Kretzoiarctos gen. nov., the Oldest Member of the Giant Panda Clade

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Abstract

The phylogenetic position of the giant panda, Ailuropoda melanoleuca (Carnivora: Ursidae: Ailuropodinae), has been one of the most hotly debated topics by mammalian biologists and paleontologists during the last century. Based on molecular data, it is currently recognized as a true ursid, sister-taxon of the remaining extant bears, from which it would have diverged by the Early Miocene. However, from a paleobiogeographic and chronological perspective, the origin of the giant panda lineage has remained elusive due to the scarcity of the available Miocene fossil record. Until recently, the genus Ailurarctos from the Late Miocene of China (ca. 8–7 mya) was recognized as the oldest undoubted member of the Ailuropodinae, suggesting that the panda lineage might have originated from an Ursavus ancestor. The role of the purported ailuropodine Agriarctos, from the Miocene of Europe, in the origins of this clade has been generally dismissed due to the paucity of the available material. Here, we describe a new ailuropodine genus, Kretzoiarctos gen. nov., based on remains from two Middle Miocene (ca. 12–11 Ma) Spanish localities. A cladistic analysis of fossil and extant members of the Ursidae confirms the inclusion of the new genus into the Ailuropodinae. Moreover, Kretzoiarctos precedes in time the previously-known, Late Miocene members of the giant panda clade from Eurasia (Agriarctos and Ailuropoda). The former can be therefore considered the oldest recorded member of the giant panda lineage, which has significant implications for understanding the origins of this clade from a paleobiogeographic viewpoint.


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Introduction

The Fossil Record of the Giant Panda Lineage

The extant giant panda from central Asia, Ailuropoda melanoleuca [1], differs from other living Ursidae by the presence of several craniodental adaptations to durophagy (i.e., feeding tough plant material, mainly bamboo) [1,2,3,4]. These adaptations are already present, to a large extent, in the Plio-Pleistocene relatives of A. melanoleuca (Ailuropoda microta and Ailuropoda wulanshanensis) [2,4], which displayed a larger distribution from northern China to Southeast Asia [3,6,7], than the living giant panda. Despite longstanding disputes about the phylogenetic position of Ailuropoda, especially due to dietary-driven convergence in the dentition with the lesser panda (Ailurus fulgens) [2,3], it is currently well-established on molecular grounds that the former corresponds to the living sister-taxon of other extant bears [8,9,10,11,12,13], being classified in its own subfamily (Ailuropodinae) [11,13] or tribe (Ailuropodini) [14]. Molecular data have estimated the divergence time between ailuropodines and other bears to correspond to the time between ailuropodines and other bears to correspond to the Early Miocene (ca. 22 mya) [8], 19 mya [12] or 18 mya [2]). However, from a paleontological perspective, ailuropodine origins are still largely uncertain, due to the paucity of the available Miocene record [14]. It is generally considered that Ailuropoda descended from the Asian, Late Miocene ursid Ailurolagus [5,7,14]. The latter genus is first recorded by Ai. yuanmouensis from the Chinese locality of Yuanmou (8.2-7.2 mya) [15], being subsequently recorded by Ai. yungchengensis from Lufeng (6.9-5.8 mya) [5,6,7,14,15]. Ailurolagus has been considered to be descended from an unidentified Miocene species of Ursavus [4,5,6,7,14], although no formal cladistic analysis had been thus far provided to substantiate such phylogenetic hypothesis.

Besides Ailurolagus, the extinct genus Ailurictus, from the Miocene of Europe [5,16,17,18], has been also attributed to the Ailuropodinae. Until recently, the scarcity of available Ailurictus material precluded a secure assessment of its phylogenetic affinities with the Late Miocene Ailurolagus, which already displays incipient adaptations to durophagy, being customarily considered the oldest undoubted member of the giant panda lineage [2,4,5,7,14]. The type species of Ailurictus, Ag. gaulisi, was originally based on a mandibular fragment with p3-m2 [16] from the Turolian (MN12, Late Miocene) locality Hatan (Hungary) [19]. Ag. vighi was also created by Kretzoi (1942), after an m1 from the Hungarian locality
of Rozsaszentmárton. With no other material for comparison, provisionally we consider these two species as valid, until a more detailed study of these fossils is carried out. Although some authors attributed the holotype of *A. gaali* to *Indarctos cf. sibiri* [19], more recently the validity of the former genus and species have been supported [20], being characterized by a strong development of the distal cusps of the premolars (usually poorly developed or absent in most of the Ursidae) and by the mesial position of the m1 metaconid [20]. *Agrictos* was first transferred to the Ailuropodinae several decades ago [17,18], by further including material from the Late Miocene localities of Soblay (MN10, France; [21]) and Wissbergh (= Gau-Weinheim; MN9, Germany), previously attributed to *Ursus arctos* [22,23,24]. The remains from Soblay differ from *Ursus* by the lengthening of the upper carnassial, due to the presence of a parastylar [22,23]. They can be therefore attributed to the genus *Agrictos* [17,20], even if indirectly (i.e., *Agrictos cf. gaali*) [18], since a direct comparison with the *Ag. gaali* holotype is not possible due to the lack of upper dentition in the latter.

Most recently, an older, new species of this genus, *Agrictos beatrix*, was described on the basis of dental remains from the late Aragonian (MN9, Middle Miocene) locality of Nombrevilla 2 (NOM2, Calatayud-Daroca Basin, Spain) [20], formerly attributed to *Ursus primaeus* [25]. Given the lack of lower dentition, however, a direct comparison with *Ag. gaali* from the type locality was not feasible. On the basis of a new mandible of the same species, recovered in a similarly-aged (MN6) locality from the Abocador de Can Mata (ACM) local stratigraphic series (Valles-Penedès Basin, Spain), here we show that “*Agrictos* beatrix” is distinct enough as to be attributed to a different genus, *Kretzoiarctos* gen. nov. This new genus represents the oldest and most basal member of the ursid clade currently represented by *Ailuropoda*, thus being of utmost significance for understanding the origin of the giant panda lineage from both a chronological and paleobiogeographic perspectives.

Age and Geological Background

The local stratigraphic series of ACM (els Hostalets de Pireloa, Catalonia, Spain), situated in the Valles-Penedès Basin (NE Iberian Peninsula), is a 250 m-thick stratigraphic composite succession including more than 200 fossil vertebrate localities and spanning about 1 myr (from ca. 12.5 to 11.5 mya; late Aragonian, Middle Miocene) [26,27,28,29]. The new material described in this paper (IPS46473) corresponds to an isolated find from sector ACM/C6-Cami, in a stratigraphic horizon situated 2 m above the formally-defined locality ACM/C6-A1. Based on litho- and magnetostratigraphic correlation [27,30], both ACM/C6-A1 and the layer where IPS46473 was found are correlated to subchron C5r.2n, with an interpolated estimated age of 11.6 mya. Although no associated small mammal remains are available for IPS46473, its age indicates that this find belongs to the *Megacricetodon ibericus* + *Democricetodon crassifons* concurrent range local biozone [29,31], which can be correlated to the MN6 sensu Mein and Ginsburg [32]. The locality of Nombrevilla 2 (NV2), situated in the Calatayud-Daroca Basin, can be correlated to the same biozone, spanning from ca. 11.8-11.2 mya [29,31], given the presence of *M. ibericus* and *D. crassifons* together with the lack of the hinnariine equid *Hipparitherium* [25].

Systematic Paleontology

Order Carnivora Bowdich, 1821; suborder Caniformia Kretzoi, 1942; infraorder Arctoidea Flower, 1869; parvorder Ursida Tedford, 1976; superfamilies Ursoidae Fischer von Waldheim, 1814; family Ursidae Fischer von Waldheim, 1814; subfamily Ailuropodinae Grevē, 1894; tribe Ailuropodini Grevē, 1894; *Kretzoiarctos* gen. nov.

Etymology: Dedicated to the paleontologist Miklos Kretzoi and from the Greek ‘*arctos*’ (bear).

Type species: *Kretzoiarctos beatrix* (Abella et al., 2011) comb. nov. [20].

Diagnosis: as for the type species.

Differential Diagnosis

*Kretzoiarctos* shares many morphological characters with the other middle and late Miocene European (*Ursus*, *Indarctos* and *Agrictos*) and Asian (*Ailuropus*) ursid genera, from which it is distinguished by a unique combination of features. *Kretzoiarctos* differs from *Ursus* in the relative development of the upper premolars and lower molars. Thus, the upper carnassial in *Ursus* is triangular and always lacks a parastylar, whereas in *Kretzoiarctos* it shows a well-developed parastylar (although not yet separated from the paracone). In turn, the lower carnassial in *Ursus* has a well-developed sectorial blade in the trigonid, whereas in *Kretzoiarctos* the m1 displays blunt and low cusps. Moreover, the m2 in *Ursus* is relatively shorter than in *Kretzoiarctos* because of the less-developed talonid. *Kretzoiarctos* differs from *Agriarctos* in the following features: smaller dental size; less developed lower premolars without strong accessory cusps; P4 with a less developed parastylar, a less complex and more mesially-situated protocone, much less developed basal labial cingulum, more labiolingually-compressed labial cusps (parastyle, paracone and metastyle), and P4 slightly longer than the M1 (instead of being similar in length). Finally, *Kretzoiarctos* can be easily distinguished from *Indarctos* by the smaller size of the former. *Indarctos sibiri* is the smallest species of this genus and the only one that could be mistaken for *Kretzoiarctos*, although they can be distinguished because *I. sibiri* has no trace of parastylar and no accessory cusps in the premolars.

*Kretzoiarctos beatrix* (Abella et al., 2011) comb. nov

Synonyms: *Ursus arctos* (in ref. [33], p. 78); *Ursus primaeus* (in ref. [25], p. 31); *Agrictos beatrix* (in ref. [20], p. 188).

Holotype: left P4, NV-2-42 (Figure 1; 3a–c) from NV II.

Paratype: right M1, NV-2-40 (Figure 1; 4a–c) from NV II.

The hypodigm also includes the new material described here: partial right mandible with c1–m2 and associated P4, IPS46473 (Figure 1; 1–2 and Figure 2) from ACM/C6-Cami.

Type locality: Nombrevilla 2 (NV II; Calatayud-Daroca Basin, Spain).

Other localities: ACM/C6-Cami (Valles-Penedès Basin, Spain).

Chronological range: the type locality has an age of ca. 11.8-11.2 mya [34], whereas ACM/C6-Cami has an estimated age of 11.6 mya, both localities thus being correlated to the MN8 sensu Mein and Ginsburg [32] (late Aragonian, Middle Miocene).

Emended Diagnosis

Small-sized ailuropodine species. P4 with a well-developed protocone situated opposite to the paracone, and parastry of moderate size but well-individualized from the protocone. M1 with a highly-developed metastyle and lingual cingulum poorly-differentiated from the protocone and hypocone [20]. Robust mandibular corpus, deepest under the m1 and m2. Low-crowned and curved lower canine. Lower premolars (p2–p4) with a single, duniform main cusp, and reduced mesial and distal accessory cusps, not separated by any diastema. Long and low-crowned m1, with the metaconid and protoconid of similar height, long and shallow talonid basin, and no cusp at the paraconid-hypoconid valley. Relatively long m2 with well-developed trigonid and talonid basins.
Description of the New Material

IPS46473 corresponds to a partial right mandible that preserves the canine and the lower cheek teeth (see measurements in Table 1). The mandibular corpus is short, displays a rounded ventral outline and is taller under the m1–m2 than at the symphyseal region. Despite some damage, a relatively tall and verticalized coronoid process may be reconstructed. The lower canine, which is slightly displaced out from its alveolus, is low-crowned and curved. The premolars (p2–p4) have a single main cusp that displays a duniform shape, as well as poorly-developed mesial and distal accessory cusps. There is no diastema between the lower premolars. The m1 is long and low-crowned; the metaconid and the protoconid are similar in height; the talonid basin is long and shallow; no cusp can be observed within the paraconid-hypoconid valley. The m2 is long, and both the trigonid and talonid basins are shallow but wide. The associated P4 is quite damaged, and only the paracone and metastyle can be observed; however, it is possible to ascertain that it was a long upper carnassial with a well-developed labial cingulum; the parastyle is preserved but it cannot be clearly observed due to breakage; both the paracone and metastyle are low and wide.

Results

Nomenclatural Statement

An LSID number was obtained for the new taxon (Genus Kretzoiarctos):
urn:lsid:zoobank.org:act:96C5EE3D-3C7B-4D5B-80FD-95C242753DFA.

Cladistic Analysis

A cladistic analysis based on a morphologic data matrix for living and fossil ursids recovered a single most parsimonious tree of 157 steps (Figure 3). Besides the outgroup (the canid Canis lupus), the basalmost taxon is Zaranocyon daamsi, a representative of the Hemicyonidae–stem Ursidae not included within the Ursidae [35]. The analysis therefore recovers the monophyly of the Ursidae, represented by three successive members of its stem lineage (Ballusia elmensis, Ursavus primaevus and Ursavus brevirhinus, the latter suggesting that the genus Ursavus is paraphyletic), and two major clades: the Ailuropodinae, including the extant giant panda (Ailuropoda); and the Ursinae + Tremarctinae, including all the extant ursids except Ailuropoda. Among the latter, the extant Tremarctos ornatus appears as the basal most species, followed by Melursus ursinus. Among the Ailuropodinae, two distinct subclades can be distinguished: that composed by Indarctos species (I. vireti, I. arctoides and I. panghrensis), which are here included in a distinct tribe, Indarctini tribe nov. (type genus Indarctos); and the one including the remaining ailuropodines (tribe Ailuropodini). The Indarctini appear as the sister-taxon of the Ailuropodini, comprising the European genera Agriarctos and Kretzoiarctos, as well as the Asian Ailuropoda and Ailurarctos. The three extinct ailuropodin genera are recovered as a monophyletic clade, sister taxon of the extant Ailuropoda.

A bootstrap analysis (Figure 3) shows that most of the clades recovered by the most parsimonious tree are relatively well supported (i.e., bootstrap values higher than 50%). The monophyly of the Ursidae total group (bootstrap value 79) and of the Ursidae crown-group (bootstrap value 66) are well supported, whereas in contrast the successive stem position of Ursavus species is not supported. Among crown ursids, the dichotomy between Ailuropodinae (bootstrap value 82) and the clade composed by Tremarctinae + Ursinae (bootstrap value 82) is also well supported. Within the latter, only the position of the ursines Helarctos malayanus and Ursus thibetanus are not supported by the 50% majority rule bootstrap analysis (see Discussion). Within the Ailuropodinae, no clade is supported by the 50% majority rule tree recovered by the bootstrap analysis. The lack of resolution among this subfamily may be attributed to the retention of many plesiomorphies shared with both the successive stem ursids Ursavus...
and Ballusia, as well as with the other most primitive members of the Ailuropodinae (Indarctos, Kretzoiarctos and Agriarctos). However, most importantly, the inclusion of Kretzoiarctos and the other putative extinct ailuropodines in this subfamily is well supported not only by the most parsimonious cladogram but also by the bootstrap analysis. Accordingly, the new genus Kretzoiarctos emerges as the oldest member of the panda lineage (either the Ailuropodini and the Ailuropodinae) thus far recorded, thus enabling to track the fossil record of this subfamily of bears back to the MN8 (11.8-11.2 mya).

**Discussion**

The results of our cladistic analysis of the Ursoidea mostly agree well with molecular analyses performed on extant ursids [13,36,37], especially regarding the fact that the giant panda appears as the sister-taxon of all remaining members of this family. There are a few particular points that should be taken into account. The first is the position of Melursus ursinus, considered a member of the Ursinae, but not showing a clear position within this clade [13]. M. ursinus displays a very autapomorphic craniodental morphology adapted to myrmecophagy [38], characterized by reduced teeth and even lacking several incisors. Accordingly, the basal position of Melursus recovered by our analysis–restricted to dental, cranial and mandibular features–even though it does not differ much from the genetic-based analysis–could be interpreted as an artifact, reflecting its very autapomorphic dietary complex relative to the remaining Ursinae. A similar anomalous result is obtained by our analysis regarding the lack of support of the Ursus clade, whose monophyly is supported by molecular studies [13]. Like above, the divergent dietary adaptations displayed among the ursine bears [39] could be the cause of this ambiguity.

With regard to Ailuropodinae, our analysis indicates that this subfamily constitutes a monophyletic clade, sister-taxon to that including the remaining ursid subfamilies (Tremarctinae and Ursinae), in agreement with molecular analyses [13]. According to our results, the Ailuropodinae would be characterized by the following synapomorphies: a tall coronoid process; a high articular process; and alisphenoid canal present. Most of the remaining characteristics of this group correspond to primitive features that have been subsequently lost in both the Tremarctinae and the Ursinae, including among others: the development of the premolars; the well-developed carnassials; the wide molars; and the curved tooth row. These primitive characters are not informative for distinguishing ailuropodines from stem ursids, although they can be used to distinguish the former from the two other, more derived subfamilies of crown ursids. Along with these features, in the Ailuropodinae there is a tendency towards an increased dental complexity, as reflected by the presence of a well-developed parastyle and a complex protocone, which is a synapomorphy shared by all ailuropodines except the primitive Indarctos vireti. A list of all the apomorphies is given in Table S2.

The most parsimonious tree delivered by our cladistic analysis recovers two distinct ailuropodine subclades, which are here distinguished at the tribe level (Ailuropodini and Indarctini). Alternatively, these subclades could be distinguished at the subfamily rank. However, it should be taken into account that the monophyly of these tribes, unlike that of the subfamily Ailuropodinae, is not supported by the bootstrap analysis, so that merely distinguishing them at the tribe level is a more conservative option. Additional studies based on more complete material would be required in order to confirm their monophyly. If confirmed, or in case one of them was shown to occupy a more basal position than the other within the Ursidae, then it might be preferable to separate them at the subfamily level. The Indarctini includes the species of the genus Indarctos [40,41,42], which was widely distributed though Northern Africa, Eurasia and North and Central America during the Late Miocene (MN9-MN13; [40,43]).

**Table 1.** Dental measurements (in mm) of Kretzoiarctos beatrix from ACM/C6-Camí (this study) and Nombrevilla 2 (from ref. 20).

<table>
<thead>
<tr>
<th>Catalogue No.</th>
<th>Tooth</th>
<th>Length</th>
<th>Width</th>
</tr>
</thead>
<tbody>
<tr>
<td>IPS46473</td>
<td>Right lower canine</td>
<td>14.73</td>
<td>8.84</td>
</tr>
<tr>
<td>IPS46473</td>
<td>Right p2</td>
<td>7.67</td>
<td>4.83</td>
</tr>
<tr>
<td>IPS46473</td>
<td>Right p3</td>
<td>5.33</td>
<td>8.87</td>
</tr>
<tr>
<td>IPS46473</td>
<td>Right p4</td>
<td>–</td>
<td>6.64</td>
</tr>
<tr>
<td>IPS46473</td>
<td>Right m1</td>
<td>22.64</td>
<td>10.80</td>
</tr>
<tr>
<td>IPS46473</td>
<td>Right m2</td>
<td>17.74</td>
<td>11.28</td>
</tr>
<tr>
<td>IPS46473</td>
<td>Right m3</td>
<td>–</td>
<td>9.73</td>
</tr>
<tr>
<td>NV-2-42</td>
<td>Left P4</td>
<td>18.5</td>
<td>13.0</td>
</tr>
<tr>
<td>NV-2-40</td>
<td>Right M1</td>
<td>17.2</td>
<td>15.4</td>
</tr>
</tbody>
</table>

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These species include *I. vireti* (Iberian Peninsula, MN9) and *I. arctoides* (Europe MN10-11), with a relatively limited temporal range and geographic distribution, as well as *I. punjabiensis* (MN10-13), with a Holarctic distribution, and the endemic insular *I. laurillardi* from Baccinello (MN12), whose taxonomic status is yet to be determined. According to our results, the Indarctini still share several plesiomorphic traits with the Ailuropodini (non-reduced carnassials, all premolars present, wide molars, etc.), whereas some of the common characters, such as the developed pterygoideus muscles, the high coronoid process and the development of the zygomaticomandibularis muscle against the reduction of the masseteris, are derived characters present in the ursids with less meat in their diet [44]. The Ailuropodini, comprising the extant giant panda and the remaining fossil Ailuropodinae—including *Kretzoiarctos*—can be distinguished from the Indarctini by a set of derived dental features (elongated P4, presence of a labial parastyle in the P4, development distal and mesial accessory cusps in the premolars), which we interpret as adaptations towards a more herbivorous diet [1, 2, 3, 4]. With a late Miocene age, *Kretzoiarctos* represents the oldest known member not only of the tribe Ailuropodini, but also of the whole subfamily Ailuropodinae (*Ailuropodinae* + *Indarctini*), substantially preceding in time the other taxa that had been previously attributed to this group. Given that *Kretzoiarctos* is only known from the Iberian Peninsula (Calatayud-Daroca and Valles-Penedes basins), a Western European origin of the giant panda lineage (Ailuropodinae) is now tentatively supported by the results of this paper. It should be taken into account, however, that the fossil record of this group is still too scarce and fragmentary, as evidenced by the various ghost lineages that must be inferred based on the Early Miocene divergence times for ailuropodines suggested by molecular data. The fossil remains of *Kretzoiarctos* reported here, however, at least conclusively document the occurrence of ailuropodines by the Middle Miocene.

**Conclusions**

A new genus of extinct ursid belonging to the giant panda lineage, *Kretzoiarctos* gen. nov. (*Ursidae: Ailuropodinae: Ailuropodini*), is here described on the basis of new fossil remains from the Spanish site ACM/C6-Cami (Valles-Penedes Basin). This new material allows a more precise taxonomic approach of the type material from the also Spanish locality of Nombrevilla 2 (Calatayud-Daroca Basin), previously attributed to the Late Miocene genus *Agriarctos*. With a late Middle Miocene age, *Kretzoiarctos* is the oldest known member of the tribe Ailuropodini, but also of the whole subfamily Ailuropodinae (*Ailuropodini* + *Indarctini*), substantially preceding in time the other taxa that had been previously attributed to this group. Given that *Kretzoiarctos* is only known from the Iberian Peninsula (Calatayud-Daroca and Valles-Penedes basins), a Western European origin of the giant panda lineage (*Ailuropodinae*) is now tentatively supported by the results of this paper. It should be taken into account, however, that the fossil record of this group is still too scarce and fragmentary, as evidenced by the various ghost lineages that must be inferred based on the Early Miocene divergence times for ailuropodines suggested by molecular data. The fossil remains of *Kretzoiarctos* reported here, however, at least conclusively document the occurrence of ailuropodines by the Middle Miocene.
Miocene of Eurasia, with *Boulaia* and *Ursinus* displaying a successive basal position with regard to crown ursids as a whole.

**Materials and Methods**

**Cladistic Analysis**

The data matrix of craniodental features employed in the cladistic analysis, including 82 characters and 19 taxa (see Table S1), was coded on the basis of original specimens of osteological and fossil material, casts, and published figures and descriptions. For *Canis lupus*, *Ursus arctos*, *Ursus americanus*, *Tremarctos ornatus*, *Ailuropoda melanoleuca*, *Ursus brennus*, *Indarctos everti*, *I. arctoides* and *J. pongebensis*, we had direct access to skulls and mandibles. For *Ursus thibetanus* and *Helarctos malayanus*, we relied on casts of mandibles and skulls. For the remaining species, we used either photographs or images from scientific papers. The character description can be consulted in the Text S1. The matrix was generated using MacClade 4.08a OS X, and was analyzed using PAUP* [Version 4.0b10 for Macintosh [45]. A maximum-parsimony analysis was performed using the branch-and-bound method, with *Canis lupus* as the outgroup. Even though *C. lupus* is not a member of the Ursidae, its cranial, mandibular and dental morphologies are supposed to be similar to the ancestor of the Arctoidea, and therefore a quite accurate choice as an outgroup for this analysis. In order to test clade robusticity, a bootstrap analysis with 1,000 replicates was performed using the branch-and-bound search option.

**Studied Material**

IPS46473 is a nearly complete right mandible with canine and p2-m3 series (Figure 1; 1a-c and Figure 2) associated to a broken P4 (Figure 1; 2a–b) from ACM/C6-Cami. These specimens are housed at the Instituto Català de Paleontologia Miquel Crusafont (ICP; Sabadell, Catalonia, Spain). The material from Nombrevilla (NON) is researcher in formation in the CSIC program JAE-PRE_CP2011. We thank the Preparation Division of the ICP for the preparation of the A.C.M specimen, and Marta Palmero for her splendid drawings. Israel Sánchez gave us crucial advice regarding the cladistics analysis. We also thank Borja Figueirido for the photographs of *U. maritimus* and *M. ursinus*, and Francisco Pastor for the loan of the skeleton of *T. ornatus*. A.V. is researcher in formation in the CSIC program JAE-PRE_CP2011.

**Nomenclatural Statement**

The electronic edition of this article conforms to the requirements of the amended International Code of Zoological Nomenclature, and hence the new names contained herein are available under that Code from the electronic edition of this article. This published work and the nomenclatural acts it contains have been registered in ZooBank, the online registration system for the ICZN. The ZooBank LSIDs (Life Science Identifiers) can be resolved and the associated information viewed through any standard web browser by appending the LSID to the prefix “http://zoobank.org/”. The LSID for this publication is: urn:lsid:zoobank.org:pub:B572EF80-998C-45D4-8364-14C0C7F299D9. The electronic edition of this work was published in a journal with an ISSN, and has been archived and is available from the following digital repositories: PubMed Central, LOCKSS.

**Supporting Information**

**Table S1** Data matrix of craniodental features employed in the cladistic analysis, including 82 characters and 19 taxa. (NON)

**Table S2** List of the apomorphies found in the cladistic analysis. Note: the changes of states are shown for each taxon. Note: Character number start in 0. Therefore Character 1 in the matrix would be character 0 in this list. (PDF)

**Text S1** Character description. (PDF)

**Acknowledgments**

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**Author Contributions**

Conceived and designed the experiments: JA PM JM. Performed the experiments: JA JM. Analyzed the data: JA DMA JMR AV. Wrote the paper: JA DMA JMR AV. Contributed reagents/materials/analysis tools: JA DMA JMR AV. Wrote the paper: JA DMA.

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