); a right radius (NKT-155); a proximal part of radius (NKT-167); two right metacarpals (NKT-137, NKT-141); a left metacarpal (NKT-131); two proximal parts of left metacarpals (NKT-26, NKT-67); a right tibia (NKT-271); two distal parts of tibia (NKT-150, NKT-154); two left astragali (NKT-163, NKT-266); a right astragalus (NKT-267); two right calcanei (NKT-153, NKT-268); four left metatarsals (NKT-133, NKT-136, NKT-139, NKT-160); two right metatarsals (NKT-138, NKT-144); a distal part of a right metatarsal (NKT-151); three proximal parts of right metatarsals (NKT-132, NKT-140, NKT-168). The taxon is represented by at least four adult and a possibly young individual. Measurements are provided in Appendix 2: Tables 8–14.

Cranium (NKT-172). The cranium is badly crashed, moderately deformed, and laterally compressed (Fig. 7). It preserves both tooththrows but in the left one the molars are very badly preserved (Fig. 7C). The mesial margin of the orbit reaches the level of M^3 . The length from the mesial edge of the orbit to the mesial root of P^2 is 154 mm. A pair of supraorbital ossicones is present, both broken near the base (Fig. 7A). It can be assumed that the ossicones are placed relatively medially compared to the dorso-lateral orbital margins, although the



FIG. 8. — *Palaeotragus* aff. *berislavicus* from Nikiti-1, Chalkidiki peninsula, Greece (latest Vallesian): **A**, radius, LGPUT NKT-156; **B**, radius, LGPUT NKT-155; **C**, metacarpal III-IV, LGPUT NKT-137; **D**, metacarpal III-IV, LGPUT NKT-131. Views: 1, proximal; 2, cranial (dorsal); 3, distal; 4, caudal (palmar) views. Scale bar: 20 cm.

skull is very deformed. The basal anteroposterior diameter of the ossicones is 43.4 mm, while the transverse diameter is *c.* 32 mm. Their cross section is oval shaped with main axis trending anteroposteriorly.

Upper dentition. The labial side of the left molars of NKT-172 is damaged (Fig. 7C; Table 8). The toothrows are not very worn. The P² and P³ have a circular occlusal outline. Labially, the parastyle and the paracone ribs are well-developed, whereas the metastyle is weak. The fossettes of the P² and the P³ are wide and U-shaped. A hypoconal fold can be seen in the distal part of the P³'s fossette. Compared to the P² and P³, the P⁴ has a more pronounced metastyle, and a less convex lingual wall giving to it a more squared occlusal outline. The central fossette of the P⁴ is wide and simple. On the disto-lingual part of the P⁴ there is a style that reaches almost to the half of the crown height.

The basic morphology of the molars is similar to those of other giraffids (Fig. 7C). The M² and M³ are slightly worn and have well-developed parastyle, paracone rib, mesostyle and metastyle. The labial rib of the metacone is almost flat. A weak labial cingulum is present on M¹ and M². On the M³ a labial style is attached on the basis of the metacone. Lingually, the protocone and the hypocone are almost equally prominent and of similar shape on the M¹ and M². In the M³ the protocone protrudes lingually more than the hypocone. The distal protocone and mesial hypocone flanges converge in the middle of the tooth. The

distal hypocone flange tends to connect with the metastyle; they are already connected in the less worn M³. The size of the mesial fossette is small. Mesially, the protocone flange is connected to the parastyle. Both fossettes are U-shaped. There is a very weak hypoconal fold in the M².

Postcranials. The preserved distal humerus (NKT-161) is crashed and deformed preventing reliable morphological observations (breadth of distal epiphysis: 85.4 mm; width of distal epiphysis: 40.94 mm). Its size is intermediate between those of *P. rouenii* and *P. coelophrys*. It has the same width as a *P. microdon* specimen described by Bohlin (1926). On the NKT specimen, the olecranon fossa is deep, wide and U-shaped.

The radii are all elongated (Fig. 8A, B) and moderately slender. Two of the NKT radii are fairly curved (concave laterally; NKT-155, NKT-156; Fig. 8A, B; Table 9), while the rest three are straighter (NKT-159, NKT-167, NKT-169). The cross section is crescent-shaped in the proximal 4/5 of bone's length, with rounded cranial and straight caudal faces; it is trapezoidal shaped in the distal most 1/5 of radius length. The two epiphyseal areas are both much wider than the shaft, and they both expand medially and laterally. In the specimen NKT-159 the olecranon is partly preserved. It seems to greatly tilt latero-distally. Proximally, the medial, rectangular/sub-rectangular shaped articular surface is much larger than the quadrangular lateral one, separated by each other by a wide, shallow furrow (Fig. 8A1, B1). The medial tuberosity is not



FIG. 9. — *Palaeotragus* aff. *berislavicus* from Nikiti-1, Chalkidiki peninsula, Greece (latest Vallesian): tibia, LGPUT NKT-271 in: **A**, proximal; **B**, cranial; **C**, distal, **D**, caudal views. Scale bar: 20 cm.

developed. A tilted narrow crest divides the distal articular surface into two equal subregions, representing the articular surfaces for the scaphoideum and semilunare, respectively. They are both round and slightly concave. The semilunare surface is

interrupted by a convex protrusion distally. The groove for the extensor carpi radialis muscle at the cranial part of the distal epiphysis is wide, and very shallow to flattened, delimited by blunt ridges (Fig. 8A2, B2).

The best preserved metacarpals (Fig. 8C, D) vary slightly in length (Table 10), having however similar robusticity indices (8.38–8.75%). The lateral and medial epicondyles are asymmetrical. The lateral epicondyle has half the size of the medial one, and it is of square or rectangular shape. The medial epicondyle is rounded dorsally and has an overall shape of a half to $\frac{1}{4}$ of a circle. The fossa in-between the two epicondyles continues in the medial epicondyle. The medial and lateral epicondyles continue to the medial and lateral ridges respectively, which are of similar width and morphology. They are both rounded near the proximal end and they become slenderer and sharper in the shaft area. The central trough is very deep near the proximal end of the bone but becomes shallower and flatter towards the distal end. The trough's width is variable. However, it can be said that the longer the bone, the wider is the trough. The pyramidal rise is absent in most of the specimens apart of NKT-137 in which a slight protrusion could be attributed to the pyramidal rise; however, it is not prominent at all. The keels of the distal epicondyles are more prominent palmarly, and they also extend onto the distal end of the palmar side of the shaft.

Three tibia specimens are preserved (Table 11). Proximally, the angle of the sulcus muscularis is obtuse when compared to the bovid or cervid anatomy (Fig. 9A, B). The tuberosity is not well preserved but does not seem to be pronounced (Fig. 9). The lateral condyle is somewhat damaged laterally and cranially. In the better preserved distal tibial fragment NKT-154 the cross section of the distal shaft is sub-rectangular. The distal articular surface has a broad rectangular shape, and the articular surface for the medial malleolus is quite prominent (Fig. 9C). The cochlear crest, that separates the cochlear furrows, is moderately prominent forming a strong caudal and a weak cranial crest.

The size of the three preserved NKT astragali (Fig. 10; Table 12) is intermediate between the groups of small and large *Palaeotragus*. The NKT astragali are rectangular in shape, as the lateral and medial lengths are of almost equal size and not thickened medially. In dorsal view the lateral edge of the trochlea tilts slightly medially. The central fossa is large, deep and triangular. The medial groove is weak in NKT-266; it is damaged in the other two specimens. The lateral notch is faint in NKT-266, prominent in NKT 267 (Fig. 10A) and intermediate in NKT-163 (Fig. 10B). The median depression is wide and deep in NKT-163 and NKT-266 but shallower in NKT-267. Ventrally, the medial ridge tilts medially in the specimens NKT-266 and NKT-267 (Fig. 10A), while it is almost vertical in NKT-163 (Fig. 10B). The intertrochlear notch is deep and narrow in NKT-163, but it is wide and shallow in NKT-266 and NKT-267 (Fig. 10). The proximal triangular fossa is prominent and the interarticular groove absent in all the specimens. The medial scala is absent in NKT-267, weak in NKT-266 (Fig. 10A) and slightly more prominent in NKT-163 (Fig. 10B). The distal intracephalic fossa is absent in NKT-266

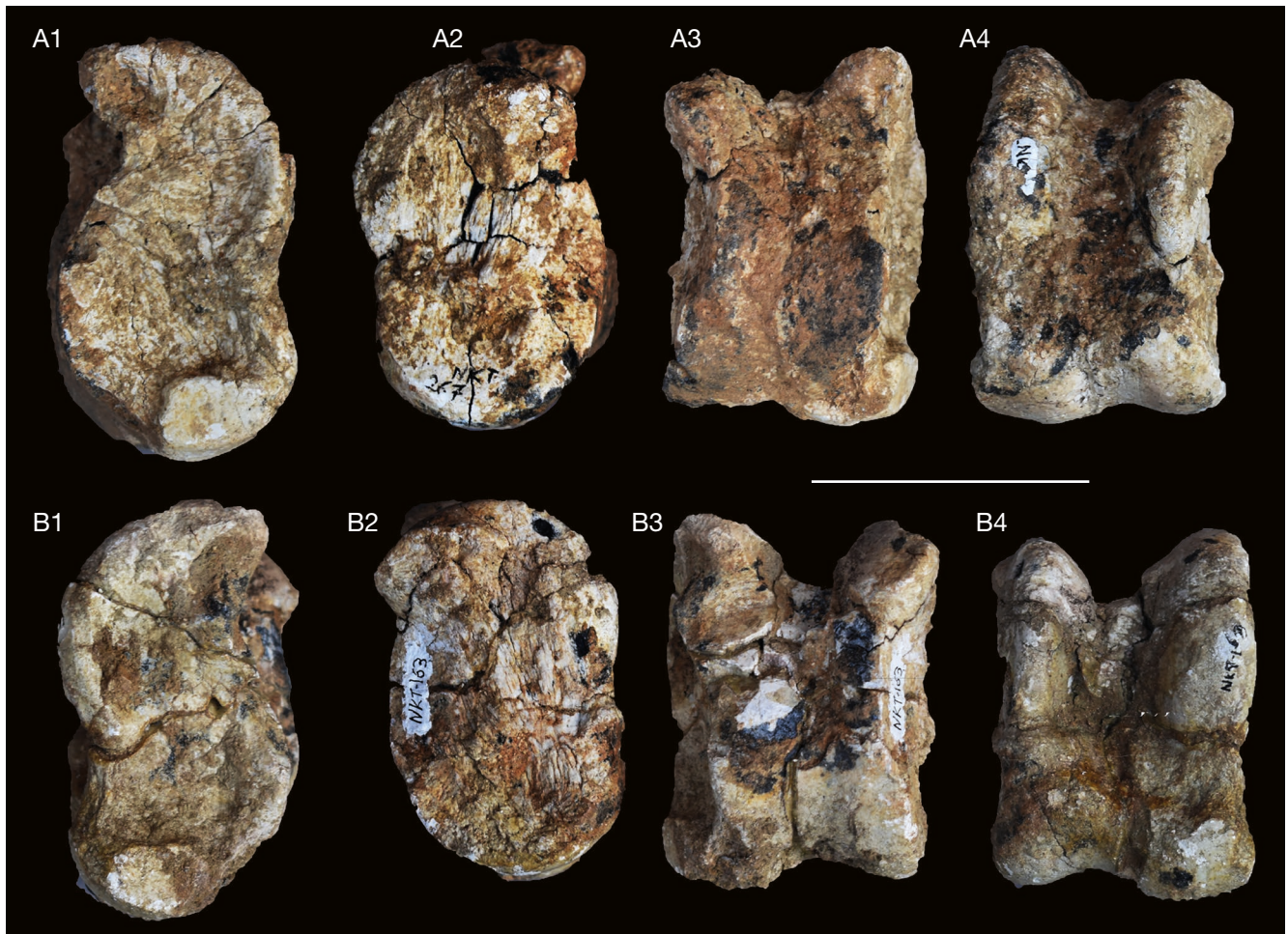


FIG. 10. — *Palaeotragus* aff. *berislavicus* from Nikiti-1, Chalkidiki peninsula, Greece (latest Vallesian): **A**, astragalus, LGPUT NKT-267; **B**, astragalus, LGPUT NKT-163. Views: 1, lateral; 2, medial; 3, posterior; 4, anterior. Scale bar: 5 cm.

and NKT-267; in NKT-163 it probably exists, although the preservation status of the specimen does not allow to be decisive (Fig. 10). The lateral and medial crests are equally thick, with the lateral pointing posteriorly, and the medial slightly tilting medially. The lateral side of the astragali is somewhat concave.

Two right calcanei were collected. The calcaneus NKT-153 (Fig. 11; Table 13) is larger, with a more robust corpus than NKT-268. However, the NKT-268 head seems to be stronger relatively to the corpus, than the NKT-153 one. Both the dorsal and plantar crests are parallel to the bone axis. The calcaneal tuberosity is prominent, though weathered in both specimens. In NKT-268 a medial crest separates the calcaneal tuberosity from the rest of the calcaneal corpus. The sustentaculum tali is somewhat damaged in NKT-153 (Fig. 10B, C), while it is robust in NKT-268. The proximo-plantar articular surface for the astragalus consists of two concave surfaces, with the plantar one being almost double in size than the dorsal surface. Medially, the articular surface for the astragalus is fairly deep and concave. The articular surface for the scaphocuboidum is located plantarly and is slightly damaged in both specimens. Dorsally, there is a well-developed, articular facet for the malleolus.

The metatarsals (Fig. 12; Table 14) demonstrate a variation in both the total length and the robusticity indices (7.3–12%), also partly due to postmortem deformation (e.g., NKT-133). The lateral epicondyle is smaller and subdivided in two regions, the dorsal and the plantar heads (Fig. 12A). In contrast, in the medial proximal epicondyle this separation is not evident. The medial epicondyle has a trapezoid shape (Fig. 12A). The lateral dorsal head is more circular. The shape of the lateral plantar head is intermediate in shape. The pygmaios is not preserved (Fig. 12A, B). The central trough varies in the studied specimens. However, it is significantly shallower and it disappears from the middle towards the distal end of the bone (Fig. 12B). The width of the trough also varies, and it seems to follow the total bone width. The proximo-plantar fossa is present, weak and communicates with the central trough in the specimen NKT-139 whereas the bad conservation status of the other specimens does not allow to identify it. In contrast to the metacarpals, a dorsal trough is evident; it is deeper at the proximal end of the bone, becomes shallower downwards and disappears at the distal end (Fig. 12D). The distal epiphysis of the metatarsals is similar to that of the metacarpals.

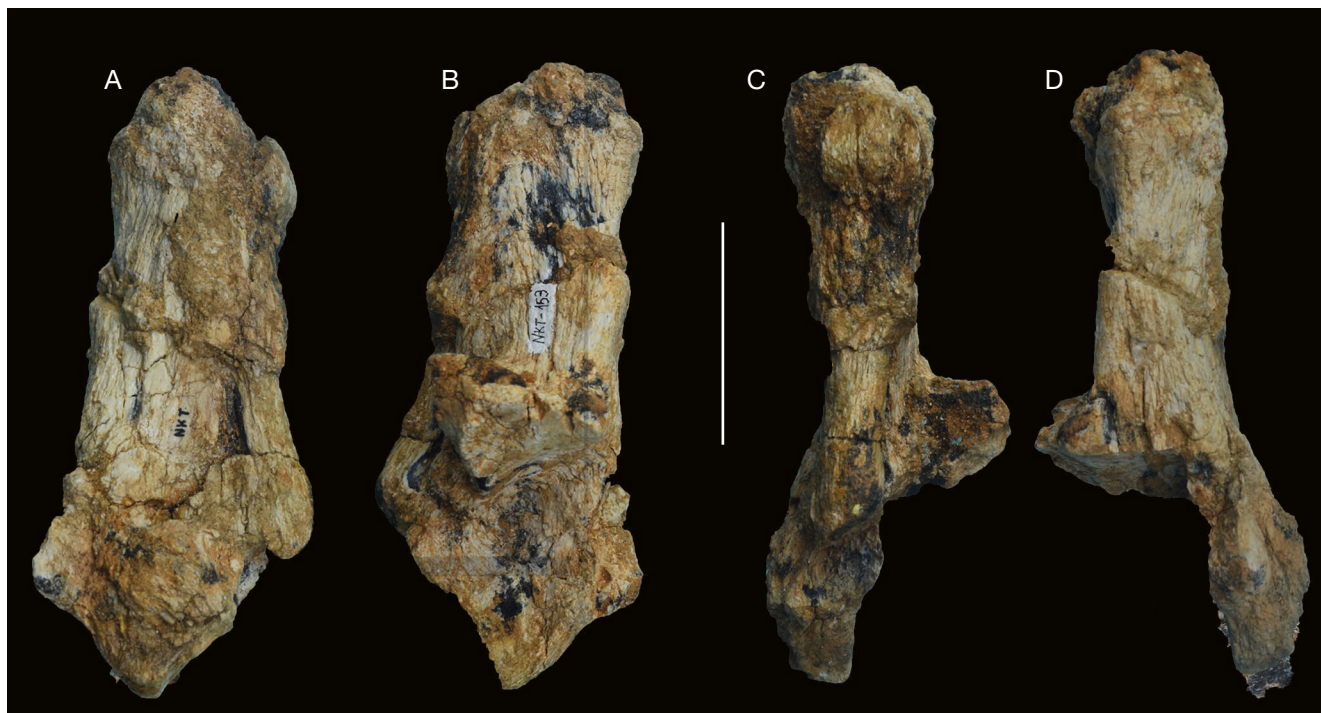


FIG. 11. — *Palaeotragus* aff. *berislavicus* from Nikiti-1, Chalkidiki peninsula, Greece (latest Vallesian). Calcaneus, LGPUT NKT-153 in: **A**, lateral; **B**, medial; **C**, dorsal; **D**, plantar views. Scale bar: 5 cm.



FIG. 12. — *Palaeotragus* aff. *berislavicus* from Nikiti-1, Chalkidiki peninsula, Greece (latest Vallesian). Metatarsal, LGPUT NKT-136 in: **A**, proximal; **B**, plantar; **C**, distal; **D**, dorsal views. Scale bar: 20 cm.

COMPARISON

THE TAXONOMY OF LATE VALLESIAN EURASIAN PALAEOTRAGUS

Assessing the relationships among the Late Miocene *Palaeotragus* taxa is a difficult task. A variety of species were described in the late 19th and early 20th centuries, in a time when communication among different scholars was quite difficult. Since then, just a few comprehensive systematic revisions have been accomplished (e.g., Godina 1979; Geraads 1986), even though the genus is the most common giraffid in the Late Miocene of Eurasia. The problem is further exacerbated by the wide geographical distribution of the relevant samples and their usually fragmentary nature.

Traditionally, *Palaeotragus* species are distinguished by means of size in two groups (Geraads 1974, 1986; Iliopoulos 2003; Kostopoulos & Saraç 2005; Kostopoulos 2009). The first group usually includes the small-sized taxa *Palaeotragus rouenii*, *P. microdon* and sometimes *P. pavlowae*. The term “*P. rouenii* group” is generally used for this ensemble of taxa, characterized by their small size and their long, slender metapodials. The second group incorporates larger species such as *Palaeotragus coelophrys*, *P. quadricornis*, *P. expectans*, *P. borissiakii*, *P. hoffstetteri*, *P. moldavicus*, *P. asiaticus* and *P. berislavicus*. The name “*P. coelophrys* group” is usually applied for larger-sized members, which are also characterized by the shorter and more robust metapodials than those of the “*P. rouenii* group”. Earlier works by Geraads (1974, 1986), do not detect any

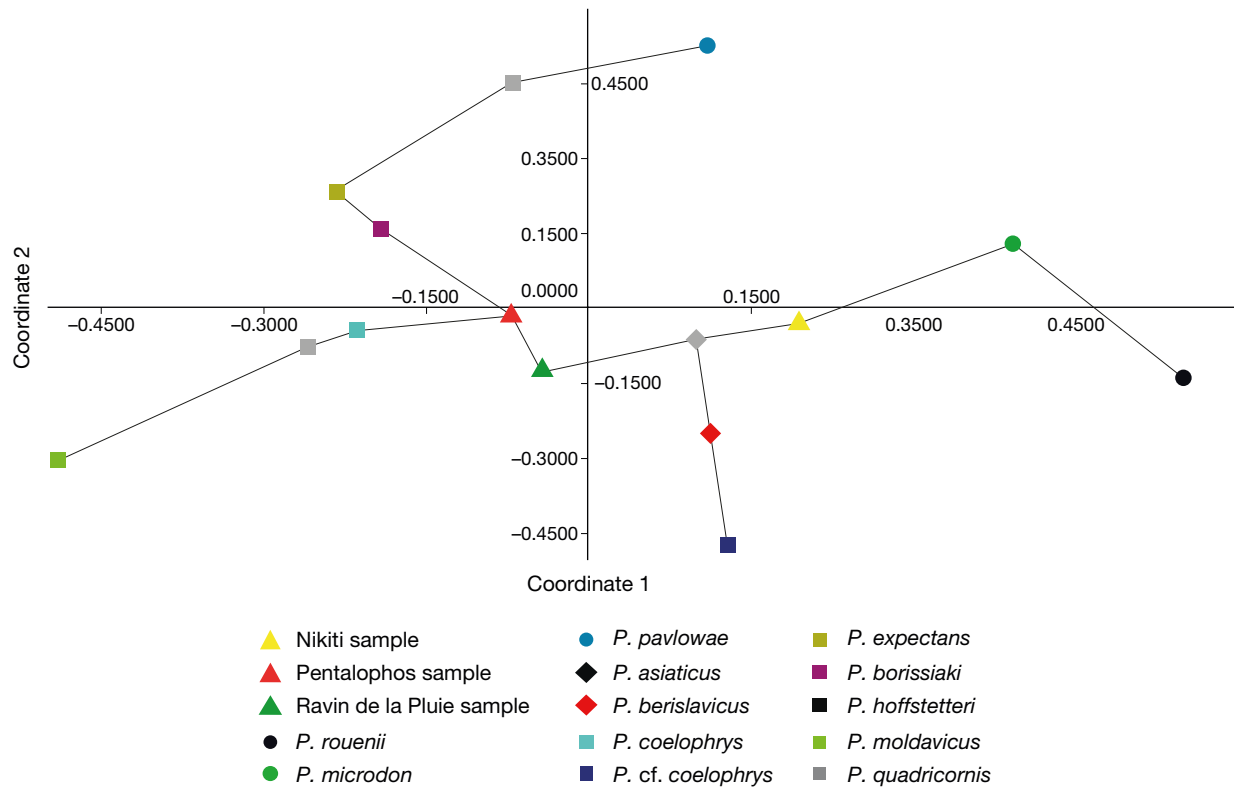


FIG. 13. — Principal Coordinates Analysis (plane of coordinates 1 and 2, representing 39,5% and 15,3% of initial data variance) comparing 27 biometrical variables and 4 selected features of premolar morphology of the Late Miocene Eurasian palaeotragines, as well as the Vallesian Greek *Palaeotragus* Gaudry, 1861 samples studied here.

significant or consistent morphometric difference among members of the “*P. coelophrys* group”, and all these taxa are commonly treated as possible synonyms under *P. coelophrys*, although the same author (Geraads *et al.* 2013) recently recognized the need for a deeper revision.

Godina (1979) on the other hand, followed a quite different approach, recognizing all species known at that time (except *P. quadricornis*) and grouping them into three distinct subgenera: *Palaeotragus* (*Palaeotragus*), with *P. rouenii* and *P. pavlowae*; *Palaeotragus* (*Yuorlovina*), with *P. coelophrys*, *P. microdon*, *P. asiaticus* and *P. hoffstetteri*; and *Palaeotragus* (*Achtiaria*), with *P. expectans*, *P. borissiakii*, *P. moldavicus* and *P. berislavicus*. According to Godina (1979) representatives of *Achtiaria* differ from those of *Yuorlovina* by a shorter cranial face, a shorter diastema compared to the tooththrow length and a non-continuous lingual wall on the P_3 . Members of *Palaeotragus* are smaller in size, the front part of the skull is elongated, and the diastema length exceeds that of the tooththrow; the lingual wall of P_3 could be either continuous or not.

Godina’s (1979) taxonomic point of view has not been widely accepted and is proving quite fragile in several respects. First, the type material of most species in question does not include the anatomical elements on which presumed taxonomic distinctions are based. Additionally, the validity of most diagnostic features is challenged by the scarcity of the material. For example, while Godina (1979) repeatedly used mandibular characters to assess similarity, for some of

the proposed species only a single mandibular specimen was known at that time per species, while for some others there were not even complete mandibular specimens preserved. The morphology of the ossicones used to distinguish some of the bulky palaeotragines by Godina, was not known for all *Palaeotragus* species (e.g., *P. coelophrys* from Maragheh), whereas sexual dimorphism proved later to have a strong effect on the ossicone size (e.g., Geraads 1978; Kostopoulos 2009).

To test previous taxonomic assumptions, we perform here an analysis of 27 available biometrical variables (absolute and relative) and of 4 selected morphological characters of premolar morphology (see Appendix 1). In order to avoid discrepancies per species introduced by later assignments our initial analysis was based exclusively on the material provided from the type localities. Our basic results are summarized and better illustrated by a PCO analysis (Fig. 13) on which the Greek material studied here is also extrapolated (see Appendices 3, 4 for additional information). All multivariate treatments of both PCA and PCO suggest the following clusters:

P. microdon and *P. rouenii* are always separated from the rest of taxa, as well as between them, keeping a marginal position on the diagrams (Fig. 13; Appendix 3: Figs S1, S2).

P. berislavicus, *P. asiaticus* and the Nikiti-1 taxon are grouped more closely together than with other compared taxa. PCO analyses constantly indicate *P. cf. coelophrys* from China as closer to this set of taxa, though in PCA the Chinese species appears isolated.

P. coelophrys from Maragheh, *P. borissaki*, and *P. expectans*, along with PNT and RPI samples are grouped closer together than to other taxa. *P. quadricornis* from Samos and the Sinap *P. hoffstetteri* always appear closer to members of this group than to the previous one. The same may apply for *P. moldavicus*, though this taxon appears in several analyses more distant.

P. pavlowae is the most debatable taxon among the compared ones but also one of the less known. It appears close to *P. asiaticus*-*P. berislavicus*-NKT ensemble in PCA, but it is quite remote in PCO.

The taxonomy of *P. rouenii* and *P. microdon* is sufficiently solid. Both taxa represent the smaller Late Miocene representatives of the genus with the Chinese taxon, *P. microdon* being distinguished from the European one, *P. rouenii* by its longer tooththrow with shorter premolars compared to molars, more advanced P₃, shorter and more robust radius and metacarpal and shorter but equally slender metatarsal (Appendices 3, 4; see also Bohlin 1926; Geraads 1974; Hamilton 1978; Kostopoulos & Saraç 2005). Among several populations of *P. rouenii* described so far, three are worthy of further mention. First, the *P. rouenii* sample from Thermopigi (Xafis *et al.* 2019) demonstrates extremely elongated postcrania, especially the radius (SIT-939), and metatarsal (SIT-307; Xafis *et al.* 2019). A similarly long radius has been described by Kostopoulos & Koufos (2006) from Perivolaki (LGPU PER-1180), whereas two very long metatarsals (but shorter than those from Thermopigi) have been described from Kryopigi (LGPU KRY-7937, KRY-7938; Lazaridis 2015). Although the extreme lengthening of these specimens cannot be easily interpreted, it is certainly indicative of the strong dolichopodality developed in some *P. rouenii* populations, allowing an overall re-consideration of its intraspecific metrical variability.

P. pavlowae was erected by Godina (1979) based on material from Grebeniki (Ukraine) originally described by Pavlow (1913) as *Camelopardalis parva*. Pavlow (1913) ascribed to this taxon only a palate (RSGU 1637) and referred to the same genus a M₂-M₃ (no catalogue number indicated). Godina (1979) uses RSGU 1637 as the holotype of *P. pavlowae* and also refers to this species an upper and a lower juvenile dentition (OSU 2376 and OSU 2375, respectively), a partial ulna, and an astragalus (OSU 2749, OSU 2746, respectively) stored in the Odessa Museum, a first phalanx (SIZK 25-118) stored in Kiev, all from Grebeniki as well as an upper P₂-M₃ (RSGU 1639) from Blagodarnenskaya, Stavropol. No lower dentition has been ascribed to *P. pavlowae* from its type locality. Godina's (1979: 44) poor species diagnosis indicates: "Length P₂-M₃ – 127 mm. Limbs of intermediate length and massiveness" (translated from Russian). The length of the holotype upper tooththrow from Grebeniki is indeed larger than that of *P. rouenii* (Appendices 3, 4) and within the metrical and proportional range of *P. microdon*. On the other hand, the postcrania ascribed by Godina (1979) to this taxon appear larger and more robust than those referred by Bohlin (1926) to *P. microdon* and are closer to those from Pavlodar [a metatarsal (PIN 2432/7005) and three astragali (PIN 2346-130, PIN 2346-151, PIN 2413-6939)] referred to as *P. asiaticus* (Appendices 3, 4). The deciduous upper dentition from Gre-

beniki assigned to the same species is comparable in size to that of *P. rouenii* (dP₂-dP₄ average length= 57 mm), but the milk teeth show a well-developed lingual cingulum; the dP₂ is more symmetric mesio-buccally; the mesial lobe of dP₃ is rather triangular than trapezoidal, the angle formed by the mesial and distal buccal flanges of the paracone is obtuse, and the mesio-buccal stylid is bifurcated; a postprotocrista and a neocrista are still detectable on the dP₃; as the dP₄, M¹ and M² show a strong hypoconal spur (pers. obs.; unfortunately, we couldn't locate OSU 2375 in the Odessa collections). By these features OSU 2376 appears closer to the dental morphology of AMNH 22807 from Samos and AMNH 26362 from Shansi (see Kostopoulos 2009:308), as well as to *P. expectans* (see Borissiak 1914: pls 1, 2). Inadequate evidence does not allow definitive conclusions about the Grebeniki taxon; we consider it more likely to be a chimeric species resulting from the mixing *P. rouenii* material with that of another palaeotrage.

Two morphometric clusters are recognized here within the group of large Late Miocene palaeotrages. *P. coelophrys*, *P. expectans*, *P. borissaki*, and *P. quadricornis* share a more primitive dP₃ and P₃ structure (the former unknown in *P. quadricornis*), larger size and more robust proportions compared to the rest of the examined taxa. The dP₃/P₃ morphology and postcranial proportions of *P. hoffstetteri* are very similar to this group of taxa, as deduced from several postcrania in MNHN, Paris and NHM Izmir (most of them belonging to the original material described by Ozansoy in 1965) and some dental remains from Loc 49 of Sinap (referred to as *P. coelophrys* by Gentry 2003). As others before us (e.g., Churcher 1970; Geraads 1974, 1978) we think that there are no sufficient ground to discriminate within this set of taxa, for which we apply by priority the name *P. coelophrys*. By its absolute dimensions and proportions *P. moldavicus* matches pretty well *P. expectans*-*P. borissaki* but differs in apparently fully molarized P₃ of the paratype mandible PIN 649/26 (Godina 1979: fig. 8). Moreover, *P. moldavicus* postcrania appear to be larger than those of other members of this group. However, the known material is so scarce, that any attempt to distinguish it from other large palaeotrages would be premature at the moment.

The second group of large Late Miocene palaeotrages recognized in our analysis includes *P. berislavicus* and *P. asiaticus*. They share a fully molarized P₃, dental proportions and limb lengths comparable to those of *P. rouenii*, but the dentition is significantly larger (c. 15%) and the postcrania stouter, though not as much as in *P. coelophrys* (Appendices 3, 4). Although suggestions about the Vallesian *P. berislavicus* morphology and proportions are relatively safe as they are based on material from a single locality, Berislav (Ukraine), the younger *P. asiaticus* is a more doubtful species. The Ortok (Kyrgyzstan) type material (Godina 1979) includes a cranium with tooththrow and a P₂-P₄ both intermediate in size and premolar/molar proportions between *P. rouenii*/*P. microdon* and *P. coelophrys* (Appendices 3, 4); a few isolated upper and lower teeth with some approaching better *P. coelophrys*, while others being closer to the upper range of *P. rouenii*/*P. microdon*; a radius similar in size and proportions to *P. coelophrys* (Appendix 3: Fig. S5); two

intermediate tibiae (Appendix 3: Fig. S7); and an astragalus comparable in size to *P. rouenii* (Appendix 3: Fig. S8). No metapodials are known from the type locality. Material from Pavlodar (Kazakhstan) ascribed by Godina (1975, 1979) to the same species includes some isolated lower teeth intermediate in size but closer to *P. coelophrys*; a metatarsal as long as that of *P. rouenii* but more robust (Appendix 3: Fig. S9); and three astragali similar to those of *P. coelophrys* (Appendix 3: Fig. S8). An intermediate in size lower dentition (length $P_2-M_3=141$ mm) and a metatarsal as long as in *P. rouenii* (453 mm) from Kalmakpaj (Kazakhstan), as well as a partial metatarsal from Pristashkent district (Uzbekistan) were also ascribed to *P. asiaticus* by Godina (1979). Although we cannot fully exclude the possibility that *P. asiaticus* may represent a distinct taxon, similar in several respects to *P. berislavicus*, it is also highly probable that the species concept is based on a mixture of material of two different species.

By its tooththrow size *P. cf. coelophrys* from China matches pretty well the *P. berislavicus*-*P. asiaticus* ensemble. Its P_3 is also molarized (Bohlin 1926: pl. 3), but the hypoconulid is communicating with the mesial lobe of the tooth. The only postcranial evidence available for this taxon is an astragalus (Bohlin 1926 – Appendices 3, 4), whose proportions do not differ from those of *P. rouenii*, or *P. berislavicus*-*P. asiaticus*. The Chinese taxon could actually belong to the same species represented by the roughly contemporaneous and geographically close Ortok-Pavlodar samples, a hypothesis already proposed by Godina (1975, 1979). For the time being and until new material is available we prefer to exclude the Asian taxa from our overall concept and we consider *P. berislavicus* as a valid species based on the Berislav material only.

Conclusion

To sum up, our overview of the Late Miocene palaeotragines from Eurasia allows recognizing four valid species

P. rouenii Gaudry, 1861; type species characterized by small size, fairly molarized P_3/dP_3 , lower premolar to molar length ratio between 57% and 62%, and strongly dolichopodial limbs. Female ossicones absent or very slim.

P. microdon (Koken, 1885); characterized by small size, longer tooththrow with shorter premolars (lower premolar to molar length ratio between 63% and 73%), more advanced P_3 , shorter and more robust radius and metacarpal and shorter metatarsal than the type species. Females bear slender ossicones.

P. berislavicus Korotkevitch, 1957 (possible synonym of *P. asiaticus*); characterized by intermediate size, fully molarized P_3 , and similarly long but stouter limbs than the type species.

P. coelophrys (Rodler & Weithofer, 1890) (synonym of *P. expectans*, *P. borissiakii*, *P. hoffstetteri*, *P. quadricornis*, and possible synonym of *P. moldavicus*); characterized by larger size, less molarized P_3/dP_3 , and more robust limbs than the type and rest of species. Known females lack ossicones.

THE VALLESIAN GREEK LARGE PALAEOTRAGES

Pentalophos sample

The dental remains from *Pentalophos* indicate a taxon significantly larger than *P. rouenii*, and *P. microdon*. In most abso-

lute size and proportions, they match those of *P. coelophrys* (Fig. 13; Appendix 3). Comparing the PNT upper teeth with those of the type cranium of *P. coelophrys* from Maragheh, premolars have in both cases a slight hypoconal fold but the P^2 from Maragheh has a somewhat prominent lingual rib in the hypocone area that it is absent from the P^2 of PNT. The P^3 of Maragheh is similar to that of PNT, except that it is squarer and bears a lingual groove. In the only available P^4 from PNT, the paracone is more centrally placed, the P^4 fossette is somewhat better developed mesially and lingually, and the occlusal outline more triangularly shaped than in Maragheh. The premolars of the Vallesian *Palaeotragus* from Sinap illustrated by Ozansoy (1965: pl. X, fig. 3) resemble a lot that of PNT, although they are significantly more square (especially the P^4) and with enlarged fossettes. The premolars of another Sinap upper tooththrow housed in MNHN Paris (MNHN.F.TRQ no catalogue number) show a much weaker hypoconal spur and a more robust paracone, while the parastyle is considerably weaker in P^2 and P^3 compared to the PNT premolars. Finally, the P^4 is squared in the MNHN.F-Sinap specimen, in contrast to the triangular P^4 from PNT. The Vallesian *P. coelophrys* from Eldari (Alexeev 1930: pl. I, figs 1-4) bears an unusually triangular P^3 (specimen PIN 1408/381), while the P^3 of the specimen PIN 1408/202 is identical to the premolars of PNT, probably bearing a prominent distolabial stylid, similar to that of the P^2 of PNT-113. The P^4 has a circular occlusal outline in PIN 1408/381 and squared in PIN 1408/202, in contrast to the triangular in PNT-113. The lower deciduous dentition from *Pentalophos* (LGPNT PNT 121F) is almost identical to that of the Vallesian *P. coelophrys* from Sebastopol (Borissiak 1914: pl. II, figs 1-5), both having a large rhomboid mesial lobe and strong ectostylids on dP_4 and a less advanced dP_3 with the metaconid being attached to the protoconid-entoconid junction and turned strongly forwards without closing the anterior valley.

The PNT metatarsals seem to approach better *P. coelophrys*, as they are considerably robust proximally (Appendix 3: Fig. S9). The morphology of the proximal articular surface of PNT differs from that of *P. rouenii* in the bone protrusion instead of light groove separating medially the plantar and dorsal heads, the non-tilt plantar head, the more robust lateral dorsal head, and the less prominent pygmaios.

Multivariate analysis (Fig. 13; Appendix 3) consistently suggests PNT taxon as grouped with the largest palaeotragines. Hence, according to both morphological and metrical evidence we attribute the *Pentalophos* taxon to *P. coelophrys*.

Ravin de la Pluie sample

Geraads (1978) originally studied the RPI-91B cranium and referred it to as *Palaeotragus cf. coelophrys*. A new morphological comparison of the worn tooththrows, indicates that the RPI palaeotrage has more rectangularly shaped and larger premolars compared to molars than the *P. coelophrys* type from Maragheh. The P^3 and P^4 of RPI are very similar in shape with the P^3 and P^4 from the Vallesian specimen PIN 1408/202 of *P. coelophrys* of Eldari (Borissiak 1914: pl. I, fig. 4). They are also very similar with the teeth of the Vallesian *P. coe-*

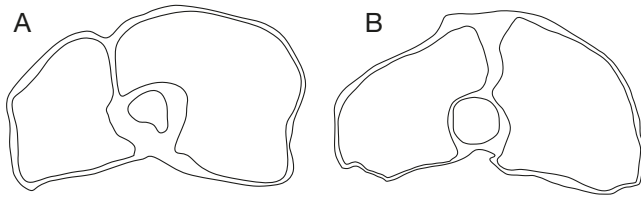


FIG. 14. — Morphological comparison of *Palaeotragus* Gaudry, 1861 proximal left metacarpal epiphyses: **A**, *P. rouenii* Gaudry, 1861 from Nikiti-2; **B**, *P. aff. berislavicus* from Nikiti-1.

lophrys from Sinap (MNHN.F.TRQ no catalogue number), although more square shaped. Most likely due to advanced wear, the paracone of the P³ and P⁴ of the RPI skull is much less prominent than the paracone of the respective teeth of the MNHN.F specimen. The premolars of the *P. coelophrys* specimen from Sinap, provided by Ozansoy (1965: pl. X, figs 1-3), are almost identical to those of the RPI, in both shape and morphology. However, the fossettes of the Sinap specimen are slightly larger, likely due to the less advanced wear stage. Although the labial ribs of the RPI molars appear less prominent than those of PNT (but different wear stage may again exaggerate differences) both samples are placed close in the PCO analysis (Fig. 13) and within the range delineated by taxa included herein into *P. coelophrys* (see also Appendix 3: Figs S1-S4).

According to our observations and measurements the RPI taxon fits better *P. coelophrys* than any other taxon recognized herein. In fact, RPI's cranium has one of the largest described tooththrows. The absence of any postcranial material from that site prohibits any further comparison. Hence, we just confirm Geraads (1978) classification of that material as *Palaeotragus cf. coelophrys*.

Xirochori sample

The size of the single specimen from Xirochori approaches the mean values of *P. coelophrys* (Appendix 3: Fig. S11). Although the Xirochori molars are less worn, their morphology is almost identical to that of RPI, from which it differs in having a less distally pointed third lobe of the M₃. In any case the material is inadequate for certain conclusions and it is therefore referred to as *Palaeotragus* sp.

Nikiti-1 sample

The cranium and associated upper dentition from Nikiti-1 are significantly larger than those of *P. rouenii* and *P. microdon* and within the size range of *P. coelophrys* – *P. berislavicus* – *P. asiaticus*. The poor preservation status of NKT's skull does not allow for an extended comparison with *P. coelophrys* holotype from Maragheh and the two specimens belong to different sexes possibly exaggerating differences.

Dentally, the upper tooththrow from NKT differs from that of PNT (and RPI to the extent that morphologies can be recognized due to advanced wear) in the weaker styles (especially the metastyle) of the premolars, the less protruding paracone rib of the more squared P⁴, and the even weaker

metacone rib of the molars. Compared to *P. berislavicus* holotype (Korotkevich 1957), the labial ribs are equally developed in both samples. However, the P² and P³ of the Berislav giraffid are circular, while the same teeth from NKT are more square shaped. *P. berislavicus* demonstrates a triangular P⁴, while *Palaeotragus* from NKT has a square P⁴. Moreover, the P⁴ from NKT bears a lingual styloid that is missing from *P. berislavicus*.

In spite the bad preservation of most NKT postcranials, they clearly indicate a taxon proportionally close to *P. rouenii* though slightly more robust and with shorter metapodials. The NKT radius falls into the lower length range of *P. rouenii* (Appendix 3: Fig. S5) but it has a broader distal part. In that feature it agrees with *P. berislavicus* and *P. asiaticus* (Appendices 3, 4) although there is only one radius specimen available for each of these two taxa. In five complete radii of *P. rouenii* from several sites (except Kemiklitepe D, Turkey), the robusticity index (DT diaphysis/Length) is always below 10; it ranges between 10.4 and 11.2 in the four radii from NKT and between 10 and 13 in *P. coelophrys*–*P. berislavicus*–*P. asiaticus* (n=5). Unfortunately, data on the cranio-caudal diameter of radius are very limited. However, some scarce measurements (Geraads 1974), seems to indicate that *P. rouenii* had a significantly slenderer radius than NKT, at least proximally. In fact, the cranio-caudal diameter of NKT's radius is closer to that reported for *P. coelophrys*. However, when NKT specimens are compared with some *P. rouenii* radii from the sites Perivolaki and Nikiti-2, they did not seem to be more robust proximally. The radius curvature is a feature which also varies among the NKT sample, as well as among *P. rouenii*. Shafts' cross section is of similar crescent shape. Distally and cranially, the V-shaped formation for the adhesion of extensor carpi radialis muscle is equally prominent in NKT and *P. rouenii*. The distal articular surface is also very similar in the NKT palaeotrage and *P. rouenii*.

For a similar length (= 395 mm at average, excluding the likely young-adult NKT-137) the metacarpals of *P. coelophrys* appear significantly more robust (RI c. 12 instead of 9.5 in NKT), whereas for a similar robusticity (9.5) the metacarpals of *P. rouenii* are usually significantly longer (>430 mm in 8 out of 12 specimens from several sites) than in NKT. Concerning the morphology of the metacarpals, there are several differences between the NKT taxon and *P. rouenii* (Fig. 14). Proximally, the bone protrusion which separates the medial and lateral epicondyles is much more intense in *P. rouenii* than in NKT. Moreover, the medial epicondyle is of trapezoidal shape and the lateral epicondyle is of square shape in *P. rouenii*, while they are both of half-circle shape in NKT. As a result, the proximal articular surface has a trapezoidal shape in *P. rouenii* and a half-circle shape in the NKT specimens. Dorsally, the shaft is parallel to the bone axis medially, while the axis of the bone and the shaft are angled laterally in *P. rouenii*. That feature is very prominent proximally and it could be said that the cross section of the shaft has a shape of a right triangle. The same feature exists in NKT's metacarpals too, although it is much less

prominent, and as a result, the cross section of the shaft is more rectangular. Palmarly, the central trough extends throughout the whole bone in *P. rouenii* reaching at the trochlear, although it is considerably shallower distally than proximally. On the contrary, in the NKT metacarpals, the central trough seems to disappear at the distal 1/3 of the bone. Thus, the distal and palmar side of the metacarpal is completely flat in NKT. Distally the lateral condyle, seems to extend slightly more laterally in *P. rouenii* than in the NKT metacarpal. Finally, distally and dorsally, the shaft is somewhat more curved in *P. rouenii*, while in the NKT metacarpals is more flattened.

The tibia data are quite rare. According to the only fully preserved tibia available from NKT, its length falls within the *P. rouenii* range. However, it seems that the NKT taxon has broader distal epiphysis than *P. rouenii*, approaching *P. berislavicus* and *P. asiaticus* (Appendix 3: Fig. S7).

The NKT astragali and calcanei are intermediate in size between those of *P. rouenii* and *P. coelophrys*; the astragalus of *P. rouenii* from Perivolaki (PER) is more elongated than the NKT one, having a more rectangular shape (more square in NKT). The two astragali referred to *P. berislavicus* are smaller than the NKT ones and cannot be separated from those of *P. rouenii* (Appendix 3: Fig. S8).

By their proportions the metatarsals from NKT are placed in between those of *P. rouenii* and *P. coelophrys*; they are as long as the shorter known samples of *P. rouenii* but as wide as the largest individuals of this species. By these features, they approach better the two known metatarsals of *P. berislavicus* (Appendix 3: Figs S9, S10, S12).

Concerning the metatarsal morphology, there are several differences between the NKT taxon and *P. rouenii* on the proximal epiphysis (Fig. 15). Medially, the proximal articular surface is continuous in *P. rouenii*, while in NKT it is separated by a groove at the point between the two heads. Hence, the two heads of the medial epicondyle are more strongly separated in NKT. The shape of the dorsal head of the lateral epicondyle is half-circular in both NKT and *P. rouenii*, but in NKT is much more elongated. Moreover, in NKT that head is placed at a more obtuse angle to the axis of the proximal articular surface. In contrast, in *P. rouenii*, it is placed parallel to the proximal articular surface axis. As a result, the lateral epicondyle is of equal or of greater width to the medial epicondyle in NKT than in *P. rouenii*. In the medial epicondyle, the highest point of the dorsal face is more medial in *P. rouenii*. The plantar head of the medial epicondyle is extended plantarly in *P. rouenii* and it tilts laterally, while in NKT it is less extended and tilts medially. Those two heads are separated by a slight medial groove that is more marked in *P. rouenii* than in NKT. The central trough seems to vary significantly among NKT specimens in terms of depth and width. Finally, the distal part, relatively to the shaft, is wider in NKT than in *P. rouenii*.

Two assumptions can be formulated about the NKT taxon. The first one could be that two populations of *Palaeotragus* were present in Nikiti-1: a population of *P. rouenii* represented exclusively by postcranials and with slightly shorter

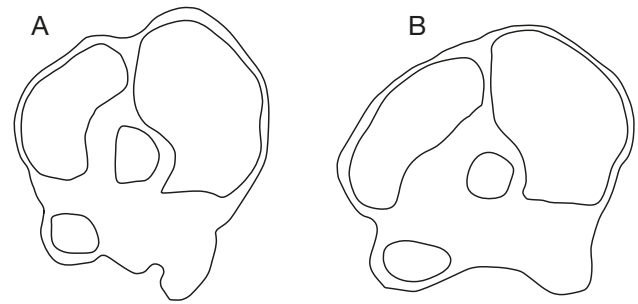


FIG. 15. — Morphological comparison of *Palaeotragus* Gaudry, 1861 proximal left metatarsal epiphyses: A, *P. rouenii* Gaudry, 1861 from Perivolaki; B, *P. aff. berislavicus* from Nikiti-1.

and more robust limbs than typically, and a population of a taxon close to *P. coelophrys* represented only by the partial cranium NKT-172. The case of coexistence of different *Palaeotragus* species in Greece is already well known (Iliopoulos 2003; Kostopoulos 2009). However, both the absence of any postcranial element from Nikiti that could be safely grouped with *P. coelophrys* and the proportional and morphological differences of the available postcranials from typical *P. rouenii* challenge this assumption. The homogeneity of the postcranial sample suggests a single medium sized *Palaeotragus* species, intermediate between *P. coelophrys* and *P. rouenii*. The combination of cranial and most of the postcranial evidence points to a population similar to *P. berislavicus*, though more data from both taxa are needed for definitive conclusions. Hence, we refer at the moment the NKT palaeotrage to *Palaeotragus* aff. *berislavicus*.

THE TUROLIAN GREEK LARGE PALAEOTRAGES

In our current knowledge large palaeotrages are relatively rare in the otherwise wealth Turolian faunas of Greece. Apart from the Samos Island (Kostopoulos 2009 and references therein) large *Palaeotragus* are reported from Kerassia (Euboea) and Thermopigi (Serres basin) sites (Iliopoulos 2003; Xafis *et al.* 2019). Samos material previously attributed to either *P. coelophrys* or *P. quadricornis* is herein considered as representing a single species, *P. coelophrys* (see also discussion in Kostopoulos 2009). Iliopoulos (2003) described two badly preserved mandibular fragments (K4/Δ8/1 and K4/Δ8/2) from Kerassia, most likely of the same individual, bearing P₃-M₃ right and P₂-part P₄ left, respectively. By its size and proportions (Lpm = 157.3 mm; Lp = 65.2 mm; Lm = 89.9 mm; data from Iliopoulos 2003), the Kerassia large palaeotrage fits within *P. coelophrys* and *P. berislavicus* size range. The rather advanced P₃ morphology and the relatively short molar row compared to the premolars might point to *P. berislavicus*, though material is not sufficient to provide any definitive determination.

Xafis *et al.* (2019) described from the Turolian fauna of Thermopigi (Greece) an upper premolar row, a left lower toothrow, an incisor and a canine as belonging to a larger palaeotrage than the well-documented *P. rouenii* from this site. Both the complete lower toothrow and the upper and lower

premolar row from Thermopigi are quite shorter than the larger palaeotragids recognized here (*P. coelophrys* and *P. berislavicus*; see Appendix 4). Metrically, the lower toothrow SIT 700 matches both *P. cf. coelophrys* from China and *P. asiaticus* from Kalmakpaj (Godina 1979, PIN 2432-84), whereas the P_3 shows similarly advanced molarization. Nevertheless, the Thermopigi specimens are just marginally longer or at the upper size range of *P. rouenii* and given the overall increased size (i.e., lengthening) of the local population of this species as indicated by the available postcranials, we could not exclude the possibility that those specimens may represent just a larger individual of *P. rouenii*.

CONCLUSION

Our analysis confirms previous suggestions that morphometrical discrimination among *P. coelophrys*, *P. expectans*, *P. borissaki*, *P. hoffstetteri*, *P. quadricornis* and possibly *P. moldavicus* is not sufficiently supported by present data, and hence all these taxa should be treated at the moment as possible synonyms under *P. coelophrys* (Rodler & Weithofer, 1890). Size differences between local samples (e.g., Maragheh vs. Starye Bogeny) might appear high in the light of extremely scarce evidence but they are not larger than size variance within the best recorded *P. rouenii* or other giraffid species (e.g., Ríos *et al.* 2016, 2017). The dP_3/P_3 molarization is usually less advanced in *P. coelophrys* than in *P. rouenii*, though morphological variation does exist and data are limited for statistically safe and definitive conclusions. Limb shortening and metapodial robusticity possibly increase from the Vallesian peri-Black Sea samples to the Turolian ones from Anatolia-Iran but the scarcity of the material does not allow to properly test this hypothesis.

In accordance with the Black Sea record, *P. coelophrys* appears to have a strong signal in the Vallesian faunas from Axios Valley, N. Greece, certainly recorded in Pentalophos and most probably in Ravin de la Pluie; the Xirochori mandible may also belong to the same species but material is insufficient. Turolian reports of the species in continental Greece are rather dubious but the taxon has a strong presence in contemporaneous faunas south of Caucasus from where probably expanded westwards in the Anatolian domain.

Palaeotragus asiaticus and *P. cf. coelophrys* from the Turolian equivalent of Central Asia (Kyrgyzstan, Uzbekistan, and Kazakhstan) and China respectively, are probably closely related to each other. They could represent an eastward offshoot of Iranian *P. coelophrys* or – as the biometric data suggest – they might originated from another taxon; in that case the Vallesian *P. berislavicus* from Ukraine would be an ideal ancestor by means of morphometry. The Berislav palaeotrage is placed morphometrically between *P. rouenii* and *P. coelophrys*, and it probably invaded Southern Balkans at the end of the Vallesian, documented in the Nikiti-1 fauna. It is also possible that the species survived here quite later, as the Kerassia record may imply, but data are once again inadequate for certain conclusions.

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APPENDICES

APPENDIX 1. — Biometrical variables and selected premolar features of Late Miocene Eurasian palaeotragines used in multivariate analyses (Fig. 11 main text; Appendix 3: Figs S1, S2).

Lpm	Lower toothrow length;	Mt-Bdia	Breadth of the metatarsals' diaphysis;
Lp	Lower premolar row length;	Mt-Ddia	Depth of the metatarsals' diaphysis;
Lm	Lower molar row length; ×100 (p/pm); ×100 (p/m); ×100 (m/pm);	Mt-R.I.	Metatarsals R.I.;
LPM	Upper toothrow length;	Mt-Bp	Breadth of the metatarsals proximal end;
LP	Upper premolar row length;	Mt-Dp	Depth of the metatarsals proximal end;
LM	Upper molar row length; ×100 (P/PM) ×100 (P/M) ×100 (M/PM)	P ₃ morphology	
R-GL	Greatest length of the radii;	0	primitive ruminant morphology;
R-Bp	Breadth of the radii proximal end;	1	molarized.
Mc-GL	Greatest length of the metacarpals;	P ₄ lingual groove	
Mc-Bdia	Breadth of the metacarpals' diaphysis;	0	absent;
Mc-R.I.	Metacarpals' robusticity index (×100);	1	present.
T-Bd	Breadth of the tibiae distal end;	P ₄ lingual stylids	
T-Dd	Depth of the tibiae distal end;	0	absent;
A-GLl	Greatest lateral length of the astragali;	1	present.
A-Bd	Breadth of the astragali distal end;	P ₄ occusal shape	
Mt-GL	Greatest length of the metatarsals	0	circular;
		1	triangular;
		2	squared.

APPENDIX 2. — Tables 1-14: Measurements of *Palaeotragus coelophrys* (Rodler & Weithofer, 1890).

TABLE 1. — Measurements of *Palaeotragus coelophrys* upper teeth from Pentalophos (PNT). Abbreviations: **LPM**, Length of upper toothrow; **LP**, Length of upper premolar row; **LM**, Length of upper molar row; **LP₂**, Length of P₂; **WP₂**, Width of P₂; **LP₃**, Length of P₃; **WP₃**, Width of P₃; **LP₄**, Length of P₄; **WP₄**, Width of P₄; **LM₁**, Length of M₁; **WaM₁**, Width of M₁ anterior lobe; **WpM₁**, Width of M₁ posterior lobe; **LM₂**, Length of M₂; **WaM₂**, Width of M₂ anterior lobe; **WpM₂**, Width of M₂ posterior lobe; **LM₃**, Length of M₃; **WaM₃**, Width of M₃ anterior lobe; **WpM₃**, Width of M₃ posterior lobe (in mm).

Measurements	Specimen Reg. no. PNT-					
	113	165	152	163	161	164
LPM	138.45	—	—	—	—	—
LP	59.5	—	—	—	—	—
LM	82.14	—	—	—	—	—
LP ₂	17.42	—	20.06	19.32	19.02	—
WP ₂	19.54	—	18.73	19.54	19.17	—
LP ₃	20.21	—	20.53	20.8	—	21.4
WP ₃	21.2	—	21.65	24.37	—	24.1
LP ₄	20.47	—	—	—	—	—
WP ₄	26.05	—	—	—	—	—
LM ₁	26.8	—	—	—	—	—
WaM ₁	28.13	—	—	—	—	—
WpM ₁	27.62	—	—	—	—	—
LM ₂	30.1	28.6	—	—	—	—
WaM ₂	31.06	28.15	—	—	—	—
WpM ₂	26.8	26.23	—	—	—	—
LM ₃	29.36	31.64	—	—	—	—
WaM ₃	18.1	30.26	—	—	—	—
WpM ₃	24.18	26.74	—	—	—	—

TABLE 2. — Measurements of *Palaeotragus coelophrys* lower teeth from Pentalophos (PNT). Abbreviations: **LM**, Length of lower molar row; **LP₂**, Length of P₂; **WP₂**, Width of P₂; **LP₃**, Length of P₃; **WP₃**, Width of P₃; **LP₄**, Length of P₄; **WP₄**, Width of P₄; **LM₁**, Length of M₁; **WaM₁**, Width of M₁ anterior lobe; **WpM₁**, Width of M₁ posterior lobe; **LM₂**, Length of M₂; **WaM₂**, Width of M₂ anterior lobe; **WpM₂**, Width of M₂ posterior lobe; **LM₃**, Length of M₃; **WaM₃**, Width of M₃ anterior lobe; **WpM₃**, Width of M₃ posterior lobe (in mm).

Measurements	Specimen Reg. no. PNT-328F
LM	97
LM ₁	28.18
WaM ₁	18.55
WpM ₁	21.45
LM ₂	28.05
WaM ₂	22.31
WpM ₂	23.5
LM ₃	38.08
WaM ₃	21.37
WpM ₃	21.96
W3M ₃	16.7

TABLE 3. — Measurements of *Palaeotragus coelophrys* lower deciduous teeth from Pentalophos (PNT). Abbreviations: **LdP**, Length of deciduous lower pre-molar row; **LdP₂**, Length of dP₂; **WdP₂**, Width of dP₂; **LdP₃**, Length of dP₃; **WdP₃**, Width of dP₃; **LdP₄**, Length of dP₄; **WdP₄**, Width of dP₄; **LM₁**, Length of M₁; **WaM₁**, Width of M₁ anterior lobe; **WpM₁**, Width of M₁ posterior lobe (in mm).

Measurements	Specimen Reg. no. PNT-121F
LdP	69.79
LdP ₂	15.82
WdP ₂	8.38
LdP ₃	22.98
WdP ₃	9.71
LdP ₄	31.03
WdP ₄	11.34
WmdP ₄	16.23
Wdpd ₄	16.14
LM ₁	28.88
WaM ₁	20.12
WpM ₁	20.06

TABLE 4. — Measurements of *Palaeotragus coelophrys* metatarsals from Pentalophos (PNT). Abbreviations: **Mt-Bp**, Breadth of metatarsal proximal end; **Mt-Dp**, Depth of metatarsal proximal end; **Mt-Bdia**, Breadth of metatarsal at the middle of the shaft; **Mt-Ddia**, Depth of metatarsal at the middle of the shaft (in mm).

Measurements	Specimen Reg. no.	
	PNT-119F	PNT-114F
Mt-Bp	62.98	60.39
Mt-Dp	60.75	57.58
Mt-Bdia	37.02	40.66
Mt-Ddia	37.20	32.81

TABLE 5. — Measurements of *Palaeotragus cf. coelophrys* upper teeth from the Ravin de la Pluie (cranium RPI-91B). Abbreviations: **LPM**, Length of upper toothrow; **LP**, Length of upper premolar row; **LM**, Length of upper molar row; **LP₂**, Length of P₂; **WP₂**, Width of P₂; **LP₃**, Length of P₃; **WP₃**, Width of P₃; **LP₄**, Length of P₄; **WP₄**, Width of P₄; **LM₁**, Length of M₁; **WaM₁**, Width of M₁ anterior lobe; **WpM₁**, Width of M₁ posterior lobe; **LM₂**, Length of M₂; **WaM₂**, Width of M₂ anterior lobe; **WpM₂**, Width of M₂ posterior lobe; **LM₃**, Length of M₃; **WaM₃**, Width of M₃ anterior lobe; **WpM₃**, Width of M₃ posterior lobe (in mm).

Measurements	Specimen Reg. no.	
	RPI-91B left	RPI-91B right
LPM	152.25	147.4
LP	68.68	62.81
LM	94.07	85.41
LP ₂	18.82	22.88
WP ₂	20.44	20.62
LP ₃	21.61	21.25
WP ₃	23.67	22.61
LP ₄	21.14	19.21
WP ₄	26.04	26.64
LM ₁	24.82	23.96
WaM ₁	26.85	27.42
WpM ₁	26.28	26.85
LM ₂	28.50	30.51
WaM ₂	29.21	28.70
WpM ₂	27.81	27.38
LM ₃	28.63	28.47
WaM ₃	28.18	26.41
WpM ₃	24.59	22.41

TABLE 6. — Measurements of *Palaeotragus cf. coelophrys* lower teeth from Ravin de la Pluie (RPI). Abbreviations: **LM**, Length of lower molar row; **LP₂**, Length of P₂; **WP₂**, Width of P₂; **LP₃**, Length of P₃; **WP₃**, Width of P₃; **LP₄**, Length of P₄; **WP₄**, Width of P₄; **LM₁**, Length of M₁; **WaM₁**, Width of M₁ anterior lobe; **WpM₁**, Width of M₁ posterior lobe; **LM₂**, Length of M₂; **WaM₂**, Width of M₂ anterior lobe; **WpM₂**, Width of M₂ posterior lobe; **LM₃**, Length of M₃; **WaM₃**, Width of M₃ anterior lobe; **WpM₃**, Width of M₃ posterior lobe (in mm).

Measurements	Specimen Reg. no. RPI-104
LM	89.59
LM ₁	24.92
WaM ₁	17.09
WpM ₁	17.38
LM ₂	25.80
WaM ₂	18.30
WpM ₂	19.24
LM ₃	39.15
WaM ₃	18.89
WpM ₃	19.91
W3M ₃	10.97

TABLE 7. — Measurements of *Palaeotragus* sp. lower teeth from Xirochori (XIR). **LM₂**, Length of M₂; **WaM₂**, Width of M₂ anterior lobe; **WpM₂**, Width of M₂ posterior lobe; **LM₃**, Length of M₃; **WaM₃**, Width of M₃ anterior lobe; **WpM₃**, Width of M₃ posterior lobe (in mm).

Measurements	Specimen Reg. no. XIR-24
LM ₂	26.93
WaM ₂	19.55
WpM ₃	19.04
LM ₃	40.33
WaM ₃	20.06
WpM ₃	20.00
W3M ₃	11.95

TABLE 8. — Measurements of *Palaeotragus aff. berislavicus* upper teeth from the Nikiti (cranium NKT-172). Abbreviations: **LPM**, Length of upper toothrow; **LP**, Length of upper premolar row; **LM**, Length of upper molar row; **LP₂**, Length of P₂; **WP₂**, Width of P₂; **LP₃**, Length of P₃; **WP₃**, Width of P₃; **LP₄**, Length of P₄; **WP₄**, Width of P₄; **LM₁**, Length of M₁; **WaM₁**, Width of M₁ anterior lobe; **WpM₁**, Width of M₁ posterior lobe; **LM₂**, Length of M₂; **WaM₂**, Width of M₂ anterior lobe; **WpM₂**, Width of M₂ posterior lobe; **LM₃**, Length of M₃; **WaM₃**, Width of M₃ anterior lobe; **WpM₃**, Width of M₃ posterior lobe (in mm).

Measurements	Specimen Reg. no.	
	NKT-172 left	NKT-172 right
LPM	141.41	146.1
LP	62.35	63.49
LM	85.10	84.9
LP ₂	22.40	21.61
WP ₂	18.93	18.91
LP ₃	20.79	21.20
WP ₃	23.5	21.18
LP ₄	20.66	20.14
WP ₄	27.26	24.72
LM ₁	25.59	29.46
WaM ₁	—	27.80
WpM ₁	—	26.30
LM ₂	29.87	30.22
WaM ₂	33.29	28.14
WpM ₂	—	28.25
LM ₃	27.66	30.58
WaM ₃	141.41	31.63
WpM ₃	62.35	27.68

TABLE 9. — Measurements of *Palaeotragus* aff. *berislavicus* radii from Nikiti (NKT). **R-GL**, Greatest Length; **R-Bp**, Breadth of the proximal end; **R-Dp**, Depth of the proximal end; **R-Bpart**, Breadth of the proximal articular surface; **R-Bdia**, Breadth in the middle of the shaft; **R-Ddia**, Depth in the middle of the shaft; **R-Bd**, Breadth of the distal end; **R-Dd**, Depth of the distal end (in mm).

Measurements	Specimen Reg. no.				
	NKT-159	NKT-167	NKT-156	NKT-155	NKT-169
R-GL	449	—	462	476	470
R-Bp	82.42	88.05	81.24	82.82	71.45
R-Dp	53.25	46.82	44.64	49.00	40.74
R-Bpart	—	—	67.95	73.83	—
R-Bdia	48.18	48.25	51.47	49.61	51.25
R-Ddia	36.03	35.02	41.64	33.26	30.77
R-Bd	75.51	—	75.66	80.59	73.95
R-Dd	55.17	—	54.47	51.78	42.60

TABLE 10. — Measurements of *Palaeotragus* aff. *berislavicus* metacarpals from Nikiti (NKT). **Mc-GL**, Greatest Length; **Mc-Bp**, Breadth of the proximal end; **Mc-Dp**, Depth of the proximal end; **Mc-Bdia**, Breadth in the middle of the shaft; **Mc-Ddia**, Depth in the middle of the shaft; **Mc-Bd**, Breadth of the distal end; **Mc-Dd**, Depth of the distal end (in mm).

Measurements	Specimen Reg. no.				
	NKT-141	NKT-137	NKT-131	NKT-67	NKT-26
Mc-GL	409.0	378.0	380.0	—	—
Mc-Bp	64.68	57.89	64.60	70.37	61.15
Mc-Dp	40.25	32.71	46.55	39.63	41.28
Mc-Bdia	35.80	31.68	32.79	42.30	—
Mc-Ddia	31.44	31.21	30.74	30.79	—
Mc-Bd	63.41	60.26	61.65	—	—
Mc-Dd	38.19	35.56	37.16	—	—

TABLE 11. — Measurements of *Palaeotragus* aff. *berislavicus* tibiae from Nikiti (NKT). **T-GL**, Greatest Length; **T-Bp**, Breadth of the proximal end; **T-Dp**, Depth of the proximal end; **T-Bdia**, Breadth in the middle of the shaft; **T-Ddia**, Depth in the middle of the shaft; **T-Bd**, Breadth of the distal end; **T-Dd**, Depth of the distal end (in mm).

Measurements	Specimen Reg. no.		
	NKT-271	NKT-150	NKT-154
T-GL	451.0	—	—
T-Bp	82.57	—	—
T-Dp	82.07	—	—
T-Bdia	50.40	45.49	47.78
T-Ddia	35.53	36.98	39.60
T-Bd	81.60	67.53	68.90
T-Dd	55.79	54.73	45.77

TABLE 12. — Measurements of *Palaeotragus* aff. *berislavicus* astragali from Nikiti (NKT). **A-GLI**, Greatest lateral Length; **A-GLm**, Greatest medial Length; **A-Bp**, Breadth of the proximal end; **A-DI**, Lateral Depth; **A-Dm**, Medial Depth; **A-Bd**, Breadth of the distal end (in mm).

Measurements	Specimen Reg. no.		
	NKT-267	NKT-163	NKT-266
A-GLI	78.58	77.83	78.64
A-GLm	70.75	72.57	67.41
A-Bp	49.29	52.01	50.17
A-DI	45.40	43.42	49.20
A-Dm	47.25	46.36	51.30
A-Bd	47.03	50.96	50.00

TABLE 13. — Measurements of *Palaeotragus* aff. *berislavicus* calcanei from Nikiti (NKT). **C-GLI**, Greatest Length; **C-GB**, Greatest Breadth (in mm).

Measurements	Specimen Reg. no.	
	NKT-268	NKT-153
C-GL	136.61	147.35
C-GB	45.76	55.00

TABLE 14. — Measurements of *Palaeotragus* aff. *berislavicus* metatarsals from Nikiti (NKT). **Mt-GL**, Greatest Length; **Mt-Bp**, Breadth of the proximal end; **Mt-Dp**, Depth of the proximal end; **Mt-Bdia**, Breadth in the middle of the shaft; **Mt-Ddia**, Depth in the middle of the shaft; **Mt-Bd**, Breadth of the distal end; **Mt-Dd**, Depth of the distal end (in mm).

Measurements	Specimen Reg. no.									
	NKT-136	NKT-160	NKT-139	NKT-133	NKT-144	NKT-138	NKT-168	NKT-151	NKT-140	NKT-132
Mt-GL	414	409	386	412	415	436	—	—	—	—
Mt-Bp	58.86	60.85	52.45	62.76	59.50	59.46	56.44	—	58.42	54.81
Mt-Dp	57.83	49.49	54.62	53.06	45.71	—	50.44	—	44.82	—
Mt-Bdia	34.22	41.25	33.65	49.60	33.35	31.94	38.48	40.73	40.98	32.80
Mt-Ddia	33.58	34.93	29.68	28.75	33.05	30.21	36.30	31.35	34.00	35.18
Mt-Bd	—	64.80	57.84	60.56	57.89	—	—	61.19	—	—
Mt-Dd	—	38.49	35.30	40.47	35.68	33.92	—	37.81	—	—

APPENDIX 3. — Figures S1 à S12.

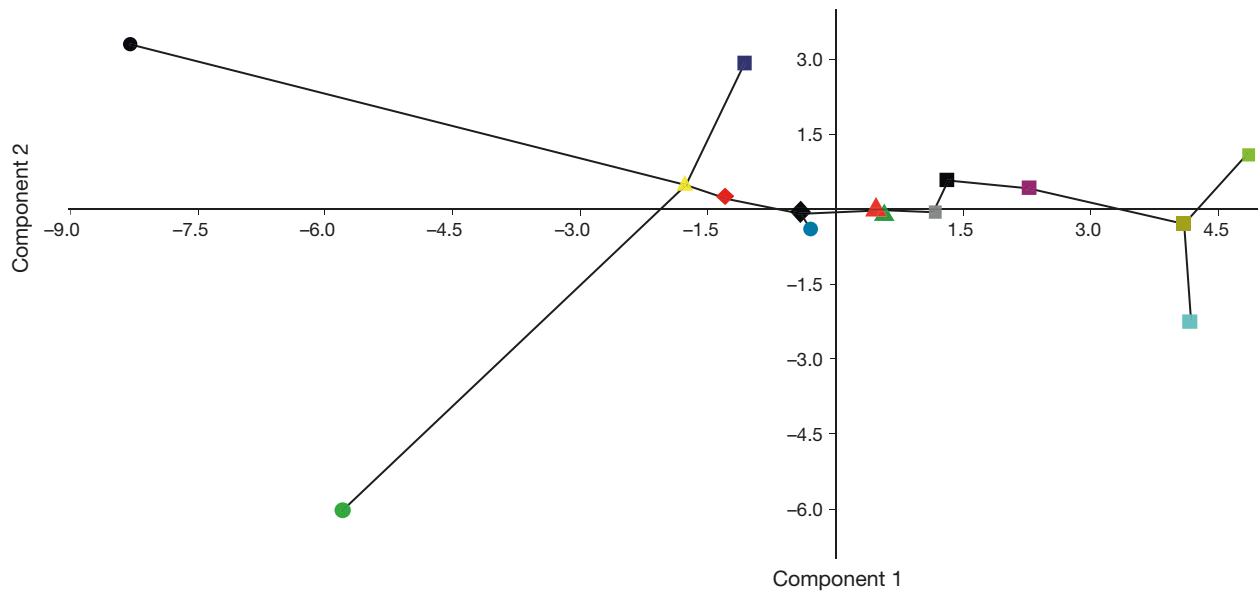


FIG. S1. — Principal Components Analysis (plane of components 1 and 2, representing 40,3% and 14,4% respectively of initial data variance) comparing 27 biometrical variables and 4 selected features of premolar morphology of the Late Miocene Eurasian palaeotragas as well as the Vallesian Greek samples studied here.

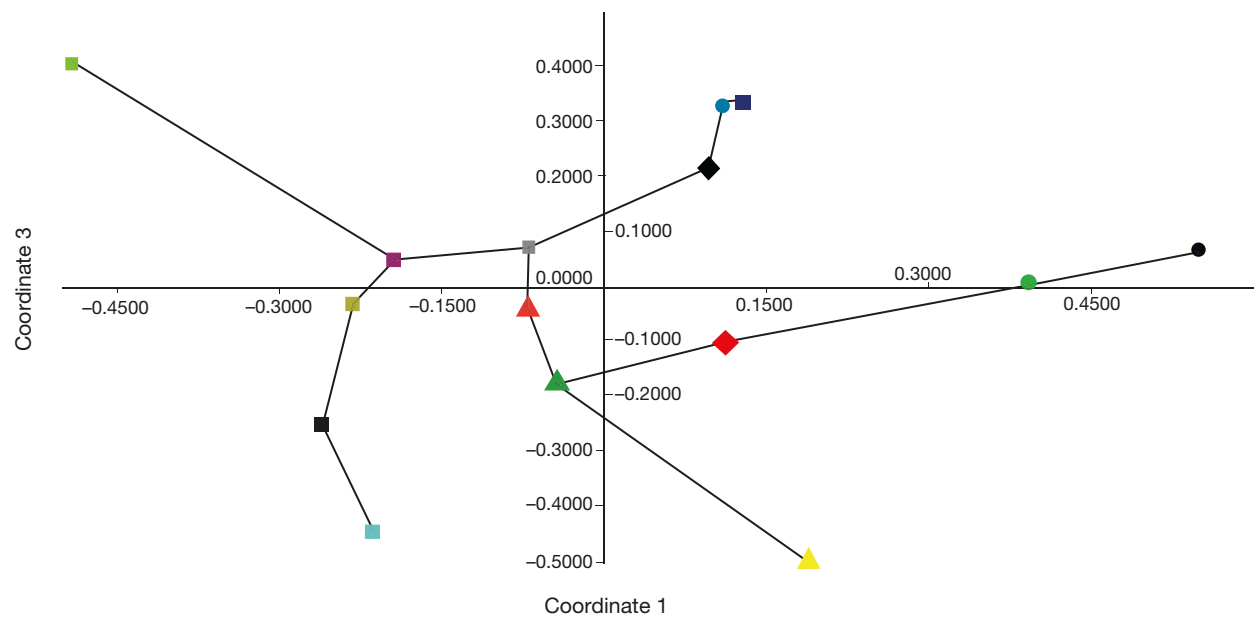
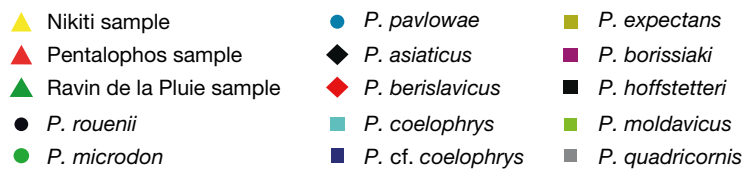


FIG. S2. — Principal Coordinates Analysis (plane of coordinates 1 and 3, representing 39,5% and 10,4% respectively of initial data variance) comparing 27 biometrical variables and 4 selected features of premolar morphology of the Late Miocene Eurasian palaeotragas as well as the Vallesian Greek samples studied here.

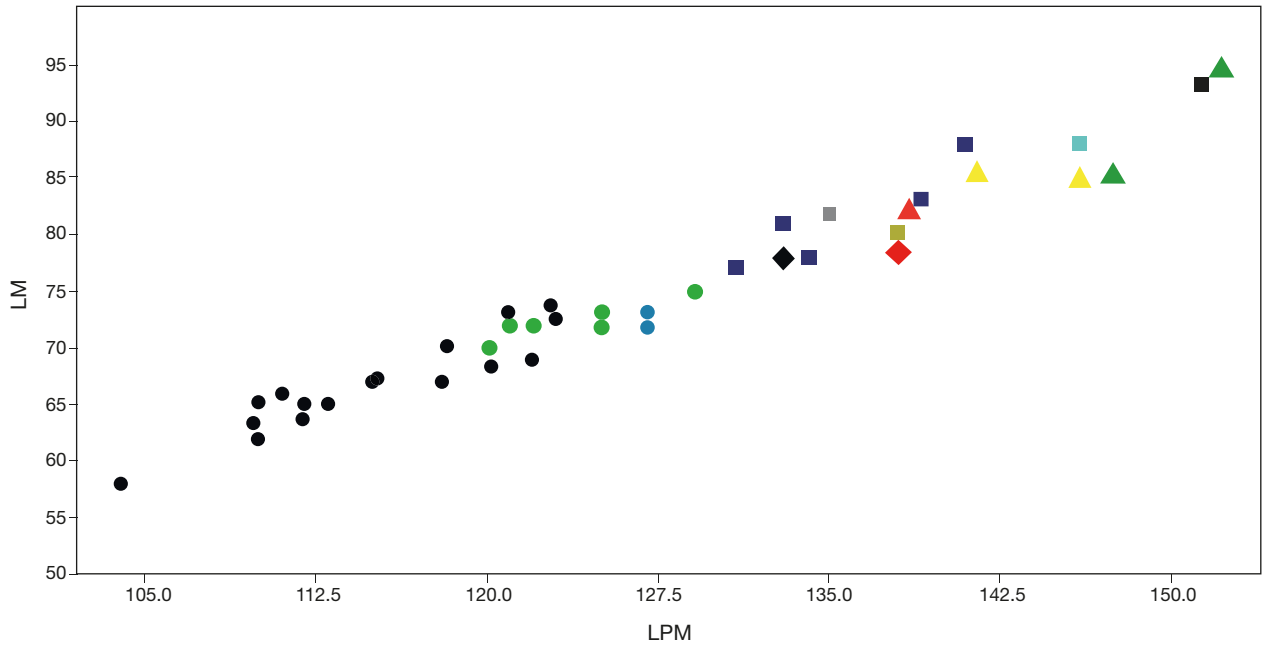


FIG. S3. — Scatter plot of upper teeth proportions of the Late Miocene Eurasian palaeotragines as well as the Vallesian Greek samples studied here. Abbreviations: **LM**, length of molar row; **LPM**, length of toothrow.

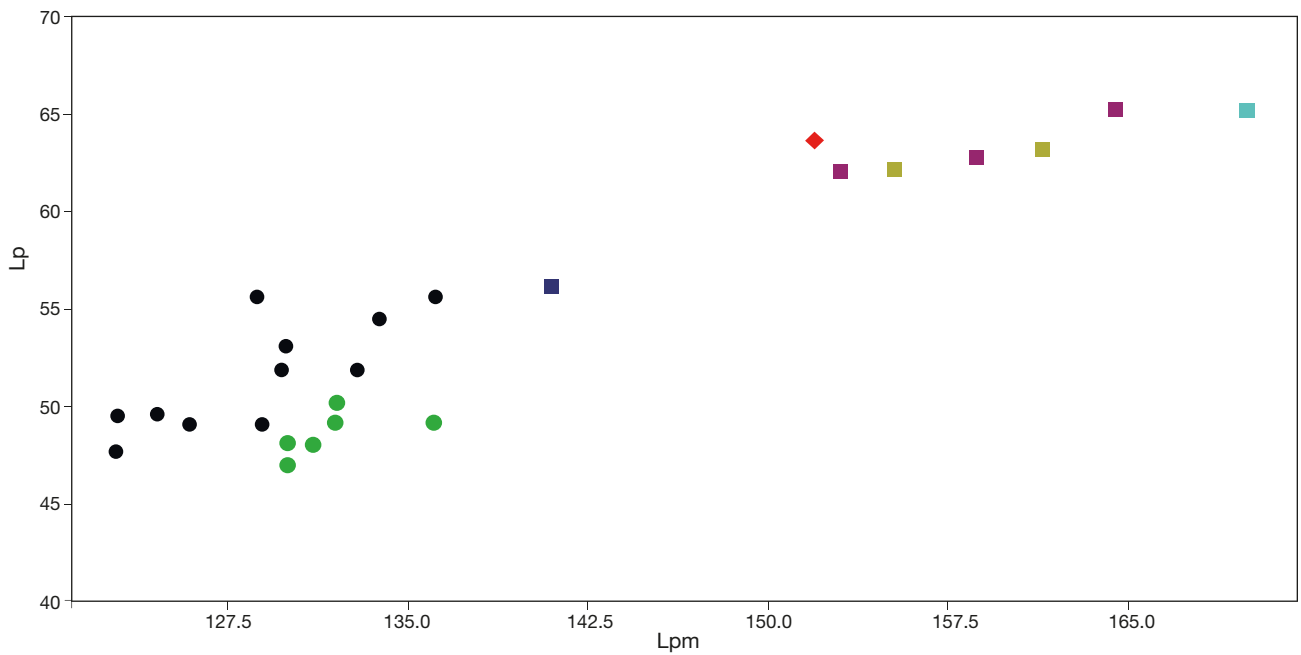
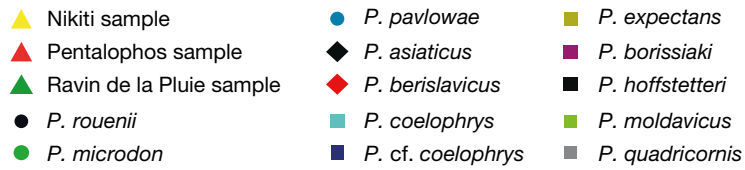


FIG. S4. — Scatter plot of lower teeth proportions of the Late Miocene Eurasian palaeotragines. Abbreviations: **Lp**, length of premolar row; **Lpm**, length of toothrow.

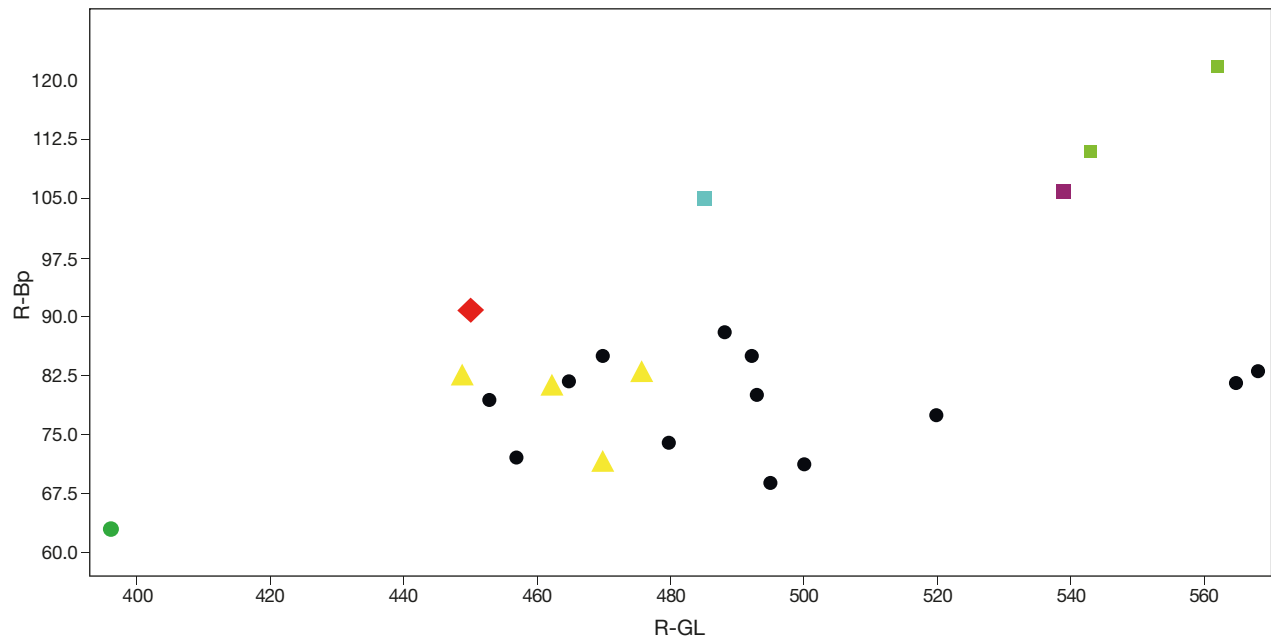


FIG. S5. — Scatter plot of radii proportions of the Late Miocene Eurasian palaeotrages as well as the Nikiti (NKT) sample. Abbreviations: **R-Bp**, breadth of the proximal epiphysis; **R-GL**, greatest length.

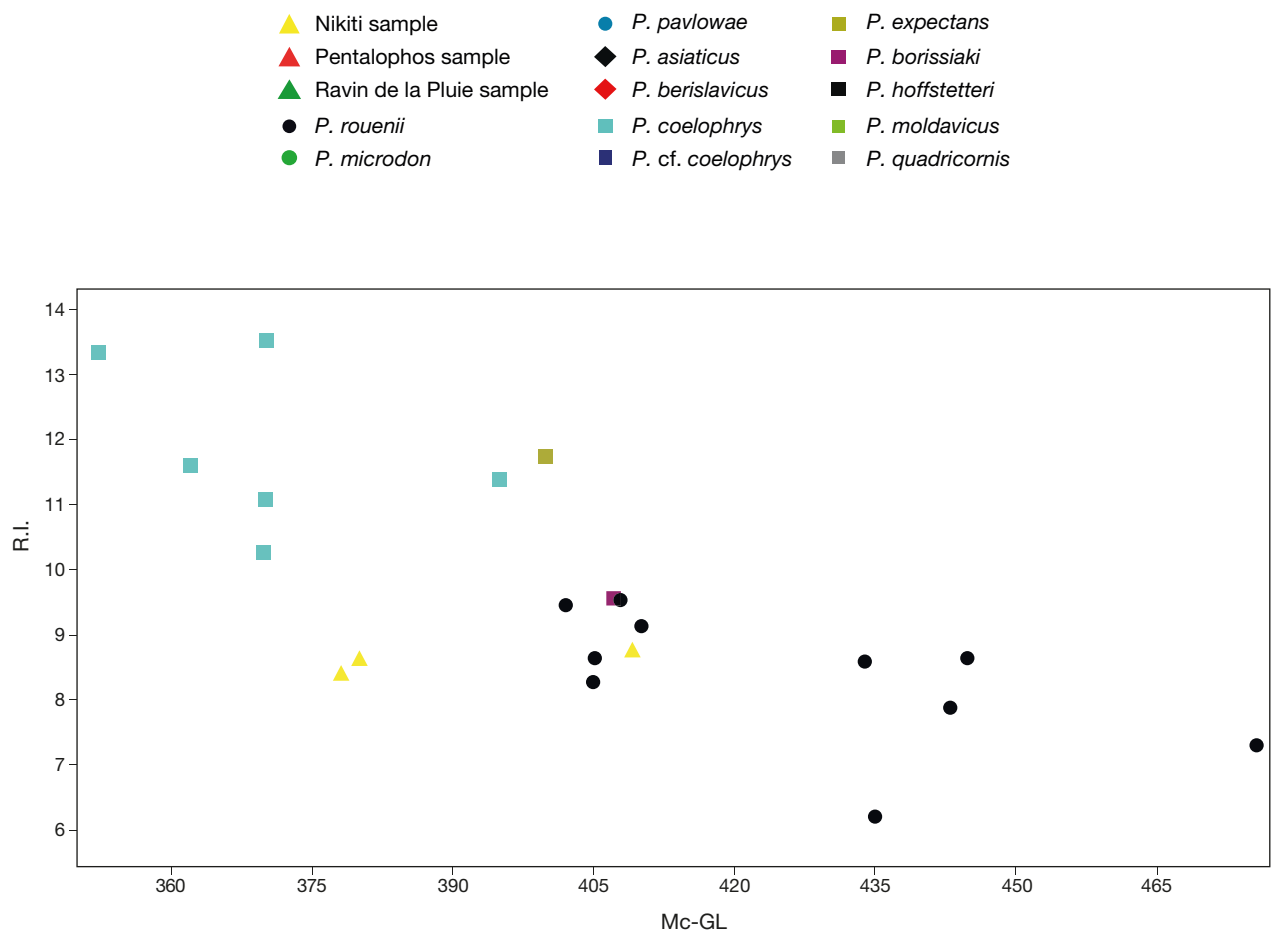


FIG. S6. — Scatter plot of metacarpal proportions of the Late Miocene Eurasian palaeotrages as well as the Nikiti (NKT) sample. Abbreviations: **R.I.**, robusticity index; **Mc-GL**, metacarpal greatest length.

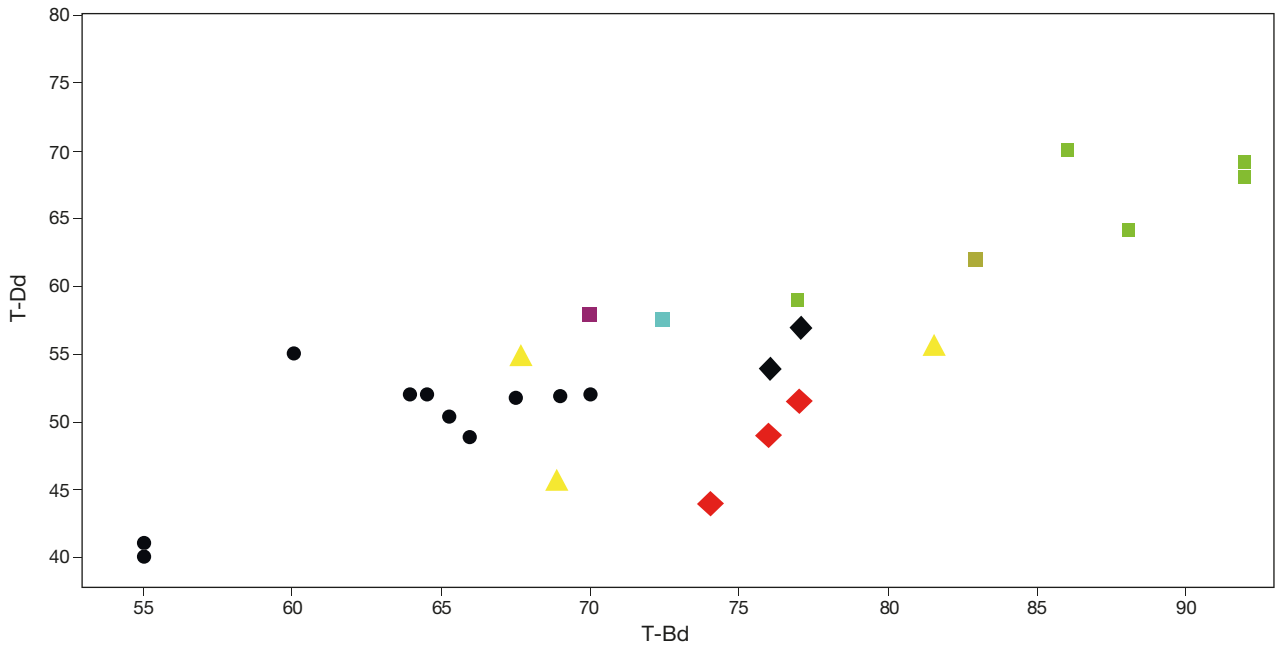


FIG. S7. — Scatter plot of distal tibiae proportions of the Late Miocene Eurasian palaeotrages as well as the Nikiti (NKT) sample. Abbreviations: **T-Dd**, depth of the distal end; **T-Bd**, breadth of the distal end.

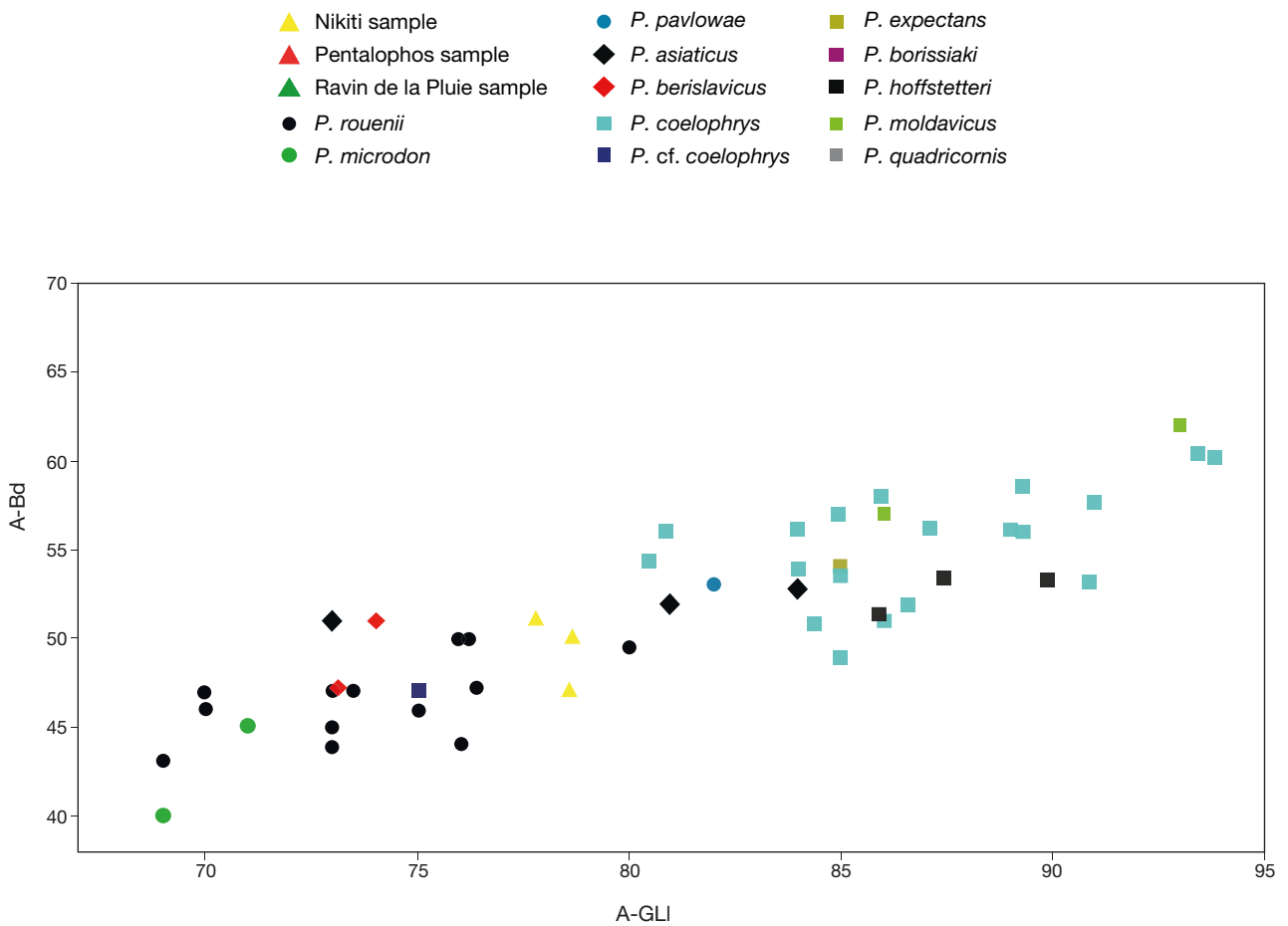


FIG. S8. — Scatter plot of astragali proportions of the Late Miocene Eurasian palaeotrages as well as the Nikiti (NKT) sample. Abbreviations: **A-Bd**, breadth of the distal end; **A-GLI**, greatest lateral length.

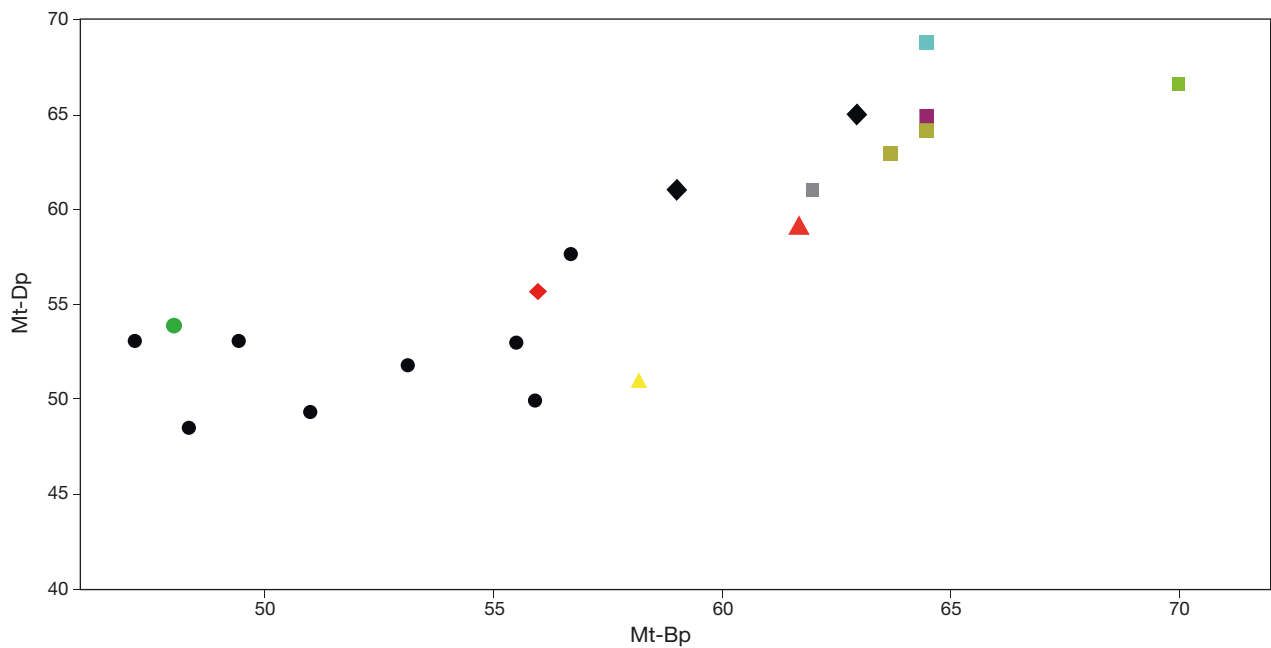


FIG. S9. — Scatter plot of proximal metatarsal proportions of the Late Miocene Eurasian palaeotragines as well as the Nikiti (NKT) and Pentalophos (PNT) samples. Abbreviations: **Mt-Dd**, depth of the proximal end; **Mt-Bd**, breadth of the proximal end.

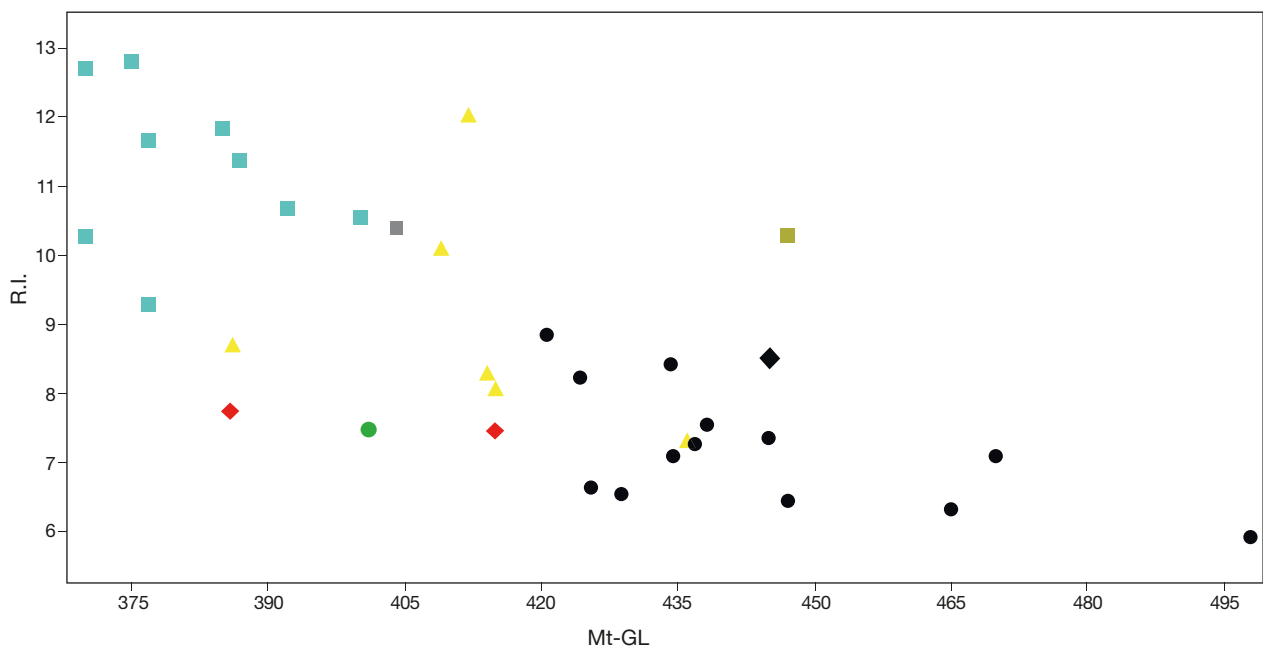
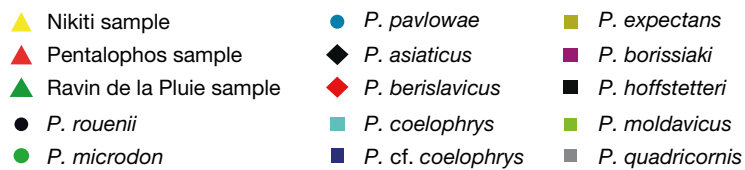


FIG. S10. — Scatter plot of metatarsal proportions of the Late Miocene Eurasian palaeotragines as well as the Nikiti (NKT) sample. Abbreviations: **R.I.**, robusticity index; **Mt-GL**, metatarsal greatest length.

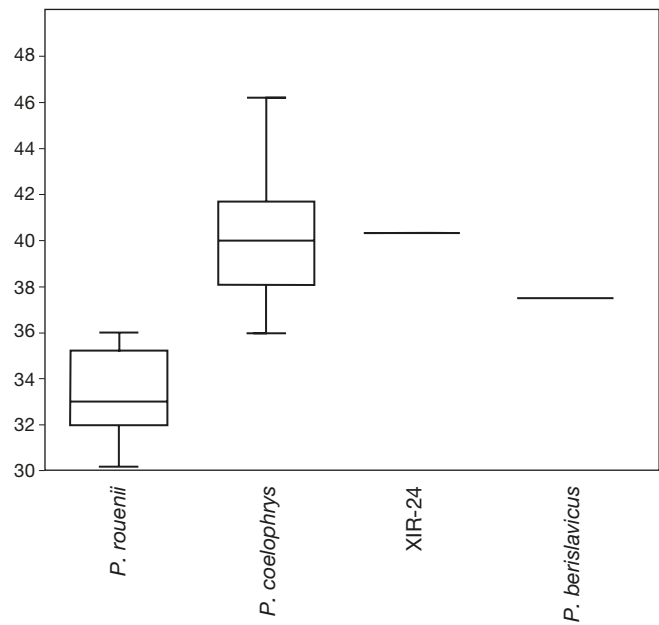


FIG. S11. — Boxplot comparing M₃ length among different Late Miocene Eurasian *Palaeotragus* Gaudry, 1861 forms.

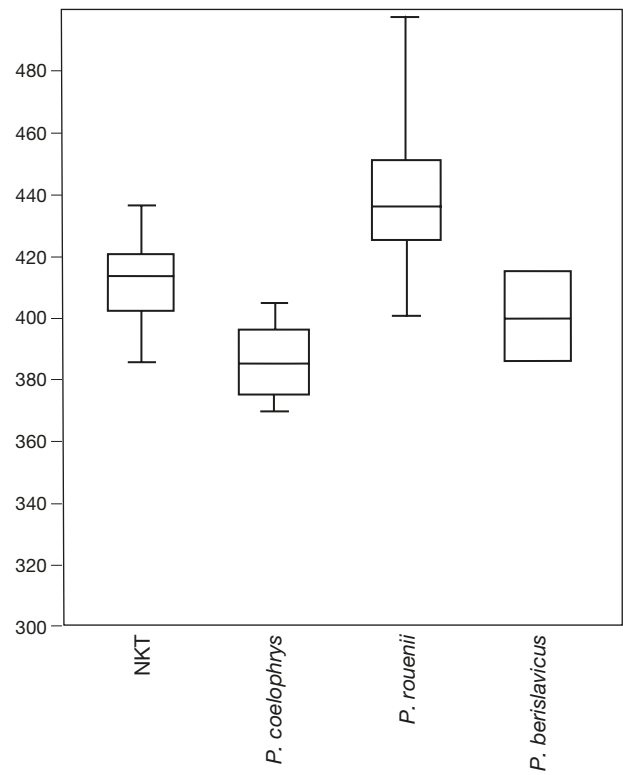


FIG. S12. — Boxplot comparing metatarsal length among different Late Miocene Eurasian *Palaeotragus* Gaudry, 1861 forms.

APPENDIX 4. — Average measurements of the biometrical variables and the selected premolar features of the Late Miocene Eurasian palaeotragids. Type localities marked in bold. For character abbreviations see Appendix 1.

	Site	Lpm	Lp	Lm	Lp/Lm	Lp/Lpm	Lm/Lpm
<i>P. rouenii</i>	Pikermi	128.7	55.5	76.6	72.45	43.12	59.51
	Cimişlia	130	53	77.8	68.12	40.76	59.84
	Taraclia	—	—	78	—	—	—
	Kerassia	134.95	54.85	80.9	67.79	40.64	59.94
	Ditiko	126	49	78	62.82	38.88	61.90
	Cioburciu	—	—	—	—	—	—
	Samos	128.8	50.6	78.75	64.25	39.28	61.14
	Todorovo	—	—	—	—	—	—
	Kemiklitepe	—	—	—	—	—	—
	Novo-Elisabetovka	—	—	—	—	—	—
	Strumyani	122.9	47.6	75.3	63.21	38.73	61.26
	Hadjidimovo	129.8	51.8	78	66.41	39.90	60.09
	Akkasdagli	129	49	77	63.63	37.98	59.68
	Thermopigi	123	49.34	73.66	66.98	40.11	59.88
	Kryopigi	—	—	77.6	—	—	—
	Perivolaki	—	78.6	—	—	—	—
<i>P. coelophrys</i>	Maragha	170	65	100	65	38.23	58.82
	Samos	—	—	—	—	—	—
<i>P. cf. coelophrys</i>	China	144	56.66	87.5	64.75	39.35	60
<i>P. berislavicus</i>	Berislav	152	63.6	89.7	70.90	41.84	59.01
<i>P. microdon</i>	Kansu/Bohlin 1926	131.5	48.16	82.16	58.61	36.62	62.48
	Shansi/1926	133	50	85.5	58.47	37.59	64.28
	China/Bohlin 1926	—	—	—	—	—	—
<i>P. quadricornis</i>	Samos	—	—	—	—	—	—
<i>P. asiaticus</i>	Ortok	—	—	—	—	—	—
	Kalmakpai	141	—	—	—	—	—
	Pavlodar	—	—	—	—	—	—
<i>P. hoffstetteri</i>	Sinap	—	—	—	—	—	—
<i>P. pavlowae</i>	Grebeniki	—	—	—	—	—	—
	Blagodarnenskaya	—	—	—	—	—	—
<i>P. moldavicus</i>	Raspopeny	—	—	98	—	—	—
	Korsakova Assembly	—	—	—	—	—	—
<i>P. borissiaki</i>	Eldari	158.75	63.5	99.4	63.88	40	62.61
<i>P. expectans</i>	Sevastopol	158.35	62.5	96.25	64.93	39.46	60.78
	Varnitsa	—	—	—	—	—	—
<i>P. aff. berislavicus</i>	NKT	—	—	—	—	—	—
<i>P. coelophrys</i>	PNT	—	97	—	—	—	—
<i>P. cf. coelophrys</i>	RPI	—	—	89.59	—	—	—

APPENDIX 4. — Continuation.

	Site	LPM	LP	LM	P/M	P/PM	M/PM
<i>P. rouenii</i>	Pikermi	118.2	51.9	67.33	77.07	43.90	56.9656
	Cimişlia	114.5	50	65.4	76.45	43.66	57.11
	Taraclia	112.5	50.25	64.75	77.60	44.67	57.56
	Kerassia	—	—	—	—	—	—
	Ditiko	121	54	73	73.97	44.62	60.33
	Cioburciu	—	—	64	—	—	—
	Samos	112.82	49.5	65.21	75.90	43.87	57.80
	Todorovo	—	—	—	—	—	—
	Kemiklitepe	—	—	—	—	—	—
	Novo-Elisabetovka	—	—	—	—	—	—
	Strumyani	—	—	—	—	—	—
	Hadjidimovo	—	—	—	—	—	—
	Akkasdagli	123	53	72.7	72.90	43.08	59.10
	Thermopigi	—	—	—	—	—	—
	Kryopigi	—	—	—	—	—	—
	Perivolaki	—	—	—	—	—	—
<i>P. coelophrys</i>	Maragha	146	66	88	75	45.20	60.27
	Samos	—	—	—	—	—	—
<i>P. cf. coelophrys</i>	China	135.6	60	81.4	73.71	44.24	60
<i>P. berislavicus</i>	Berislav	138	63	78.5	80.25	45.65	56.88
<i>P. microdon</i>	Kansu/Bohlin 1926	—	—	—	—	—	—
	Shansi/1926	—	—	—	—	—	—
	China/Bohlin 1926	123.42	54.57	71.6	76.21	44.21	58.00
<i>P. quadricornis</i>	Samos	135	58.6	81.8	71.63	43.40	60.59
<i>P. asiaticus</i>	Ortok	133	59	78	75.64	44.36	58.64
	Kalmakpai	—	—	—	—	—	—
	Pavlodar	—	—	—	—	—	—
<i>P. hoffstetteri</i>	Sinap	151.3	61.6	93.2	66.09	40.71	61.59
<i>P. pavlowae</i>	Grebeniki	127	54	72	75	42.51	56.69
	Blagodarnenskaya	127	55	73	75.34	43.30	57.48
<i>P. moldavicus</i>	Raspopeny	—	—	—	—	—	—
	Korsakova Assembly	—	—	—	—	—	—
<i>P. borissiaki</i>	Eldari	—	—	77	—	—	—
<i>P. expectans</i>	Sevastopol	138	62	80	77.5	44.92	57.97
	Varnitsa	—	—	—	—	—	—
<i>P. aff. berislavicus</i>	NKT	143.75	63	85	74.11	43.90	59.23
<i>P. coelophrys</i>	PNT	138.45	59.5	82.14	72.43	42.97	59.32
<i>P. cf. coelophrys</i>	RPI	149.82	65.74	89.74	74.15	44	59.33

APPENDIX 4. — Continuation.

	Site	R-GL	R-Bp	Mc-GL	Mc-Bdia	Mc-R.I.	T-Bd	T-Dd
<i>P. rouenii</i>	Pikermi	477.4	71.5	420.15	34.8	8.28	55	40.65
	Cimişlia	492.5	83.5	425.5	37	8.69	66.6	51
	Taraclia	488	86.5	—	—	—	70	52
	Kerassia	—	—	445	38.4	8.62	65.3	50.4
	Ditiko	—	—	—	—	—	—	—
	Cioburciu	—	—	—	—	—	—	—
	Samos	483.75	79.58	422	37.85	8.96	62.25	53.5
	Todorovo	—	—	402	38	9.45	—	—
	Kemiklitepe	453	79.25	405	33.5	8.27	—	—
	Novo-Elisabetovka	—	—	—	—	—	—	—
	Strumyani	—	—	—	—	—	—	—
	Hadjidimovo	—	—	—	—	—	—	—
	Akkasdagli	88.2	—	34.4	—	67.5	51.7	—
	Thermopigi	568	83.21	475.79	34.5	7.25	—	—
	Kryopigi	500.7	—	—	—	7.1	—	—
	Perivolaki	565	84.65	—	—	—	—	—
<i>P. coelophrys</i>	Maragha	485	100.5	369.8	41	11.08	72.5	57.5
	Samos	—	—	—	—	—	—	—
<i>P. cf. coelophrys</i>	China	—	—	—	—	—	—	—
<i>P. berislavicus</i>	Berislav	450	91	—	—	—	75.6	48.16
<i>P. microdon</i>	Kansu/Bohlin 1926	—	—	—	—	—	—	—
	Shansi/1926	—	—	—	—	—	—	—
	China/Bohlin 1926	396	63	368	35	9.51	55	—
<i>P. quadricornis</i>	Samos	—	—	—	—	—	—	—
<i>P. asiaticus</i>	Ortok	501	—	—	—	—	76.5	54.5
	Kalmakpai (Kazakhstan)	—	—	—	—	—	—	—
	Pavlodar	—	—	—	—	—	—	—
<i>P. hoffstetteri</i>	Sinap	—	—	—	—	—	—	—
<i>P. pavlowae</i>	Grebeniki	—	—	—	—	—	—	—
	Blagodarnenskaya	—	—	—	—	—	—	—
<i>P. moldavicus</i>	Raspopeny	552.5	114	—	45	—	87	66
	Korsakova Assembly	107.5	—	—	—	—	—	—
<i>P. borissiaki</i>	Eldari	539	106	407.5	39	9.57	70	58
<i>P. expectans</i>	Sevastopol	108.5	400	47	11.75	83	62	—
	Varnitsa	—	—	—	—	—	—	—
<i>P. aff. berislavicus</i>	NKT	464.25	81.196	389	35.79	9.2	72.67	52.09
<i>P. coelophrys</i>	PNT	—	—	—	—	—	—	—
<i>P. cf. coelophrys</i>	RPI	—	—	—	—	—	—	—

APPENDIX 4. — Continuation.

	Site	Mt-Bp	Mt-Dp	Mt-Bdia	Mt-Ddia	Mt-GL	Mt-R.I.
<i>P. rouenii</i>	Pikermi	48.33	48.5	32	31.5	426	7.51
	Cimişlia	55.5	53	33.5	39.5	425	7.88
	Taraclia	56.67	57.67	33.25	34	436.75	7.61
	Kerassia	55.95	49.9	36.5	41.2	—	—
	Ditiko	—	—	—	—	—	—
	Cioburciu	—	—	—	—	—	—
	Samos	53.1	51.8	31.77	35.35	445	7.37
	Todorovo	—	—	—	—	—	—
	Kemiklitepe	—	—	—	—	—	—
	Novo-Elisabetovka	56.67	57.67	34	34	—	—
	Strumyani	—	—	—	—	—	—
	Hadjidimovo	—	—	—	—	—	—
	Akkasdagli	—	—	—	—	—	—
	Thermopigi	47.14	53.1	31.37	35.26	497.79	6.13
<i>P. coelophrys</i>	Kryopigi	51	49.325	31.4	37.35	467.5	6.71
	Perivolaki	49.45	53.05	29	33.3	447	6.48
<i>P. cf. coelophrys</i>	Maragha	56.74	54.92	36.47	38.46	382.55	9.53
	Samos	54	—	35	—	377	9.28
<i>P. cf. coelophrys</i>	China	—	—	—	—	—	—
<i>P. berislavicus</i>	Berislav	55.96	55.62	30.5	35.67	400.5	7.61
<i>P. microdon</i>	Kansu/ Bohlin 1926	—	—	—	—	—	—
	Shansi/1926	—	—	—	—	—	—
	China/Bohlin 1926	48	54	30	38	401	7.48
<i>P. quadricornis</i>	Samos	62	61	42	42	404	10.39
<i>P. asiaticus</i>	Ortok	—	—	—	—	—	—
	Kalmakpai	—	—	—	453	—	—
	Pavlodar	63	65	38	45	445	8.53
		59	61	37	42	—	—
<i>P. hofstetteri</i>	Sinap	—	—	—	—	—	—
<i>P. pavlowae</i>	Grebeniki	—	—	—	—	—	—
	Blagodarnenskaya	—	—	—	—	—	—
<i>P. moldavicus</i>	Raspopeny	70	66.67	46.5	44	—	—
	Korsakova Assembly	—	—	—	—	—	—
<i>P. borissiaki</i>	Eldari	64.5	65	44.5	—	—	—
<i>P. expectans</i>	Sevastopol	64.5	—	40.5	50.5	—	—
	Varnitsa	—	—	46	—	447	10.29
<i>P. aff. berislavicus</i>	NKT	58.17	50.85	37.33	31.7	412	9.06
<i>P. coelophrys</i>	PNT	61.68	59.16	38.93	35	—	—
<i>P. cf. coelophrys</i>	RPI	—	—	—	—	—	—

APPENDIX 4. — Continuation.

	Site	A-GLI	A-Bd	P3 morphology	P4 lingual groove	P4 stylids	P4 shape
<i>P. rouenii</i>	Pikermi	71	44	1	0	1	2
	Cimişlia	73	45	–	–	–	–
	Taraclia	73	45.89	–	–	–	–
	Kerassia	–	–	1	–	–	–
	Ditiko	–	–	1	1	0	2
	Cioburciu	–	–	–	–	–	–
	Samos	75	47.1	1	0	0	2
	Todorovo	–	–	–	–	–	–
	Kemiklitepe	–	–	–	–	–	–
	Novo-Elisabetovka	45	–	–	–	–	–
	Strumyani	–	–	–	–	–	–
	Hadjidimovo	–	–	–	–	–	–
	Akkasdagli	78.1	49.75	–	0	0	2
	Thermopigi	–	–	0	0	1	–
	Kryopigi	–	–	–	–	–	–
	Perivolaki	–	–	–	–	–	–
<i>P. coelophrys</i>	Maragha	86.81	55.20	0	0	0	0
	Samos	–	–	–	–	–	–
<i>P. cf. coelophrys</i>	China	75	47	1	0	0	2
<i>P. berislavicus</i>	Berislav	73.5	49	1	0	0	1
<i>P. microdon</i>	Kansu/Bohlin 1926	–	–	0	0	2	–
	Shansi/1926	–	–	0	0	2	–
	China/Bohlin 1926	70	39.5	–	0	0	2
<i>P. quadricornis</i>	Samos	–	–	0	1	0	2
<i>P. asiaticus</i>	Ortok	73	51	1	0	0	2
	Kalmakpai	–	–	–	–	–	–
	Pavlodar	83.5	52.5	–	–	–	–
<i>P. hofstetteri</i>	Sinap	87.5	51.58	0	0	0	1
<i>P. pavlowae</i>	Grebeniki	82	53	–	1	0	2
<i>P. moldavicus</i>	Raspopeny	89.5	59.5	1	0	0	0
	Korsakova Assembly	–	–	–	–	–	–
<i>P. borissiaki</i>	Eldari	–	56	0	0	0	2
<i>P. expectans</i>	Sevastopol	85	54	0	–	–	–
	Varnitsa	–	–	0	–	–	–
<i>P. aff. berislavicus</i>	NKT	78.35	49.33	–	0	1	2
<i>P. coelophrys</i>	PNT	–	–	0	0	1	–
<i>P. cf. coelophrys</i>	RPI	–	–	–	0	0	2