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A new and primitive species of *Protophiomys* (Rodentia, Hystricognathi) from the late middle Eocene of Djebel el Kébar, Central Tunisia
A new and primitive species of *Protophiomys* (Rodentia, Hystricognathi) from the late middle Eocene of Djebel el Kébar, Central Tunisia

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Abstract: Based on fossil discoveries and phylogenetic studies, an Eocene Asian origin for hystricognathous rodents and anthropoid primates has gained strong support in recent years. The two groups then invaded both Africa and South America, which promoted their evolutionary success. However, the fossil record has so far failed to constrain the nature and precise timing of these pivotal dispersal events. In Africa, given the apparent absence of hystricognaths and anthropoids in early to early middle Eocene localities, it is suggested that these mammal groups dispersed from Asia to Africa sometime during the middle Eocene. In this paper, we report the discovery of several isolated teeth of a rodent from a new vertebrate locality situated in central Tunisia (Djebel el Kébar, KEB-1), dating from the late middle Eocene (Bartonian, ~39.5 Myr). These fossils document a diminutive new species of *Protophiomys (P. tunisiensis nov. sp.)*, a basal genus of hystricognathous rodents which is well known from several North African mammalian-bearing localities of the end of the Eocene. The teeth of *P. tunisiensis* display a suite of anatomical details comparable with those observed in the other species of the genus, but with a lesser degree of development. Such an apparent primitive evolutionary stage is corroborated by the greater antiquity of this Tunisian species. *P. tunisiensis* nov. sp. is so far the most ancient representative of hystricognaths in Africa. However, it can be expected that hystricognaths were already present on that landmass given the new data on early caviomorphs recently reported from South America (at ~41 Myr). The arrival of hystricognaths in Africa from South Asia certainly predates the depositional period of the Kébar sediments, but perhaps not by much time.

Keywords: North Africa, Bartonian, Dental morphology, Paleobiogeography, Adaptive radiation

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INTRODUCTION

Hystricognaths are a natural group of rodents (infra-order Hystricognathi Tullberg, 1899) that are today represented by the Afro-Asian porcupines, African cane-, mole- and dassie-rats, and the South American guinea pigs, chinchillas, capybaras, pacas, agoutis, etc. From a paleontological perspective, hystricognaths are so far absent (unknown) from the earliest Tertiary fossil record at a global scale. Their earliest occurrences date only from the late middle or early late Eocene in both Africa and South America (Jaeger et al., 1985, 2010a; Sallam et al., 2009; Antoine et al., 2012). Such a record either indicates the existence of a significant Eocene (Lutetian) gap in the hystricognath record, or suggests that this group achieved a rapid widespread distribution in the Old and New Worlds just after its emergence. The record of the earliest hystricognaths was long exclusive to Africa (e.g., Osborn, 1908; Wood, 1968; Jaeger et al., 1985). However, over the past few decades, fossil discoveries in South Asia have demonstrated that this tropical province was also a major centre of an early hystricognathous rodent radiation (the “baluchimyine” radiation), and most likely the ancestral homeland of the Hystricognathi clade (Hussain et al., 1978; Marivaux et al., 2002, 2004; Dawson et al., 2006; Sallam et al., 2009, 2012; Barbière & Marivaux, in press). From a historical biogeographic perspective, in this context of an Asian origin, it is worth noting that if ancestral Asian hystricognaths dispersed across the Tethys Sea to colonize Africa sometime during the middle Eocene, they were also able to continue their pattern of intercontinental dispersal across the South Atlantic to invade South America as well (e.g., Antoine et al., 2012). Interestingly, the same seems to be true for anthropoid primates (stem Afro-Asian anthropoids and the origin of platyrhines; e.g., Antoine et al., 2012; Chaimanee et al., 2012; Seiffert, 2012). However, critical factors such as the mode and precise timing of these Asio-African and Afro-South-American dispersals, and the pathways by which these dispersals occurred remain entirely unknown.

In the framework of our paleontological program in the early Tertiary of North Africa, since 2008 we have focused our field research in Tunisia, notably on the geological outcrops exposed in the center of the country (Fig. 1). In the vicinity of the village of Soug-Jedid (Sidi Bouzid Township), the survey of the Eocene shallow marine variegated deposits that are accessible at the foothill of the south-eastern flank of Djebel el Kébar, has allowed for the discovery of a new vertebrate-bearing locality (KEB-1). From this locality, a few isolated teeth of mammals were sorted from several
hundred teeth of selacians by wet-screening about three tons of sediment during the course of three field seasons. Among these mammalian teeth, one can be attributed to a primate (Marivaux et al., 2014), another to a hyracoid (unpublished material), and several others (more than fifty) to a rodent. In this paper, we analyze the dental material of that rodent. These teeth document a diminutive and primitive new species of *Protophiomys*, a basal genus of hystricognathous rodents which is well known from Bir el Ater in eastern Algeria (Nementcha mountains: *P. algeriensis*; Jaeger et al., 1985), Birket Qarun 2

Figure 1. Location map and stratigraphic position of the vertebrate-bearing Djebel el Kébar locus 1 locality (KEB-1) in central Tunisia. A, simplified map of Tunisia showing the location of Djebel el Kébar in central Tunisia (Sidi Bouzid Township, Amanria hamlet – Soug-Jedid village); B, lithostratigraphical log of the Tertiary sedimentological sequence of Djebel el Kébar (south eastern part), indicating the position of the fossiliferous KEB-1 locality (log from Gilles Merzeraud; modified after Merzeraud et al., submitted); C, photograph showing the typical badlands of variegated clays from the early Tertiary sequence of Djebel el Kébar, which corresponds to shallow marine environmental deposits (picture by Gilles Merzeraud). The red arrow indicates the level that has yielded KEB-1; D, photograph of the fossiliferous glauconitic clays of KEB-1 (picture by Monique Vianey-Liaud).
in Egypt (Fayum BQ-2: *P. aegyptensis*; Sallam et al., 2009), and possibly from Dur At-Talah in Libya (DT-Loc 182: “P. durattalahensis”; Jaeger et al., 2010a; see discussion section), three mammalian-bearing localities very close in age at the end of the Eocene (latest middle [?] to early late Eocene). A few teeth, tentatively referred to *Protophiomys* (*P. algeriensis*), were also reported from Sperrgebiet (Silica South and Silica North) in Namibia, two localities that are considered to be Lutetian in age (Pickford et al., 2008; but see discussions about the age of these localities by Marivaux et al., 2012, Coster et al., 2012a, and this paper).

At KEB-1 in Tunisia (Fig. 1), the sediment consists of glauconitic clays, which were formed locally in lagoonal conditions (see Merzeraud et al., submitted). Radiometric K-Ar analyses performed on few grains of glauconite extracted from the glauconitic clays embedding the fossils, indicate that KEB-1 is late middle Eocene in age (Bartonian; datings ranging from 38.7 ± 1.0 Ma to 40.7 ± 1.1 Ma; see Yans in Marivaux et al., 2014). This new species of *Protophiomys* from KEB-1 (*P. tunisiensis* sp. nov.) is therefore among the most ancient representatives of the hystricognathous rodents to be recorded in Africa thus far. Here, we describe and compare these new fossils from Tunisia, and discuss the affinities of this new rodent species, with a special emphasis on possible paleobiogeographic and macroevolutionary implications. The dental terminology (Fig. 2) follows that of Marivaux et al. (2004) (modified after Wood & Wilson, 1936).

**SYSTEMATIC PALEONTOLOGY**

Class MAMMALIA Linnaeus, 1758
Order RODENTIA Bowdich, 1821
Suborder CTENOHYSTRICA Huchon, Catzeflis & Douzery, 2000
Infraorder HYSTRICOGNATHI Tulberg, 1899
*Protophiomys* Jaeger, Denys & Coiffait, 1985 (Fig. 3)

*Type species.* *Protophiomys algeriensis* Jaeger et al., 1985.

*Included species.* *Protophiomys aegyptensis* Sallam et al., 2009; *Protophiomys tunisiensis* nov. sp.; “*Protophiomys* durattalahensis” Jaeger et al., 2010, should be referred to another genus sensu Antoine et al. (2012), Sallam et al. (2012) and Barbière & Marivaux (in press), but see Coster et al. (2012b); *Waslamys attiai* Sallam et al., 2009, should be reassigned to the genus *Protophiomys sensu* Coster et al. (2012b).

**Range and distribution.** Late middle Eocene (Bartonian) of Tunisia to early late Eocene (early Priabonian) of eastern Algeria and northern Egypt (and possibly Lutetian of Namibia; Pickford et al., 2008 [but see discussion in this paper]).

**Emended diagnosis.** Brachydont upper and lower cheek teeth with low but well-defined cusps (-ids) and lophs (-ids); upper molars without protocone but with a small to indistinct central metaconule; metaloph joining the vestigial metaconule and never connecting to the posteroloph; mesolophule non-existent or variably developed (weakly to moderately developed); complete mure absent; complete neo-endoloph-enterocrest complex present on M3, variably developed (complete or incomplete) on M2 (hypoflexus partially or entirely closed lingually), but absent on M1 (hypoflexus open lingually); buccal and lingual walls of upper molars and lower molars, respectively, absent to strongly developed; lower molars with complete or subcomplete ectolophid without well-defined mesoconid (vestigial or absent); posterior arm of the protoconid very short (incomplete metalophulid II, buccally limited); anterior cingulid either very weak or absent.

**Upper tooth**

![Upper tooth diagram](image)

**Lower tooth**

![Lower tooth diagram](image)
Protophiomys tunisiensis nov. sp.

urn:lsid:zoobank.org:pub:8D979CB2-625F-41D7-80E5-AD338BE5DE2

Holotype. KEB-1-014, an upper right M2 (Fig. 3M).

Hypodigm. 1 right dp4 (KEB-1-006), 1 left p4 (KEB-1-002), 3 right p4 (KEB-1-033-035), 6 left m1 (KEB-1-003, 036-40), 4 right m1 (KEB-1-029-032), 8 left m2 (KEB-1-004, 041-046, 067), 7 right m2 (KEB-1-023-28, 069), 3 left m3 (KEB-1-005, 047-048), 6 right m3 (KEB-1-017-022), 1 left DP4 (KEB-1-058), 1 right DP4 (KEB-1-073), 1 right P4 (KEB-1-007), 7 left M1 (KEB-1-007, 010-011, 059-061, 074), 8 right M1 (KEB-1-009, 054-057, 070-072), 5 left M2 (KEB-1-012, 015, 062-063, 075), 4 right M2 (KEB-1-013, 050-052), 4 left M3 (KEB-1-016, 064-066), 2 right M3 (KEB-1-048-049). Dental measurements are reported in Table 1. All these fossils are housed in the paleontological collections of the museum of the “Office National des Mines” of Tunis, Tunisia.

Type locality. Kébar 1 (KEB-1), southern flank of Djebel el Kébar, Amanuria hamlet, Song-Jedid village (Sidi Bouzid region), central Tunisia (Fig. 1).

Age. Late middle Eocene (Bartonian). Dated at ~39.5 Ma by radiometric K-Ar analyses performed on grains of glauconite extracted from the glauconitic clays embedding the fossils of KEB-1 (see Yans in Marivaux et al., 2014).

Etymology. Refers to the provenance of this species (i.e., Tunisia).

Diagnosis. Tiny species, about half the dental size of P. algeriensis or P. aegyptensis. Differs from these latter in having a dental pattern characterized by low and thin transverse crests (-ids), tetralophodont (absence of mesolophule) and less asymmetric in occlusal outline due to its trigonid, which is pinched buccolingually with respect to the talonid (Fig. 3E). The protoconid and metaconid are equal in size, buccolingually well separated, and slightly more elevated than the talonid cuspsids. There is neither metalophulid I nor metalophulid II. Mesially, the tooth bears a small but distinct cuspidate anteroconid. This mesial cusp is linked to the metaconid by a short but high anterior cingulid. This cingulid is developed lingually but not buccally, and the trigonid basin remains open buccally as a result. On the talonid, the entoconid and hypoconid are buccolingually opposed and linked by a thin, low (somewhat faintly visible) and straight hypolophid. The anterior arm of the hypoconid is moderately long and connects both the hypolophid and a mesiodistally elongate ectolophid. This latter longitudinal cristid is thin, straight and complete, being connected to the distal flank of the protoconid. There is neither development of a mesoconid nor an incipient mesolophid in the middle of the ectolophid. The entoconid extends mesially via the development of a strong anterior arm. This latter joins the posterior arm of the metaconid, which is also well-marked. These two arms form a lingual wall, which closes the talonid basin lingually. Distally, the crown displays a low posterior cingulid, which bears a strong hypoconulid near the midline. This distal cusp is virtually as large as the hypoconid, and both cuspsids are linked by a trenchant post-hypoconid cristid. The posterior cingulid remains very low between the hypoconid and hypoconulid, but forms a posterolophid between the hypoconulid and entoconid.

Deciduous lower premolar. Only one dp4 was discovered at KEB-1. This tooth is noticeably longer than wide, and mesiodistally asymmetric in occlusal outline due to its trigonid, which is pinched buccolingually with respect to the talonid (Fig. 3E). The protoconid and metaconid are equal in size, buccolingually well separated, and slightly more elevated than the talonid cuspsids. There is neither metalophulid I nor metalophulid II. Mesially, the tooth bears a small but distinct and cuspidate anteroconid. This mesial cusp is linked to the metaconid by a short but high anterior cingulid. This cingulid is developed lingually but not buccally, and the trigonid basin remains open buccally as a result. On the talonid, the entoconid and hypoconid are buccolingually opposed and linked by a thin, low (somewhat faintly visible) and straight hypolophid. The anterior arm of the hypoconid is moderately long and connects both the hypolophid and a mesiodistally elongate ectolophid. This latter longitudinal cristid is thin, straight and complete, being connected to the distal flank of the protoconid. There is neither development of a mesoconid nor an incipient mesolophid in the middle of the ectolophid. The entoconid extends mesially via the development of a strong anterior arm. This latter joins the posterior arm of the metaconid, which is also well-marked. These two arms form a lingual wall, which closes the talonid basin lingually. Distally, the crown displays a low posterior cingulid, which bears a strong hypoconulid near the midline. This distal cusp is virtually as large as the hypoconid, and both cuspsids are linked by a trenchant post-hypoconid cristid. The posterior cingulid remains very low between the hypoconid and hypoconulid, but forms a posterolophid between the hypoconulid and entoconid.

Permanent lower premolar. The p4 is clearly shorter than dp4, and it exhibits a much simpler dental pattern (Fig. 3A). This is particularly shown in the absence of an anteroconid and a hypolophid, but also in the absence of development of anterior and posterior cingulids. As on dp4, p4 is also slightly asymmetric in having its talonid broader than the trigonid. The protoconid, metaconid, hypoconid, entoconid and hypoconulid are equal in size, but the mesial cuspsids are slightly more elevated. The lingual cuspsids are opposed to the buccal cuspsids. As dp4, p4 displays a long and complete longitudinal ectolophid without a mesoconid. The protoconid and metaconid appear to be more separated than they are on dp4. Both mesial cuspsids are linked by a thin but complete metalophulid II. Distally, the hypoconulid is in a more internal position (more aligned on buccal wall, mesoflexus open) with a well-defined and individualized mesostyle, a trigonid basin open lingually (absence of elevated lingual wall), the absence of neo-endoloph on M2, and by the presence (sometimes) on M1-2 of a very low, incipient enterocrest (hypoflexus always open lingually even with that crest).

Nomenclatural remark. This new species must be referred to as Protophiomys tunisiensis Marivaux 2014, following the article 50.1 and the “recommendation 50A concerning multiple authors” of the International Code of Zoological Nomenclature (ICZN, 1999: 52, 182).

Description

Although fragmentary, the dental material documenting this new rodent species is abundant enough to reconstruct complete but composite upper and lower tooth rows.

Table 1. Dental measurements (in mm) of Protophiomys tunisiensis nov. sp. (L, maximum mesiodistal length; W, maximum buccolingual width; n, total number of specimens [unbroken]; x, mean; SD, standard deviation; OR, observed range).

<table>
<thead>
<tr>
<th>Tooth</th>
<th>n (n')</th>
<th>L</th>
<th>W</th>
<th>x</th>
<th>SD</th>
<th>OR</th>
</tr>
</thead>
<tbody>
<tr>
<td>dp4</td>
<td>1(1)</td>
<td>1.42</td>
<td>-</td>
<td>-</td>
<td></td>
<td></td>
</tr>
<tr>
<td>p4</td>
<td>4(3)</td>
<td>1.29</td>
<td>0.08</td>
<td>1.21-1.37</td>
<td></td>
<td></td>
</tr>
<tr>
<td>m1</td>
<td>10(8)</td>
<td>1.63</td>
<td>0.06</td>
<td>1.56-1.69</td>
<td></td>
<td></td>
</tr>
<tr>
<td>m2</td>
<td>15(13)</td>
<td>1.72</td>
<td>0.07</td>
<td>1.61-1.86</td>
<td></td>
<td></td>
</tr>
<tr>
<td>m3</td>
<td>9(7)</td>
<td>1.72</td>
<td>0.14</td>
<td>1.50-1.89</td>
<td></td>
<td></td>
</tr>
<tr>
<td>DP4</td>
<td>2(2)</td>
<td>1.27</td>
<td>0.05</td>
<td>1.24-1.31</td>
<td></td>
<td></td>
</tr>
<tr>
<td>P4</td>
<td>1(1)</td>
<td>1.42</td>
<td>-</td>
<td>-</td>
<td></td>
<td></td>
</tr>
<tr>
<td>M1</td>
<td>15(12)</td>
<td>1.50</td>
<td>0.06</td>
<td>1.39-1.54</td>
<td></td>
<td></td>
</tr>
<tr>
<td>M2</td>
<td>10(9)</td>
<td>1.56</td>
<td>0.09</td>
<td>1.41-1.72</td>
<td></td>
<td></td>
</tr>
<tr>
<td>M3</td>
<td>6(4)</td>
<td>1.57</td>
<td>0.08</td>
<td>1.44-1.64</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

4
the hypoconid-entoconid axis), and as such it does not form a distinct heel as it does on dp4. Lingually, the talonid remains open due to the absence of development of both the posterior arm of the metaconid and the anterior arm of the entoconid.

**Lower molars.** The three molars are brachydont, longer than wide, and exhibit very similar cuspid and cristid arrangements (Fig. 3B-D). The first molars (m1; Fig. 3B) are distinguishable by the position of the protoconid, which is slightly more lingual than the hypoconid on a mesiodistal axis (i.e., talonid barely broader than the trigonid). The third molars (m3; Fig. 3D) are also clearly recognizable by the position of the entoconid, which is more buccal with respect to the metaconid along the mesiodistal axis (trigonid barely broader than the talonid). The second molars (m2; Fig. 3C) are in contrast perfectly

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**Figure 3.** Lower and upper teeth of *Protophiomys tunisiensis* nov. sp. from the late middle Eocene of Djebel el Kébar locus 1 (KEB-1), central Tunisia (dimensions: length x width, in mm). A, KEB-1-002, left lower p4 (1.30 x 1.13); B, KEB-1-003, left lower m1 (1.67 x 1.45); C, KEB-1-004, left lower m2 (1.73 x 1.53); D, KEB-1-005, lower left m3 (1.78 x 1.46); E, KEB-1-006, lower right dp4 (1.42 x 1.02); F, KEB-1-073, right upper DP4 (1.31 x 1.36); G, KEB-1-008, right upper P4 (1.42 x 1.55); H, KEB-1-009, right upper M1 (1.52 x 1.50); I, KEB-1-010, left upper M1 (1.51 x 1.58); J, KEB-1-011, broken left M1 (1.62 x -); K, KEB-1-012, left upper M2 (1.57 x 1.65); L, KEB-1-013, right upper M2 (1.60 x 1.82); M, KEB-1-014, right upper M2 (1.72 x 1.96); N, KEB-1-015, left upper M2 (1.59 x 1.83); O, KEB-1-016, left upper M3 (1.64 x 1.60).
rectangular, with the distal cusps mesiodistally opposed to the mesial cusps. On the three molars (m1-3), the metaconid and entoconid occupy a position slightly more anterior to the protoconid and hypoconid, respectively. These four main cusps are acute and roughly equal in size. Distally, the hypoconulid is strong, but barely smaller than the other main cusps. This distal cusp occupies a position slightly buccal to the midline. On each dental locus, mesiobuccally one can observe the presence of a short and not very extensive anterior cingulid. On some specimens, this anterior cingulid can reach the mesial base of the metaconid, but its development remains subtle. The three molars are tetrалophodont and characterized by transverse cristids that are sub-parallel, very thin, and relatively low compared to cusp height. The metalophulid I is complete and forms the mesial margin of the teeth. This mesial transverse cristid is slightly curved in its buccal part due to the small distal displacement of the protoconid. The second transverse cristid is a very short posterior arm of the protoconid, which does not reach the lingual margin (or the metaconid), and therefore does not form a metalophulid II. The third transverse cristid is the hypolophid, which links the entoconid to the junction between the ectolophid and the anterior arm of the hypoconid. This latter is distinct, being long and oblique (directed mesiolingually). The fourth transverse cristid is the posterolophid, which is trenchant, curved and bears the hypoconulid. Lingually, the posterolophid weakens but reaches the distal base of the entoconid. On m3, the posterolophid exhibits a shorter radius of curvature than that observed on m1-2, due to the buccal displacement of the entoconid (Fig. 3D). The ectolophid is long, oblique and complete, running from the mesial extremity of the anterior arm of the hypoconid to the protoconid. There is neither a mesoconid nor a mesolophid. Lingually, the talonid basin remains open due to very weak development of the posterior arm of the metaconid and the absence of an anterior arm of the entoconid.

Deciduous fourth upper premolar. Two specimens have been identified, but only one provides occlusal details (Fig. 3F). This specimen is, however, slightly worn and broken mesially, and although appearing to be quadrangular (length = breadth), it was probably trapezoidal in occlusal outline (buccal length > lingual length) judging from the surface of breakage mesiobuccally. This tooth is low crowned, tetrалophodont, and displays four bulbous main cusps (protocon, paracon, metacon and hypocon), plus a minute cusp (metaculide, which is in nearly a central position on the crown. The transverse cristids are low and parallel. The metaconule bears a very short and buccally oriented enamel wrinkle that could correspond to an incipient mesolophid. The anterior crest is not an anteroloph, but a low cingulum that is not connected to the protoconid lingually. Mesiobuccally, this anterior cingulum is worn and broken, but it was probably more extensive. The protoloph is rectilinear and links the lingual side of the paracon to the protoconid mesiobuccally. The metaloph runs lingually from the metaconid but makes a sharp bend at midline toward the metaconule, and it connects to this latter cusp. The anterior arm of the hypoconid is short, oblique, and attached mesially to the metaconule. The posteroloph, connected lingually to the hypoconid, runs buccally and weakens progressively to end at a point directly distal to the metaconid. On the buccal margin, the posterior arm of the paracon and the anterior arm of the metaconid can be weakly to moderately developed, but the mesolophid remains open buccally. Lingually, the protoconid does not show any distal outgrowth (i.e., no neo-endoloph), but a small and low enteroecrest can be observed. This short crest closes partially the hypoflexus lingually. The metaconule also shows a very short and longitudinal wrinkle (incipient mure) that closes the hypoflexus buccally (no connection between the hypoflexus and mesolophid).

Permanent fourth upper premolar. This tooth is less low-crowned than DP4, and is sub-oval in occlusal outline, being much broader transversely than it is long (Fig. 3G). However, its dental pattern is basically similar to that of DP4, although differing in the position and development of some cusps and crests. As DP4, P4 displays a well-defined hypoconid, which develops a strong and short anterior arm that connects a minute metaconule. The protoconid and hypoconid are mesiodistally opposed, while the paraconid is lingually displaced with respect to the metaconid. The anteroloph is very low and very limited buccolingually, being neither connected to the protoconid nor to the paraconid. The protoloph is straight and trenchant, and links the paraconid to the protoconid. The metaloph is oblique (mesiolingually oriented), running from the metaconid to the metaconule. At the metaconule-metaloph connection, there is a low, thin, but long mesolophid, which runs buccally to reach a minute mesostyle in buccocentral position (located between the paraconid and metaconid). The posterolophid is also trenchant, running buccally from the hypoconid to the distobuccal border of the crown. It connects the distal side of the metaconid. Lingually, there is no trace of neo-endoloph (bulbous protoconid), but a very low and oblique enterolophid, running between the protoconid and metaconule, partially closes the hypoflexus.

Upper molars. Although differing in size and occlusal outline shape, the three upper molars share the same dental structure, which is in fact similar to that of P4 and DP4. M1 (Fig. 3H-J) and M3 (Fig. 3O) are roughly equal in size and smaller than M2 (Fig. 3K-N). M3 differs substantially from M1-2 in having a hypoconid strongly displaced buccally, thereby lending a characteristic heart-shaped occlusal outline to the tooth. M3 differs also from M1-2 in having a smaller metaconid, a protoconid buccolingually compressed, a short but well-defined mesolophid, and in showing the development of a long and high neo-endoloph/enteroloph complex, which appears hook-like. This long crest complex connects a minute metaconula (faintly distinguishable) situated at the mesial extremity of a long and longitudinal anterior arm of the hypoconid, thereby closing the hypoflexus lingually. M1s are quadrangular (length = breadth), while M2s are more rectangular, being much broader transversely than long. M1s and M2s differ also by the position of the hypoconid, which is more buccally positioned to the protoconid in M2s, while M1s the hypoconid is distally opposed to the protoconid. As for the premolars, M1-3 are brachydont and tetrалophodont. The main cusps are well-defined from the transverse crests (moderately bunodont). The anterolophid (strongly connected to the protoconid) and posterolophid (strongly connected to the hypoconid) are both trenchant in their lingual part, but weakly progressively in their buccal part. They end at a point directly mesial and distolingual to the paraconid and metaconid, respectively. There is neither a parastyle nor a metastyle, and the paraflexus and posteroflexus remain open buccally. On some M1 and M2, the anterolophid bears a small enamel swelling in its most lingual part that could indicate the presence of a minute anteroconid. Buccally, there is a minute and isolated mesostyle, which is situated between the paraconid and metaconid. There is no buccal wall (absence of posterior arm of the paraconid and anterior arm of the metaconid) and the mesolophid is open buccally. The metaloph is generally oriented buccolingually in its buccal part (parallel to the protolophid). At midline, it can
end abruptly (Fig. 3I, L) or it can bend mesially and sharply toward the metaconule (Fig. 3H, J, K, M, N). In this latter case, the metaloph can either reach the metaconule, or not. Some specimens display an enamel wrinkle emerging from the metaconule (Fig. 3H, I, J, M). This wrinkle is directed buccally, but remains short, thin, and low (highly variable), and it might indicate the presence of an incipient mesolophule. One M2 (KEB-1-013; Fig. 3L) exhibits a longer but low mesolophule, this latter being buccodistally oriented (toward the metacone). This tooth shows also a very short metaloph (limited in its buccal part). Unlike M3, the protocone of M2 and M1 are not clearly buccolingually compressed. On these two dental loci, the protocone remains well-defined (rounded) and there is no development of neo-endoloph. The hypoflexus is always transversely open. On some teeth (most often M2), the presence of a very low and oblique enterocrest can be observed, running between the protocone and metaconule.

Figure 4. Optical and scanning electron photomicrographs of KEB-1-053, an upper incisor for which the enamel microstructure is detailed at different magnification. A, longitudinal section of the incisor showing the enamel layer on the top of the tooth (optical photograph); B, studied area: the portio externa (PE) represents 23% of the total enamel thickness and displays radial enamel, the portio interna (PI) displays multiserial Hunter-Schreger bands (HSBs), which are inclined at 70-80° to the enamel-dentine junction (EDJ); C, microstructural detail of ‘B’ showing five decussating HSB with three (often four) prisms per band, and the inter-prismatic matrix (IPM) anastomosing regularly between prisms, and with its crystallites running nearly parallel or at a low angle to the prism direction; D, microstructural detail of ‘B’ showing two HSBs with three prisms and similar IPM arrangement than ‘C’; E, microstructural detail of ‘B’ showing three HSBs but more particularly the IPM, which anastomoses regularly between prisms; F, microstructural detail of ‘B’ showing one HSB, notably the prism shape in section, which is suboval. The pattern of the IPM arrangement typifies a primitive multiserial condition designated as multiserial subtype I (see Martin, 1994).
**Incisor Enamel Microstructure.** Among the dental material collected from wet-screening of the KEB-1 sediment are a few fragments of rodent incisors. Although incomplete, the size of these incisors is compatible with that of the molars of *Protophiomyx tunisiensis*, thereby suggesting that these incisors could tentatively be referred to this taxon (the sole rodent recorded at KEB-1). One fragment of upper incisor (KEB-1-053) was embedded in artificial resin and subsequently polished longitudinally (Fig. 4A). The polished section was then etched for 30 sec with H$_2$PO$_4$ (37%) to make the microstructural details visible (Fig. 4B-F).

In longitudinal section (Fig. 4B), the total enamel thickness is about 0.21 mm and is formed by two distinct layers: the *portio interna* (PI) and the *portio externa* (PE). The PE has radial enamel and represents about 23% of the total enamel thickness. The PI is very thick (≥ 0.16 mm) and consists of decussating layers of prisms, which are Hunter-Schreger bands (HSBs). There are generally 3-4 prisms wide per band (Fig. 4C-F). The prisms are slightly flattened, thereby appearing suboval in cross section (Fig. 4E-F). The HSBs are inclined at 70-80° to the enamel-dentine junction (EDJ). The inter-prismatic matrix (IPM) in the PI is thin and never surrounds the prisms. The IPM crystallites run nearly parallel or at a very low angle to the prism direction, and they anastomose regularly between the prisms (Fig. 4D-F). There is no transition zone between adjacent decussating HSBs. Such a crystallite arrangement in PI determines a multiserial enamel (Korvenkontio, 1934). The pattern of the IPM arrangement typifies a primitive multiserial condition, which is recognized as multiserial subtype I (see Martin, 1994). This enamel microstructure is therefore slightly less advanced (more primitive) than that observed in the incisor analyzed from Bir el Ater (Algeria; Martin, 1993), which show an IPM arrangement typifying a multiserial subtype II (acute angular and anastomosing IPM). This incisor from Bir el Ater is probably referable to *Protophiomyx algeriensis* (Table 2; Jaeger et al., 1985). Regrettably, the incisor enamel microstructure has not been investigated in hystricognaths from Dur At-Talah (DT-Loc 1&2) and Fayum Birket Qarun 2 (BQ-2) localities, thereby restricting the scope of comparisons that can be made with the other known species of *Protophiomyx*, but also other species of close genera.

**Comparisons**

**Remark.** We provide in Table 2 a list of early Afro-Asian hystricognathous rodents with information regarding their geographical provenance, age, and references.

**Dental comparisons.** Among the different species attributed to the genus *Protophiomyx* (*P. algeriensis*, *P. aegyptensis*, “P.” *durattalahensis*; Figs 5-6), *P. tunisiensis* nov. sp. is relatively smaller in size (roughly similar in size to “P.” *durattalahensis*) and slender in overall morphology. Compared with the other species, *P. tunisiensis* displays a dental pattern that is characterized by low and thin transverse crests (-ids) and by well-defined and salient cusps and cuspids (Fig. 5A-B). Unlike *P. algeriensis*, *P. aegyptensis* and “P.” *durattalahensis*, the trigon basin of upper molars of *P. tunisiensis* is open buccally. This is due to the lack of strong development of the posterior arm of the paracone and the anterior arm of the metacone (lack of buccal wall) as is observed in the other species. There is only a minute, low and isolated mesostyle between the paracone and metacone, which slightly obstructs the mesoflexus buccally. In the other species of the genus, a mesostyle is also distinguishable (somewhat larger), but it is entirely included in the buccal wall. Upper molars (notably M1-2) of *P. tunisiensis* do not exhibit a mesiodistal pinch of the protocone associated with the development of a distal outgrowth (i.e., neo-endoloph) as it is observed in all other species of the genus (Fig. 5C, E), but also in the *Protophiomyx*-like taxa such as *Waslamys* (*W. attiai*; Fig. 6C) or *Baluchimys* (*B. krambiense*, *B. barryi*), and in more recent hystricognath taxa (Table 2) such as *Phiomys* (*Ph. hammodulai*, *Ph. andrewsi*), *Acritophiomys* (*A. bouni*), *Metaphiomys* (*M. schaubi*), *Hodsahibia* (*H. azrea*, *H. beamshaiendri*, *H. gracilis*), *Bugtmys* (*B. zafarullahi*), or *Lophibaluchalia* (*L. pilbeamii*). This simple and primitive structure of the protocone is observed in bunodont taxa of the group such as *Talaphiomyx* (*T. lavocati*, *T. libycus*; Fig. 6E) and *Lindsaya* (*L. derabugtiensis* (Table 2). However, *P. tunisiensis* differs substantially from these two latter genera in having a more crested dental pattern with less inflated cusps and cuspids, and in having a metacone on upper molars, which is virtually undistinguishable (vestigial) compared to the prominent metacone present in *Talaphiomyx* and *Lindsaya*. The condition of the metacone in *P. tunisiensis* differs also from the condition observed in upper molars of some Asian hystricognaths (such as *Ottomania*, *Bugtmys* and *Hodsahibia*), in which this central cusps is still distinct.

On the upper molars, *P. tunisiensis* also differs from *P. algeriensis* and *P. aegyptensis* in lacking the strong development of the oblique enterocrest (notably on M2). In *P. tunisiensis*, this crest is variably developed and can be absent (Fig. 3). When it is present, it remains very low and it never closes the hypoflexus lingually as it does on M2 of *P. algeriensis* and *P. aegyptensis* (Fig. 5C, E). It is noteworthy that the enterocrest is absent on M1-2 of “P.” *durattalahensis*, and as such the hypoflexus remains always open lingually in this species despite the strong and long development of the neo-endoloph (Fig. 6A).

As in *P. algeriensis* and *P. aegyptensis*, upper molars of *P. tunisiensis* are primarily tetralophodont, a condition which is described by the presence of an anteroloph, a protoloph, a metaoloph and a posteroloph (Fig. 5A, C, E). However, some upper molars of these three species can bear a short and buccally oriented enamel wrench that is attached to the vestigial metacone. This short crest could represent an incipient mesoloph. The presence of this character is rare on upper molars of *P. tunisiensis* and *P. aegyptensis* but it is more frequent on molars of *P. algeriensis*. In contrast, this character is constant and well-developed (long mesoloph reaching the buccal margin [mesostyle]) in “P.” *durattalahensis* (Fig. 6A) but also in *Waslamys attiai* (Fig. 6C), a condition that completes the derived pentalophodont pattern in these two latter species. The pentalophodonty of upper molars, characterized by a strong development of that central transverse crest (i.e., mesoloph), is common in several more advanced Afro-Asian hystricognathous taxa (e.g., *Acritophiomys*, *Phiomys*, *Metaphiomys*, *Turkanamys*, *Lophibaluchalia*, *Hodsahibia*, *Bugtmys*...), but also in the earliest known South American hystricognaths (*Cachiyacui*, *Canaanimys*; Antoine et al., 2012). Interestingly, one upper molar identified as an M2 (KEB-1-013; Fig. 3L) of *P. tunisiensis*, exhibits an unusual tetralophodont pattern compared to the other upper teeth referred to this species. This tooth has a very short metaloph (virtually non-existent, limited to its buccal part) but in contrast displays a long mesoloph, which is oblique (buccodistally oriented). This tooth is likely a variant of the other upper molars. However, in developing such an arrangement of the metaoloph-mesoloph, the dental
<table>
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<th>Provence</th>
<th>Age</th>
<th>References (e.g.)</th>
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<td>Tunisia (Odhéle Kébén, KEB-1)</td>
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<td>This paper, Jaeger et al., 1985, Sallam et al., 2009, Marivaux et al., 2010, Coster et al., 2015</td>
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<td>Egypt (Fayoum, BO-2)</td>
<td>late middle Eocene to early late Eocene</td>
<td>Sallam et al., 2009, Sallam et al., 2011, Wood, 1968, Coster et al., 2015</td>
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Table 2. List of most of the early hystricognathous rodents from the Paleogene of Africa, South Asia, Asia Minor, and South America that are mentioned in the comparison section.
pattern of this tooth, although less bunodont, is somewhat reminiscent of that on the M2 of *Talahphiomys libycus* (Jaeger et al., 2010a: 203, fig. 5u).

The P4 attributed to *P. tunisiensis* (KEB-1-007; Fig. 3G) exhibits a long but very low central transverse crest, which runs from the mesial extremity of the anterior arm of the hypocone to a minute and buccal mesostyle. The presence of a mesolophule is not observed on P4 of the other species of the genus. Additionally, the anteroloph of KEB-1-007 is not developed and not connected to the protocone, and in fact, it appears as a low cingulum buccolingually limited. This condition differs from that observed on P4s of *P. algeriensis*, *P. aegyptensis* or “P.” *Durattalahensis*, in which the anterior cingulum is longer and more elevated, thereby forming a functional and trenchant

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**Figure 5.** Schematic line drawings of composite upper and lower dentitions of early stem hystricognaths from Africa. **A-B**, *Protophiomys tunisiensis* nov. sp.: DP4 and P4, M1-3 (A), dp4 and p4, m1-3 (B); **C-D**, *Protophiomys algeriensis* Jaeger et al., 1985: DP4 and P4, M1-3 (C), dp4 and p4, m1-3 (D); **E-F**, *Protophiomys aegyptensis* Sallam et al., 2009: DP4 and P4, M1-3 (E), dp4 and m1-3 (F). Some isolated right teeth are reversed to be included in the composite left tooth rows. Scale bars: 1 mm. Drawings by Monique Vianey-Liaud.
anteroloph. Finally, KEB-1-007 differs also from its counterparts in the other species of the genus in showing a well-defined and cuspatc hypocone as well as a stronger and longer anterior arm of that hypocone, which delimits (with the protocone) a shallow internal sinus.

As in *P. algeriensis*, *P. aegyptensis* and “*P.*” *durattalahensis*, the anteroloph and posteroloph on the upper molars of *P. tunisiensis* are both trenchant in their lingual part, but in *P. tunisiensis* they weaken progressively in their buccal part. These two lophs also expand less buccally (buccolingually shorter) than in molars of the other species. Indeed, the anteroloph and posteroloph end at a point directly mesial and distolingual to the paracone and metacone, respectively. This arrangement of these two lophs make the crown more ovoid in occlusal outline, a condition which clearly differs from the more quadrangular occlusal outline characterizing upper teeth of the other species of the genus (Figs 5A, C, E and 6A). In *P. tunisiensis*, there is neither a parastyle nor a metastyle, the anteroloph and posteroloph end buccally close to the base of the paracone and metacone, respectively (not connected to them), and the paraflexus and posteroﬂexus remain open buccally as a result.

![Figure 6](image-url)
As in *P. algeriensis* and *P. aegyptiensis* (but also in *Ottomania, Baluchimys, Hodshahibia, Lophibaluchia, Talahphiomys*), upper molars of *P. tunisiensis* do not show the development of a central longitudinal crest (the mure; Fig. 2), which is generally short and links the metaconule (or the mesial extremity of the anterior arm of the hypocone) with the protoloph (i.e., complete mure) in more advanced species such as *Acriropheiomys, Phiomys, Metaphiomys, Turkanamys* or even *Bugtimys*. In the absence of this character, the hypoflexus remains open buccally. A rough stage of this longitudinal crest (incipient, very short mure) can be however observed in some upper molars of *P. algeriensis* and *P. aegyptiensis*, but never in *P. tunisiensis*. In contrast, in “P” *durattalahensis*, the mure is often complete, incomplete or absent on upper molars (Fig. 6A).

Concerning the lower teeth, *P. tunisiensis* differs from the other species of the genus in having the cristids less developed (i.e., weakly elevated), and in having more defined and bulbous cusps. The hypocone is not mesiodistally compressed and has neither a mesial outgrowth nor an ectostylid, three characters that can be observed in lower molars of *P. aegyptiensis* and “P” *durattalahensis* (Figs 5D and 6B). On lower molars of *P. tunisiensis*, the posterolophid is trenchant in its buccal part and then weakens progressively in its lingual part. This cristid ends at the distal base of the entoconid (without connection), and as such the metalaxid remains open lingually. Lower molars of the other species exhibit a lingual wall made by the strong and high posterolophid (connected to the entoconid) and a strong, long and high posterior arm of the metaconid (including sometimes a mesostylid). This lingual wall is particularly elevated and continuous (including the entoconid) in *P. algeriensis* and in “P” *durattalahensis* (Figs 5D and 6B), while it can be interrupted (narrow notch, directly mesial to the entoconid) in *P. aegyptiensis* (Fig. 5F). Despite the presence of a notch in some case, the lingual wall closes the talonid basin and the metalaxid lingually in *P. algeriensis*, *P. aegyptiensis* and “P” *durattalahensis*. A continuous and elevated lingual wall is also observed in *Waslamsys* (Fig. 6D). In *P. tunisiensis*, there is no elevated lingual wall; the posterior arm of the metaconid is present but moderately long and low, there is no defined mesostylid, and the talonid basin remains open lingually.

Mesially, lower molars of *P. tunisiensis* display a short (limited buccally) but distinct anterior cingulid (Figs 3B-D and 5B). This cingulid is not observed in the other species of *Protophiomys*, nor in *Waslamsys, Gaudeamus*, or the earliest known caviomorphs (*Canaanymi* and *Cachiyacui*). In contrast, this character is widespread and often well-developed among early hystricognaths such as the African *Acriropheiomys, Talahphiomys, Phiomys, Metaphiomys*, but also in the Asian *Ottomania, Confiniummys, Baluchimys, Hodshahibia, Bugtimys*, and *Lophibaluchia*. The absence of this character in *P. algeriensis*, *P. aegyptiensis* and “P” *durattalahensis* is likely a derived condition, a loss which also occurred in *Waslamsys, Gaudeamus* and the earliest caviomorphs.

Finally, *P. tunisiensis* differs also from the other species of the genus, notably in some morphological details of the permanent and deciduous premolars. As in the other species, the dp4 of *P. tunisiensis* displays a long and complete longitudi- nal ectolophid, but does not bear any trace of a mesoconid or a short mesolophid as can be observed on dp4s of *P. algeriensis* and *P. aegyptiensis* (Fig. 5D, F). This lower deciduous premolar (KEB-1-006; Fig. 3E) displays also a slender and incomplete (interrupted buccally) hypolophid, while this cristid is generally strong and complete in dp4s of the other species. There is also neither a metalophulid I nor a metalophulid II (no posterior arm of the protoconid), whereas in the other species, dp4s display either a complete or an incomplete metalophulid II, and a complex anterior cristid (composite of the metalophulid I and the anterolophid). In contrast, dp4 of *P. tunisiensis* bears a small anterocristid, a character which is not observed in the other species of the genus, but present in dp4 of *Talahphiomys lavocati* (Fig. 6F). The p4s of *P. tunisiensis* differ also from the p4s in the other species in the absence of metalophulid I, while this cristid is present and complete in *P. algeriensis*, “P” *durattalahensis* and also in *Waslamsys* for which the p4 is known. In contrast, they display a complete metalophulid II, while this cristid is generally incomplete (short posterior arm of the protoconid) in the other species of the genus, as well as in *Waslamsys*.

**DISCUSSION AND CONCLUSIONS**

This new rodent species from Tunisia that we refer to the genus *Protophiomys* is one of the most slender species of that genus. *Protophiomys tunisiensis* is roughly equivalent in size to “P” *durattalahensis*, and half the dental size of *P. algeriensis* and *P. aegyptiensis*. Its dental pattern is characterized by well-defined and salient cusps and cuspsids, and by low and thin transverse and longitudinal crests and cristids. Buccal margins of upper molars and lingual margins of lower molars do not exhibit the enamel walls as developed as those observed in the other species of the genus, as well as in *Waslamsys*, a coeval *Protophiomys*-like genus (see discussion below about this genus). Upper molars of *P. tunisiensis* are tetralophodont and do not show any marked tendency toward pentalophodonty, an occlusal pattern which is in contrast particularly well-developed in “P” *durattalahen- sis*, and somewhat “incipient” in *P. algeriensis* and to a lesser extent in *P. aegyptiensis* (presence in some upper molars of these two latter taxa of a short mesolophe). Upper molars of *P. tunisiensis* do not exhibit a strong neo-endoloph (except on M3) as observed in the other species of the genus, but they present a very low and thin entocrist, a character which is particularly strong in *P. algeriensis* and *P. aegyptiensis*. “P” *durattalahensis* does not show this character on its upper molars. In fact, “P” *durattalahensis* displays a dental pattern that appears to be evolutionarily more advanced and somewhat divergent from that characterizing *P. algeriensis*, *P. aegyptiensis* and *P. tunisiensis*. In particular, this is shown in the overall aspect of the teeth, which are less brachydont with crests and cristids more elevated and thicker (robust), but also in the development of a fully achieved pentalophodont pattern of upper molars, the presence of a strong mure (complete or incomplete), and in the lack of entocrist. By this suite of derived dental traits and the lack of some characters otherwise found only in *Protopho- imys*, in our opinion the species “P” *durattalahensis* could be regarded as belonging to a genus close to, but distinct from *Protophiomys*. This hypothesis was recently suggested by cladistic assessments of the dental evidence. Indeed, recent phylogenetic analyses (e.g., Antoine et al., 2012; Sallam et al., 2012; Barbière & Marivaux, in press) advocate the para- or polyphyly of the genus *Protophiomys*, in placing “P” *durattalahensis* clearly apart from the other species of the genus. In contrast, analyses performed by Coster et al. (2010, 2012b) rather suggest that the three species of *Protophiomys* (*P. algeriensis, P. aegyptiensis*, and “P” *durattalahensis*) are closely related. Sallam et al. (2012: 297) have suggested that “P” *durattalahensis* would rather be “a form that is very similar to *Waslamsys* from BQ-2, and possibly congeneric”.

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Figure 7. Temporal distribution of hystricognathous rodents during the Paleogene. Asian “ctenodactyloids” (Ct) are the sister group of the Hystricognathi clade. Zegdoumyid anomaluroids (Zeg) are the only rodents recorded in Africa during the early Paleogene (late early to early middle Eocene [Chambi, Gour Lazib, Aznag and BC-Sperrgebiet]). For South America, we have mentioned only the most ancient taxa known thus far. TIHD, time interval of hystricognath dispersals.

Protophio-

mys tunisiensis nov. sp. from KEB-1 is the most ancient representative of hystricognaths in Africa, but it can be expected that hystricognaths were already present on that landmass given the caviomorph fossil evidence from the late middle Eocene of Peruvian Amazonia. In the context of an Asian origin for the Hystricognathi clade: the ancestral Asian hystricognaths likely colonized Africa sometime during the middle Eocene (TIHD, end of the Lutetian); hystricognaths were also able to continue at that time (during the TIHD) their pattern of intercontinental dispersal across the South Atlantic to invade South America. Africa might have played the role of a stopover for pioneer New World hystricognaths, “within a land-mass-hopping” process between Asia and South America (Antoine et al., 2012).

However, Waslamys lacks several diagnostic characters of “P.” durattalahensis (e.g., presence of a complete mure, thick and tall crests and cristids, hypoflexus open lingually on M2 [no enterocrest]), but in turn exhibits anatomical details that are otherwise found only in Protophionys (e.g., enterocrest, weak and variable development of the mesolophule). These differences are admittedly tenuous but in our opinion would rather suggest that Waslamys attiai is congeneric with Protophionys. This hypothesis was recently advocated by Coster et al. (2012b), who proposed the subfamily Protophionyinae (a clade including Protophionys + “Waslamys”). In case Waslamys would be a junior synonym of Protophionys, and given that “P.” durattalahensis appears to us somewhat autapomorphic, this latter species from Dur At-Talah could then be the type and only species of a new genus. A revision of the species “P.” durattalahensis is therefore required, but this study is beyond the scope of this paper. Regardless of the ongoing controversies over the alpha taxonomy of these early hystricognaths from Africa, the new rodent species from Tunisia we describe here is particularly critical for a better understanding of the morphological evolutionary tendencies observed in this rodent group. The dental specimens collected from KEB-1 display a suite of anatomical details that are primarily found in the different species of Protophionys, but with a lesser degree of development compared with the other species included in the genus or in the close genera (i.e., Waslamys attiai and “P.” durattalahensis). Such an apparent primitive evolutionary stage is corroborated by the greater antiquity of this Tunisian species (~39.5 Ma, late middle Eocene) with respect to that of the Algerian and Egyptian species (i.e., earliest late Eocene; Seiffert, 2006, 2012).

Even if for a long time the record of earliest hystricognaths from the Old World was exclusive to Africa, over the past decades, hystricognaths (“baluchimyines”) have been also recorded in the Paleogene of southern Asia (including the Indian Subcontinent and South-East Asia) and Asia Minor, notably from Thailand (Krabi, latest Eocene, Marivaux et al., 2000), Turkey (Süngülü, late Eocene/early Oligocene; de Bruijn et al., 2003), and from Pakistan (Bugti Hills, early Oligocene, Flynn et al., 1986; Marivaux et al., 2002; Marivaux & Welcomme, 2003; Zinda Pir, early Oligocene, Flynn & Cheema, 1994). These Asian fossil discoveries have demonstrated that this tropical province was also a major centre of an early hystricognathous rodent radiation during the Paleogene, and most likely the ancestral homeland of the Hystricognathi clade owing to the sister-group relationships with the Asian “ctenodactyloid” rodents (i.e., Ctenodactylidae, Tamquammyiidae, Chapattimyidae, Yuomyidae, and Diatomyidae; Marivaux et al., 2002, 2004; Dawson et al., 2006; Sallam et al., 2009, 2012; see also Huchon et al., 2000, 2007 for the Ctenohystri-

ca). However, the first representatives of the Hystricognathi clade in Asia date only from the end of the Eocene epoch (e.g., Baluchimys krabiensis [Krabi, Thailand], Ottomania proavita
[Süngülü, Turkey]; Table 2; Fig. 7). Given the phylogenetic relationships between Hystricognathi and Eocene Asian “ctenodactyloids”, a much greater antiquity for the Hystricognathi clade might then be expected in Asia, an antiquity which might be traced back at least to the early middle Eocene (Antoine et al., 2012; Barbieri & Marivaux, in press). Such a great phylogenetically implied antiquity for hystricognaths points out the incompleteness of their Asian fossil record, a long temporal gap (>10 Myr; Fig. 7) that must be better documented, in order to understand the origin and early adaptive radiation of that rodent group in Asia.

According to our current knowledge of the fossil record in Africa, hystricognaths are absent in significantly older, but reasonably well sampled, Eocene North African localities such as Chambi in western central Tunisia, or those from the Gour Lazib in the Sahara of western Algeria; localities dating from the late early to early middle Eocene (Hartenberger et al., 1997; Adaci et al., 2007; Coster et al., 2012a). These localities have yielded only zegdoumyid anomalurids, a group of rodents endemic to Africa that is not closely related to the clade Hystricognathi (Vianey-Liaud et al., 1994; Marivaux et al., 2004, 2011; Marivaux, unpublished new data). Another mammal-bearing locality from North Africa, Aznaz, Morocco, dating from the middle middle Eocene (middle Lutetian), has yielded one isolated rodent incisor (Tabuce et al., 2005). The material is meagre but the study of the enamel microstructure of this tooth has revealed a Schmelzmuster transitional from the pauciserial to the uniserial condition, a primitive type of enamel which is typical to that found in zegdoumyids from Algeria and Tunisia (e.g., Martin, 1993; Marivaux et al., 2011). A few teeth, tentatively referred to Propothiomys (P. cf. algeriensis), are reported from Sperrgebiet (Silica North and Silica South) in Namibia, two localities which are considered as dating from the middle middle Eocene (middle Lutetian; Pickford et al., 2008). However, there is an uncertainty regarding the Eocene age of these localities (see Coster et al., 2012a; Marivaux et al., 2012) since “Propothiomys” from Silica North is found in association with taxa that show closer affinities to Miocene rodents: a daimantomyid (Prepononomys bogenfelsi), a thryonomyid (Apoderes cf. stromeri), a myiophyiomyd (Silicamys cingulatus), and a bathyergid (cf. Bathyergoides sp.). Furthermore, the robustness of the crests and cristids, and the moderate brachydonty observed in the few teeth referred to Propothiomys from Sperrgebiet do not match the conditions found in the different Eocene species of that genus. For this reason and because of the abundance of associated rodent species that are clearly of Miocene affinities, the presence of Propothiomys in Sperrgebiet, and the middle Eocene age of the Silica rodent-bearing localities are questionable. These Silica localities are most likely Miocene in age, and as such the alleged “Propothiomys”-like teeth are certainly referable to another genus. In contrast, the Black Crow locality from Sperrgebiet (close to the Silica localities; Pickford et al., 2008) displays a diverse assemblage including macro- and micromammals, which are undoubtedly of Eocene affinity. This locality has notably yielded a zegdoumyid rodent (Glibia namibiensis [Zegdoumys namibiensis sensu Marivaux et al., 2011]), but so far no hystricognaths. Although the Paleogene fossil record of North and Sub-Saharan Africa is far from being well-documented, there is currently no evidence of hystricognaths on that landmass during the early Paleogene. Given the age of the KEB-1 locality from Tunisia (~39.5 Ma, late middle Eocene), Propothiomys tunisiensis nov. sp. is therefore the most ancient representative of hystricognathous rodents in Africa that is known thus far. The same is true for the primate recorded at KEB-1 (i.e., Amamria tunisiensis), which represents the most ancient occurrence of anthropoids in Africa (Marivaux et al., 2014).

In the context of an Asian origin for both hystricognath rodents (see aforementioned references on that subject) and anthropoid primates (e.g., Beard et al., 1994, 1996, 2009; Chaimanee et al., 1997, 2012, 2013; Jaeger et al., 1999, 2010b; Takai et al., 2001; Beard & Wang, 2004; Jaeger & Marivaux, 2005; Marivaux et al., 2005a; Kay, 2012; Seiffert, 2012), and given their apparent absence in early to middle middle Eocene localities of North Africa (e.g., Tabuce et al., 2005, 2009; Marivaux et al., 2011, 2013), these groups would then have dispersed from Asia to Africa sometime during the middle Eocene (Fig. 7). But when exactly these dispersals occurred remains unclear. Indeed, the temporal gap (~5 Myr), spanning from the early middle Eocene (i.e., after Chambi, Glib Zegdou, Gour Lazib) to the late middle Eocene (i.e., before Kébar) in the African mammal fossil record, prevents any clear understanding of the nature and timing of middle Eocene immigrations of Asian terrestrial mammals into Africa (Fig. 7). Regarding hystricognathous rodents and particularly the close relationships between Old and New World hystricognaths, it is worth noting that these Asian stem hystricognaths that colonized Africa were in fact able to continue their pattern of intercontinental dispersal across the South Atlantic to invade South America as well. This scenario relies notably on the recent discovery of stem caviomorph rodents in ~41 Myr deposits from Peruvian Amazonia (Antoine et al., 2012). These early caviomorphs exhibit a dental pattern strikingly reminiscent (but somewhat autapomorphic and advanced in some dental traits) of that of their Paleogene African counterparts, a statement which suggests that this Amazonian locality “probably documents the earliest stages of caviomorph evolution” (Antoine et al., 2012: 1321). In a notable implication, this discovery indicates also that the dispersal of stem hystricognaths to South America from Africa must have occurred sometime during the middle Eocene, and a fortiori that hystricognaths must have been present in Africa at least just before 41 Myr. Interestingly, the middle Eocene was marked by a global warming recorded at the beginning of the Bartonian (“Mid Eocene Climatic Optimum” – MECO; Zachos et al., 2008; Bohaty et al., 2009; Vandenberghe et al., 2012). In this climatic context, the question arises as to whether these changes in the global paleoenvironmental conditions at that time have played a significant role in the dispersals of this rodent group but also other mammal groups from Asia to Africa, and then to South America (Marivaux et al., 2000, 2005b, 2011, 2014; Tabuce & Marivaux, 2005; Sallam et al., 2009, 2010a-b; Antoine et al., 2012; Chaimanee et al., 2012; Barbieri & Marivaux, in press). In all cases, the much warmer and wetter conditions recorded at that time have certainly factored into the initial and successful adaptive radiations of hystricognaths in both Gondwanian landmasses.

In sum, if Propothiomys tunisiensis nov. sp. is so far the most ancient representative of hystricognaths in Africa, it can however be expected that this rodent group was already present on that landmass given the new caviomorph fossil evidence. The arrival of hystricognaths in Africa from South Asia certainly predates the depositional period of the Kébar sediments (KEB-1), but perhaps not by much time (Fig. 7). P. tunisiensis was perhaps morphologically not so far from the pioneer hystricognath that colonized Africa from South Asia. Only continuing field efforts in Lutetian deposits of Africa, South Asia but also South America will allow for substantial
discoveries of early hystricognathous rodents. These new data should provide us a more accurate temporal framework of the hystricognath dispersal events, but also further our understanding of the mode and pathways of these dispersals, as well as the macroevolutionary and phylogenetic patterns of their initial radiations.

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