New phiocricetomyine rodents (Hystricognathi) from the Jebel Qatrani Formation, Fayum Depression, Egypt

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ABSTRACT

Background: The rich rodent assemblages from the Eocene–Oligocene deposits of the Jebel Qatrani Formation (Fayum Depression, Egypt) have important implications for our understanding of the origin and paleobiogeography of Hystricognathi, a diverse clade that is now represented by the Afro-Asiatic Hystricidae, New World Caviomorpha, and African Phiomorpha.

Methods: Here we present previously undescribed material of the enigmatic hystricognath clade Phiocricetomyinae, from two stratigraphic levels in the lower sequence of the Jebel Qatrani Formation—a new genus and species (Qatranimys safroutus) from the latest Eocene Locality 41 (~34 Ma, the oldest and most productive quarry in the formation) and additional material of Talahphiomys lavocati from that species’ type locality, early Oligocene Quarry E (~31–33.2 Ma).

Results: The multiple specimens of Qatranimys safroutus from L-41 document almost the entire lower and upper dentition, as well as mandibular fragments and the first cranial remains known for a derived phiocricetomyine. Specimens from Quarry E allow us to expand comparisons with specimens from Libya (late Eocene of Dur at-Talah and early Oligocene of Zallah Oasis) that have been placed in T. lavocati, and we show that the Dur at-Talah and Zallah specimens do not pertain to this species. These observations leave the Fayum Quarry E as the only locality where T. lavocati occurs.

INTRODUCTION

Hystricognathi is a diverse clade of rodents that is characterized by a mandibular angular process situated lateral to the long axis of the lower incisor, multiserial Hunter-Schreger bands of incisor enamel, and enlarged infraorbital foramina, among other features.
Hystricognaths likely originated in the middle Eocene (Marivaux & Boivin, 2019), and in a short time window diversified and radiated across three continents: Asia, South America, and Afro-Arabia (Marivaux & Boivin, 2019). Each of these epicenters housed a distinctive clade—Hystricidae (Old World porcupines), Caviomorpha (New World hystricognaths), and Phiomorpha (African cane, dassie, and mole rats), respectively (Singleton, Dickman & Stoddart, 2006). The Asian tropics are considered to have been the ancestral homeland for Hystricognathi (Sallam, Seiffert & Simons, 2011; Barbière & Marivaux, 2015) despite the fact that the oldest known fossil occurrences of hystricognaths are from Africa (Sallam et al., 2009; Marivaux et al., 2014). Numerous molecular studies have placed Hystricidae as the sister group of a Caviomorpha-Phiomorpha clade (e.g., Huchon et al., 2007; Meredith et al., 2011; Patterson & Upham, 2014; Campbell et al., 2021). Caviomorpha and Phiomorpha are estimated to have split around 39–43 Ma (Sallam & Seiffert, 2016; Patterson & Upham, 2014).

Phiocricetomyinae is an enigmatic Afro-Arabian clade of small hystricognaths with bunodont and simple low-crested cheek teeth whose core members are currently known solely from dental remains. The fossil record of this group is very limited, and its phylogenetic position relative to Phiomorpha is uncertain; phiocricetomyines have variously been placed outside of the Phiomorpha-Caviomorpha clade (Sallam et al., 2009; Sallam, Seiffert & Simons, 2011; Sallam & Seiffert, 2016; Marivaux & Boivin, 2019) or as stem phiomorphs (Sallam & Seiffert, 2016, 2019), suggesting that this group is of key importance for understanding polarities of dental characters near the base of the hystricognath radiation. In 1968, Wood described the first genus and species of this group, a peculiar and highly derived form represented by a mandible with dP₄-M₂ that he named Phiocricetomys minutus; it remains the youngest known member of Phiocricetomyinae, being from one of the youngest fossil-bearing levels ( Quarry I, ~29–30 Ma) (Fig. 1) in the Fayum Depression. Lavocat (1973) later created the subfamily.
Phiocricetomyinae to contain *P. minutus*. Phylogenetic analyses of early Afro-Arabian Hystricognathi (*Holroyd, 1994; Sallam et al., 2009; Sallam, Seiffert & Simons, 2011, 2012; Sallam & Seiffert, 2016, 2019*) have since placed another taxon described by *Wood (1968)—early Oligocene “Phiomys” lavocati* from the ~31–33.2 Ma Fayum Quarry E (Fig. 1)—with *Phiocricetomys* to the exclusion of other hystricognaths. Published remains of “*P.*” lavocati from Quarry E have thus far been limited to dP₄-M₃, and published figures of these specimens are highly schematic line drawings.

In her doctoral dissertation, *Holroyd (1994)* was the first to propose a relationship between “*Phiomys*” lavocati and *Phiocricetomys*, wherein she proposed the new genus *Elwynomys* for “*P.*” lavocati—a decision based on insights provided by new material from L-41 (see below) and the type locality for “*P.*” lavocati (Quarry E) (Fig. 1). However, the name *Elwynomys* was never published in a way that satisfies the criteria of the International Commission on Zoological Nomenclature (ICZN). *Jaeger et al. (2010)* subsequently erected the generic name *Talahphiomys* for “*P.*” lavocati, based on their study of isolated teeth collected from the Idam Unit of the Dur at-Talah escarpment in central Libya that those authors considered to be conspecific with “*P.*” lavocati. Using the new material from Quarry E described here, we show that the specimens from Dur at-Talah actually do not belong in the species *lavocati*, but *Talahphiomys* nevertheless has priority as the generic replacement name for “*P.*” lavocati. *Coster et al. (2012)* has since identified isolated lower teeth from the early Oligocene “rodent locality 5” (ZR5) in the Zallah Oasis of Libya as *T. lavocati*, but we are able to demonstrate that the ZR5 specimens also do not belong in that species, and that *T. lavocati* is restricted to the Fayum Quarry E. Extensive new material of an additional new phiocricetomyine genus and species from the latest Eocene Quarry L-41 (herein named *Qatranimys safroutus*, see below) provides the first detailed insights into the craniodental morphology of, and intraspecific variation within, a phiocricetomyine species.

**Fossil localities**

The late Eocene–early Oligocene Jebel Qatrani Formation (*Seiffert, 2006*) that is exposed north-northwest of Birket Qarun in the Fayum Depression, Egypt (Fig. 1), has produced an extensive fossil record of distinctive and diverse clades of terrestrial mammals. The Jebel Qatrani Formation has been interpreted as a primarily fluvial deposit and is characterized by abundant weathering horizons and root traces indicative of a tropical monsoon climate regime (*Bown & Kraus, 1988*). The formation has been separated into an upper sequence and a lower sequence, with the division between the two sequences marked by a cliff-forming unit called the “Barite Sandstone”. The fossils described here come from the type locality for *T. lavocati* (Quarry E) and the older Quarry L-41 (Fig. 1). Based on the preferred paleomagnetic correlation of *Seiffert (2006)*, L-41 is estimated to be between 33.9–35 Ma, while Quarry E is estimated to be between 31 and 33.2 Ma. L-41 is located approximately 48 m above the base of the Jebel Qatrani Formation and just below a major unconformity that has been identified as the most likely site of near-shore erosion during early Oligocene sea level fall (*Seiffert, 2006*). L-41 is a well-consolidated deposit that is dominated by clay and post-depositional salt, contrasting with the fine-
medium-grained sandstones of younger Fayum quarries (Bown & Kraus, 1988). L-41 is the richest Paleogene vertebrate site in Africa and preserves hundreds of thousands of fossils such as bats (Gunnell, Simons & Seiffert, 2008), rodents (Sallam, Seiffert & Simons, 2011, 2012; Sallam & Seiffert, 2016), and primates (Simons, 1990, 1997; Simons et al., 2001; Seiffert et al., 2018). Quarry E is located approximately 90 m above the base of the Jebel Qatrani Formation and is composed of unconsolidated gravelly sandstones that were deposited as point bars in large meandering rivers (Simons & Rasmussen, 1990). Quarry E has yielded a great diversity of vertebrate fossils such as rodents (Wood, 1968), birds (Stidham & Smith, 2015), and anthropoids (Simons, 1962; Simons & Kay, 1983).

MATERIALS & METHODS

Taxonomy
The electronic version of this article in Portable Document Format (PDF) will represent a published work according to the International Commission on Zoological Nomenclature (ICZN), and hence the new names contained in the electronic version are effectively published under that Code from the electronic edition alone. This published work and the nomenclatural acts it contains have been registered in ZooBank, the online registration system for the ICZN. The ZooBank Life Sciences Identifiers (LSIDs) 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:79E437BD-03EA-42BD-B341-D77EF2AC37F7. The online version of this work is archived and available from the following digital repositories: PeerJ, PubMed Central SCIE and CLOCKSS.

Dental cusp and crest nomenclature
Terminology follows Marivaux & Boivin (2019) (Fig. 2). Teeth are referred to as I, P, and M (for incisors, premolars, and molars, respectively), with upper and lower teeth designated by superscript and subscript numbers (respectively) (e.g., the second lower molar is referred to as $M_2$).

CT scanning and rendering: μ-CT scans of the original fossils of Talahphiomys lavocati and Qatranimys safrouitus, and of casts of the original fossils of the Dur at-Talah specimens of “T. lavocati”, were collected at either the Duke University Shared Materials Instrumentation Facility or the USC Molecular Imaging Center using a Nikon XT H 225 ST micro-CT scanner. Three-dimensional surface models were constructed using ImageJ and Avizo v. 8 and saved in Stanford “ply” format. Additional surface model manipulation and measurements were conducted in Avizo and MeshLab. Digital models of all specimens scanned as part of this study are available on MorphoSource (see Data S1).

RESULTS

Systematic Paleontology
Class Mammalia Linnaeus, 1758
Order Rodentia Bowdich, 1821
Infraorder Hystricognathi Tullberg, 1899
**Parvorder** Phiomorpha *Lavocat, 1967*

**Family** Incertae sedis

**Subfamily** Phiocricetomyinae *Lavocat, 1973*

*Talahphiomys lavocati*, *Wood, 1968* (Fig. 3; Figs. 5A–5E in *Wood, 1968*)

**Synonymy**

*Elwynomys lavocati* (in part) in *Holroyd (1994*, specimen in lowest frame of her Fig. 4.10)  
*non Elwynomys lavocati* in Lewis and Simons (2001)  
*non Talahphiomys lavocati* in Jäger et al. (2010, Figs. 6K–6V)  
*non Talahphiomys lavocati* in Coster et al. (2012, Fig. 4L)  
*non Talahphiomys lavocati* in Marivaux et al. (2014, Figs. 6E–6F)

**Revised diagnosis**

*T. lavocati* (Figs. 3–5) differs from *Talahphiomys libycus* from Dur at-Talah DT-LOC-1 in having a relatively long dP¹ with a larger paraflexus. The mesoflexus bears a weak
metaloph connecting to the metaconule. *T. lavocati* differs also in lacking a mesostyle, mesolophule, and isolated metaloph on M\(^1\) (the latter being submerged into the posteroloph in *T. lavocati*). *T. lavocati* differs from *Qatranimys safroutus* (new genus and species, see below) (Figs. 3–5) in having inflation of the enamel surrounding the base of the protoconid and to a lesser extent the hypoconid of M\(^1\)–\(^2\), forming an incipient labial cingulid; broad overlap in the size and proportions of M\(^1\) relative to M\(^2\) (see bivariate plot in Fig. 6); a lingually open M\(^2\) metaflexid, with no tall connection of the posterolophid to the entoconid; mesial and lingual margins of the mesiolingual corner of M\(^1\)–\(^2\) that form roughly a 90 degree angle, rather than a relatively obtuse angle; a relatively long dP\(^4\) with a labial margin longer than the lingual margin, a relatively capacious paraflexus, and a relatively lingually placed paracone; and an M\(^1\) that is relatively quadrate in occlusal view. Differs from the “*T. lavocati*” specimens from Dur at-Talah Locality (DT-LOC-2) (Figs. 4–5) in having inflation of the enamel surrounding the base of the protoconid and to a lesser extent the hypoconid of M\(^1\)–\(^2\), forming an incipient labial cingulid; smaller M\(^1\)–\(^2\) with different proportions (being mesiodistally longer than buccolingually broad; see bivariate plot in Fig. 6); a more distally placed dP\(^4\) protoconid, and a deep sulcus between that cusp and the adjacent metaconid and anteroconid; a relatively well developed posterior arm of the metaconid and anterior arm of the entoconid on M\(^1\)–\(^2\), closing the mesoflexid lingually; relatively deep indentations on the crown wall of M\(^1\) mesial to the protocone, forming an incipient anterocingulum; dP\(^4\) that is relatively trapezoidal in occlusal outline, with a shorter anterior arm of the hypocone, larger paraflexus, and incipient connection of the metaloph with the metacone. Differs from the “*T. lavocati*”
specimens from Locality ZR5 (Zallah Oasis) in having M<sub>1-2</sub> that are relatively broad compared to mesiodistal length (see bivariate plot in Fig. 6), and in having a relatively well-developed posterior arm of the metaconid and anterior arm of the entoconid on M<sub>2</sub>, closing off the mesoflexid lingually. *T. lavocati* differs from *Phiocricetomys minutus* (Wood, 1968; Fig. 16) in retaining M<sub>3</sub> and in having quadrangular (rather than mesiodistally elongate) lower dP<sub>4</sub>-M<sub>2</sub> with less bulbous cusps and well-developed metalophulid I, ectolophid and posterolophid crests, as well as lingual closure of the trigonids through connection of the posterior arm of the metaconid and anterior arm of the entoconid.

**Holotype**
CGM 26903, right mandible with dP<sub>4</sub>-M<sub>3</sub> (early Oligocene Quarry E, Jebel Qatrani Formation, Egypt).

**Revised hypodigm**
YPM 18011, left mandible with dP<sub>4</sub>-M<sub>1</sub>; YPM 18057, left mandible with dP<sub>4</sub>-M<sub>2</sub>; DPC 4275, left maxilla with dp<sup>4</sup>-M<sub>1</sub>; DPC 5057, right mandible with dp<sub>4</sub>-M<sub>2</sub> and incisor; DPC 8181, left mandible with M<sub>1-2</sub> and incisor (See Table 1 & Data S1).
Description of new specimens

The mandible (Fig. 7) is fully hystricognathous, owing to the placement of the angular process lateral to the long axis of the incisor, leaving a distinct groove between the angular process and the incisor alveolus. On the lateral aspect of the mandible, the mental foramen is relatively small, roughly oval in shape and situated directly under the mesial part of dP<sub>4</sub> (Fig. 7H). The masseteric fossa is defined dorsally by a weakly-developed dorsal masseteric ridge that fades below the anterior part of M<sub>1</sub>. The ventral masseteric ridge is well developed and originates below the anterior part of M<sub>1</sub> and continues...
posteroventrally towards the angular process. The dorsal and ventral ridges meet inferior to the distal part of dP₄. The posterior portion of the ascending ramus is not preserved in any of the specimens, so the morphology of the coronoid, condylar and angular processes are not known; however, the coronoid process seems to be higher than the tooth row, rising lateral to the third molar and leaving a deep fossa. On the medial
surface of the mandible, the angular process originates ventral to M₃ (Fig. 7I). There are some nutrient foramina scattered on the corpus. The outline of the ventral surface of the corpus is convex, with the deepest point being beneath the distal portion of the diastema. The diastema is slightly deeper than the alveolar plane and makes up about one-third of the tooth row. The mandibular symphysis is unfused and extends posteriorly to the position below the dP₄. The symphysis has an anterior part that is broader than its posterior portion (Fig. 7I).

DPC 8181 shows that the lower incisor’s alveolus extends posteriorly to end behind the tooth row. The tooth is covered anteriorly by smooth enamel that extends to the labial and lingual surfaces. On the labial side of the incisor, the enamel covers about one-third, but only a quarter of the medial side. The pulp cavity is exposed as a small slit situated at the middle of the incisor.
A well preserved dP₄ is implanted in DPC 5057 (Fig. 7F). Its trigonid is narrower than the talonid, and the crown is longer mesiodistally than buccolingually broad. The crown displays five major bulbous cusps (metaconid, protoconid, entoconid, hypoconid and hypoconulid) that are more or less equal in size. On the mesial portion of the crown, the protoconid is positioned distolabially with respect to the metaconid, leaving a somewhat broad mesial shelf; within this shelf there is a short low cristid that is protruding from an incipient anteroconid to reach the metaconid. The anteroconid is placed mesial to the protoconid. The crown lacks a metalophulid I, and there is a notch separating the protoconid from the metaconid and anteroconid that continues into the central basin. The entoconid is placed mesial to the hypoconid, and they are linked by the hypolophid and a well-developed anterior arm of hypoconid. The junction of these two cristids is also the point of connection of a relatively short ectolophid. There is no trace of a mesostylid. On the very distal portion of the crown at its midpoint, there is a well-developed hypoconulid. This hypoconulid connects to the hypoconid via a short posterolophid but does not reach the entoconid. The posterior basin is open lingually.

The M₁ (Figs. 7F–7G) is roughly square in shape, and has five distinct major cusps (protoconid, metaconid, hypoconid, entoconid, and a well-developed hypoconulid). The labial cusps are larger in size and slightly displaced distally with respect to the lingual cusps. Three transverse cristids (metalophulid I, hypolophid and posterolophid) and one longitudinal cristid (ectolophid) are present, the latter of which meets an incipient posterior arm of the protoconid. Along the labial portion of the tooth the base is inflated, particularly around the protoconid, forming an incipient cingulum. The metalophulid I delimits the mesial wall of the crown and connects the mesiolingual side of the protoconid with the labial side of the metaconid. The anterior basin is wide and closed by a low lingual wall formed by the posterior arm of the metaconid and the anterior arm of the entoconid. The ectolophid is complete and courses from the protoconid to reach the junction of the hypolophid and the anterior arm of the hypoconid. The posterolophid runs from the hypoconid to meet the hypoconulid. The posterolophid has a robust labial portion, whereas its lingual portion tapers towards the entoconid. The hypoflexid is large and deep.

The second lower molar (M₂) is similar in morphology to the first lower molar but differs in having more robust and well developed lophids (Figs. 7F–7G). Furthermore, the lingual wall, formed by the posterior arm of the metaconid and the anterior arm of the entoconid, is relatively taller than in M₁. The labial cusps (protoconid and hypoconid) are larger and more labial in position than those on M₁, with relatively less basal inflation of the enamel. The hypoconulid is weakly developed and there is no depression between the hypoconid and the hypoconulid.

There is only one maxillary specimen in the hypodigm (DPC 4275), a fragment of a left maxilla with dP₄ and M₁ and an alveolus for dP₃ (Figs. 7A–7C). The infraorbital foramen is only partially preserved, but it is clearly broad and hystricomorphous. The margins of the incisive foramen cannot be traced with confidence due to damage in this area. The ventral ramus of the zygomatic process is thick. On the ventral view of the
maxilla at the base of the ventral ramus of zygomatic process there is a ridge defining a broad fossa for the attachment of the superficial masseter.

The dP⁴ is somewhat trapezoidal in shape and broader labially than lingually (Fig. 7C). The crown of the tooth bears four major cusps (paracone, metacone, protocone and hypocone) and a well-developed metaconule. The protoloph is a well-developed, transverse cristid which courses labially from the submerged paracone and thins toward the labial portion of the protocone. The anteroloph is lower than the protoloph and runs from the labial aspect of the protocone to terminate near the mesial aspect of the paracone, delimiting a large paraflexus. There is no mesostyle. A small centrally-placed metaconule is connected to the hypocone via a short but robust anterior arm of the latter cusp. A very thin and incomplete mure is faintly visible. There is a remnant of a metaloph that turns mesially from the metacone to meet the metaconule, delimiting a deep but small fovea (posterofossette) on the distal portion of the tooth. From the hypocone, the posteroloph runs labially to connect to the base of the metacone. The posteroloph is relatively weakly developed when compared with the anteroloph. The labial wall is formed by a long posterior arm of the paracone that terminates at the base of the metacone. The hypoﬂexus is deep and no endoloph is present. There is a very small accessory cusp (enterostyle?) in the distolingual portion of the sinus. The M¹ has the same basic occlusal configuration as dP⁴, but is larger, is transversely broader, has no mure, and has relatively well-developed cusps, including an incipient anterostyle (Fig. 7C).

**Comparison of T. lavocati with other possible phiocricetomyines**

The phylogenetic analyses of *Marivaux & Boivin (2019)* placed ten taxa other than *T. lavocati* within Phiocricetomyinae, significantly expanding the possible membership of the subfamily (Table 2). Here we expand our comparisons with these possible relatives of *T. lavocati*.

*T. lavocati* differs from *Birkamys* in having a more posteriorly placed mental foramen; smaller metaconids and entoconids relative to protoconids and hypoconids; relatively well developed dP₄-M₂ hypoconulids; a more distally placed dP₄ protoconid; and in lacking M₁–₂ anterior cingulids. The dP⁴-M¹ of *T. lavocati* differ from those of *Birkamys* in having

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**Table 2** Possible members of Phiocricetomyinae based on the phylogenetic results of *Marivaux & Boivin (2019).*

| Age               | taxon                        | Locality      | Reference                        |
|-------------------|------------------------------|---------------|----------------------------------|
| late middle Eocene| “Protophiomys” tunisiensis   | Tunisia       | Marivaux et al. (2014)           |
| late Eocene       | *Talahlphiomys libycus*      | Libya         | Jaeger et al. (2010)             |
| latest Eocene     | *Birkamys korai*             | Egypt         | Sallam & Seiffert (2016)         |
| latest Eocene     | *Mubhammys vadumensis*       | Egypt         | Sallam & Seiffert (2016)         |
| earliest Oligocene| *Mubhammys atlanticus*      | Moroco        | Marivaux et al. (2017)           |
| earliest Oligocene| *Neophiomys minutus*         | Moroco        | Marivaux et al. (2017)           |
| earliest Oligocene| *Phenacophiomys occidentalis*| Morocco       | Marivaux et al. (2017)           |
| early Oligocene   | *Neophiomys paraphiomyoides* | Egypt and Libya| Coster et al. (2012) and Wood (1968) |
| early Oligocene   | *Neophiomys dawsonae*        | Libya         | Coster et al. (2012)             |
| early Oligocene   | *Phiocricetomys minutus*     | Egypt         | Wood (1968)                      |

Al-Ashqar et al. (2021), PeerJ, DOI 10.7717/peerj.12074
a distinct metaconule (rather than being submerged into the mure); more labially placed protocones and hypocones, particularly on dP^4; relatively trenchant posterolophs; and in lacking a distinct metaloph on M^1. T. lavocati differs from Neophiomys minutus in having little or no development of the metalophulid II on M_{1-2}; well-developed dP^4 metaconule; a relatively large dP^4 paraflexus; a relatively well developed posterior arm of the paracone on dP^4; and a relatively weak and more labially placed mure on dP^4.

T. lavocati differs from Neophiomys dawsonae in being relatively small and in having little or no development of the metalophulid II on M_{1-2}; a well-developed metaconule on M^1; and no M^1 mure or metaloph. T. lavocati differs from Neophiomys parapihyoides in being smaller; in having an anteroconid on dP_4, and in lacking any development of metalophulid II on the lower molars. The M^1 of T. lavocati differs in having a well-developed metaconule, and in lacking a mure and a distinct metaloph. T. lavocati is smaller than Mubhammys, and differs in having a more posteriorly placed mental foramen; a more distally placed dP_4 protoconid and relatively short dP_4 (but more trenchant) ectolophid; lingually closed M_{1-2} mesoflexids; relatively well developed dP^4-M^1 metaconules; no mesostyles on dP^4-M^1; and a metaloph, anterostyle, and more distinct anteroloph on dP^4. T. lavocati is smaller than Phenacophiomys occidentalis and differs in having labial cusps larger in size than the lingual cusps; a more distally placed dP_4 protoconid and relatively short dP_4 ectolophid; and no anterior cingulid on dP_4-M_1.

T. lavocati further differs in lacking any connection between the metaloph and the metaconule on M^1 and in having no development of metalophulid II on dP_4-M_2.

T. lavocati differs substantially from “Protophiomys” tunisiensis in having more bunodont cusps and more robust crests; lingually closed mesoflexids; no development of metalophulid II on dP_4-M_2; no anterior cingulids on dP_4-M_2; no connection between the metaloph and metaconule on M^1; and no development of a mesostyle on M^1.

Finally, compared to early Oligocene Phiomys andrewsi (the type species of Phiomys), T. lavocati is smaller in size, and has a mental foramen at the level of the premolar rather than mesial to it; a relatively short ectolophid; more bulbous cusps; and lacks a mesoconid, metalophulid II, posterior arm of protoconid, and anterior cingulid. In the upper dentition, T. lavocati differs from P. andrewsi in having a well-developed metaconule and anteroloph, and in lacking a mesolophule; on dP_4 the protocone and the hypocone are more labially placed.

**Remarks**

Based on morphological and metric grounds (Figs. 3–6), we are able to demonstrate that T. lavocati is restricted to Quarry E in the lower sequence of the Jebel Qatrani Formation of northern Egypt. The specimens referred to T. lavocati from the Libyan sites DT-LOC-2 and ZR5 by Jaeger et al. (2010) and Coster et al. (2012), respectively, do not belong to that species and are now in need of revision. Among other things, with the new information provided by Qatranmys safrouus (see below), we do not consider the specimen identified by Jaeger et al. (2010; p. 206; Fig. 6M) as an M^2 of T. lavocati (DT-2-103) to be an M^2 (we identify it as an M^1) or to belong to T. lavocati.
Family *Incertae sedis*

Subfamily Phiocricetomyinae *Lavocat, 1973*

*Qatranimys*, new genus [urn:lsid:zoobank.org:act:3866127A-97CF-43A1-B7DC-F042805AE197](urn:lsid:zoobank.org:act:3866127A-97CF-43A1-B7DC-F042805AE197) (Figs. 8–10)

Type and only species

*Qatranimys safroutus*, new species [urn:lsid:zoobank.org:act:65B76CA2-42C9-4674-8C2A-A146994DAD3B](urn:lsid:zoobank.org:act:65B76CA2-42C9-4674-8C2A-A146994DAD3B) (Figs. 3 and 5)

Etymology

Combination of ‘qatrani’, Arabic for tar, and in reference to the Jebel Qatrani (“tar hills”) region where the species is found, and ‘mys’, Greek meaning mouse.

Diagnosis

As for the type and only species.

*Qatranimys safroutus*, new species [urn:lsid:zoobank.org:act:[ID]](urn:lsid:zoobank.org:act:[ID]) (Figs. 3 and 5)

Etymology

From colloquial Egyptian Arabic *safrout* (سَفْرَوت), meaning tiny.

*Diagnosis*

*Q. safroutus* differs from *T. lavocati* in having a relatively short dP$_4$ with a metalophulid I; no inflation of the enamel surrounding the bases of the protoconid and hypoconid on M$_1$–2

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**Figure 8** DPC 17813, complete left mandible with I and P$_2$–M$_3$ of *Qatranimys safroutus*, new genus and species, from Quarry L-41. (A) Lateral, (B) medial, (C) occlusal, and (D) ventral views; and (E) occlusal surface. Each of the grey and/or white division in the scale bar refers to 1 mm.  
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and no incipient cingulid around the M$_{1-2}$ protoconids); an M$_{1}$ that is, on average, smaller than M$_{2}$ (see bivariate plot in Fig. 6); a lingually closed M$_{2}$ metafl, with a relatively high connection of the posterolophid to the entoconid; mesial and lingual margins of the mesiolingual corner of M$_{1-2}$ that form a relatively obtuse angle, rather than a ~90 degree angle; a relatively short dP$_{4}$ with a labial margin approximately equal in length to the lingual margin, and with a smaller parafl, a more distally placed hypocone, and a relatively buccally placed paracone; and an M$_{1}$ with a relatively narrow distal moiety.

The M$_{1}$ of Q. safroutus differs from the possible M$_{1}$ of T. libycus (identified as an M$_{2}$ by Jaeger et al., 2010; p. 203; Fig. 5X) in lacking an anterior cingulid, and in having a more obtuse angle between the mesial and lingual margins of the crown and a shorter posterior arm of the metaconid. The M$_{1}$ of Q. safroutus differs from that of T. libycus in lacking both a mesostyle and a long mesolophule that meets the buccal margin of the tooth, and in having a larger anterostyle, a metaloph that is curved toward the metaconule, and a lingually positioned metacone. The M$_{2}$ of Q. safroutus differs from the M$_{2}$ of T. libycus in having a low mure, a relatively well developed metaconule, and a relatively small metacone. Q. safroutus differs from the “T. lavocati” specimens from Dur at-Talah Locality (DT-LOC-2) (Figs. 4–5) in having a more distally placed dP$_{4}$ protoconid, with a distinct metalophulid I; M$_{1-2}$ with different proportions (being mesiodistally shorter than buccolingually broad; see bivariate plot in Fig. 6); a relatively well developed
Figure 10  DPC 10300 and DPC 16815, crania of *Qatranimys safroutus* (new genus and species) from Quarry L-41. (A) Lateral view of right side of DPC 10300; (B) inferior view of DPC 10300; (C) dorsal view of DPC 16815; (D) dorsal view of DPC 10300; (E) inferior view of DPC 10300; (F) lateral view of right side of DPC 16815. Each of the grey and/or white divisions in the scale bar represents 1 mm.

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posterior arm of the metaconid and anterior arm of the entoconid on M₁–₂, closing off the mesoflexid lingually; no anterior cingulid on M₁–₂; a more obtuse angle between the mesial and lingual margins of M₁; dP⁴ that is relatively broad in occlusal outline, with a more restricted paraflexus; a connection of the dP⁴ metaloph with the metaconule; a mure connecting to the protoloph on dP¹; M¹ that is relatively broad, with a connection between the metacone and metaconule, and an incipient mure.

**Holotype**
CGM 83743, right mandible with P₄-M₃ (Fig. 3F).

**Hypodigm**
The holotype; DPC 8825, right mandible with P₄ and M₁–₃; DPC 10300, rostrum with right and left upper incisors and dP³-M²; DPC 14243, partial right edentulous maxilla; DPC 16815, cranium with two incisors, right dP⁴-M² and left dP³-M²; DPC 20965, right maxilla with dP³-M²; DPC 10710, left mandible with P₄-M₃; DPC 11345, left mandible with M₃; DPC 14056, left mandible with M₁–₃; DPC 14187, right mandible with dP₄-M₂ and incisor; DPC 14393, left mandible with P₄-M₂; CGM 83743, right mandible with P₄-M₃; DPC 17813, right mandible with P₄-M₃ and incisor; DPC 17947, left mandible with dP₄-M₂ and incisor; DPC 20659, right mandible with M₁–₃; DPC 21818, left M₂–₃ (See Table 3 & Data S1).

**Type locality**
Locality 41, Jebel Qatrani Formation, Fayum Depression, Egypt.

**Description**
The mandible is similar to those of other Fayum hystricognaths in having an angular process that is placed lateral to the plane of the incisor and tooth row, leaving a wide groove between the angular process and the incisor alveolus in ventral view; this area provides the insertion for the *pars reflexa* of the superficial masseter muscle (*Hautier & Saksiri, 2009*). The ascending ramus is posteriorly inclined and originates lateral to the alveolar plane near the base of the M₁ and M₂ (as in DPC 17813, Fig. 8). The tip of the coronoid process is not preserved. The horizontal ramus is robust and ventrally convex, with its deepest point being below the P₄. The diastema is slightly deeper than the alveolar plane. On the lateral surface, the mental foramen is relatively small and varies from being oval to round in outline; it is situated directly under the mesial part of P₄.

The masseteric fossa is deep, posteriorly broad and tapering anteriorly to terminate below the M₁. The dorsal masseteric ridge is weakly developed and crosses the dorsal surface of the horizontal ramus under the posterior part of M₁. There are some nutrient foramina scattered on the horizontal ramus. The ventral masseteric ridge is well developed, originates laterally from the area beneath the anterior part of M₁, and continues posteroventrally towards the angular process, which is not preserved (Fig. 8A). On the medial side, the angular process initiates beneath the area of M₃. The mandibular foramen is not preserved. The symphysis is partially preserved in DPC 14056 and DPC 17947.
The lower incisor is well preserved in many specimens. The tip of the incisor projects above the tooth row and extends distally to terminate posterior to $M_3$ (Fig. 8B). It is covered anteriorly by smooth enamel that extends to the labial and lingual sides, covering about one third and one fourth of the labial and lingual sides of the incisor, respectively, as seen in all Fayum hystricognaths. On the occlusal surface, the pulp cavity is preserved, has an oval shape, and is posteriorly placed.

The $dP_4$ is only known from two specimens in the hypodigm (DPC 17947 and DPC 14187) (Figs. 3C and 3D). The $P_4$ is present in several specimens, indicating that $dP_4$ is replaced by the permanent premolar. The $dP_4$ is generally pear-shaped in outline and longer mesiodistally than labiolingually. The occlusal pattern is trilophodont (with metaenulophulid I, hypolophid and posterolophid) and displays five major cusps (metaconid, protoconid, entoconid, hypoconid and hypoconulid) that are more or less equal in size and

### Table 3

| Specimen | Side | Upper teeth | $dP^3$ | $dP^4$ | $M^1$ | $M^2$ | $M^3$ |
|----------|------|-------------|-------|-------|-------|-------|-------|
|          |      |             | Length | Width  | Length | Width  | Length | Width  |
| DPC 10300| right| 0.40        | 0.43   | 0.98   | 1.03   | 1.14   | 1.23   | 1.14   | 1.32   | –     | –     |
|          | left | 0.37        | 0.40   | 0.98   | 1.00   | 1.12   | 1.23   | 1.13   | 1.34   | –     | –     |
| DPC 16815| right| –           | –      | 0.96   | 0.97   | 1.11   | 1.20   | 1.05   | 1.22   | –     | –     |
|          | left | 0.40        | 0.43   | 0.98   | 0.98   | 1.10   | 1.20   | 1.04   | 1.22   | –     | –     |
| DPC 20965| right| 0.56        | 0.60   | 1.34   | 1.29   | –      | –      | –      | –      | –     | –     |
|          |      |             |        |        |        |        |        |        |        |        |        |
| No.      |      |             | 4      | 5      | 4      | 4      | 0      |        |        |        |        |
| Mean     |      |             | 0.433  | 0.465  | 1.048  | 1.054  | 1.117  | 1.215  | 1.09   | 1.275  | –     | –     |

| Specimen | Side | Lower teeth | $dP_4$ | $P_4$ | $M_1$ | $M_2$ | $M_3$ |
|----------|------|-------------|-------|-------|-------|-------|-------|
|          |      |             | Length | Width  | Length | Width  | Length | Width  |
| CGM 83743| right| –           | –      | 1.03   | 0.90   | 1.17   | 1.08   | 1.19   | 1.08   | 1.24   | 1.05   |
| DPC 8825 | right| –           | –      | 0.95   | 0.88   | 1.14   | 1.07   | 1.19   | 1.05   | –      | –      |
| DPC 10710| left | –           | –      | 0.96   | 0.91   | 1.16   | 1.09   | 1.13   | 1.08   | 1.22   | 1.14   |
| DPC 11345| left | –           | –      | –      | –      | –      | –      | –      | –      | –      | –      |
| DPC 14056| left | –           | –      | –      | –      | –      | –      | 1.14   | 1.1    | 1.24   | 1.16   | 1.28   | 1.17   |
| DPC 14187| right| 1.01        | 0.78   | –      | –      | 1.11   | 0.99   | 1.22   | 1.17   | –      | –      |
| DPC 14393| left | –           | –      | 0.98   | 0.96   | 1.13   | 1.11   | 1.15   | 1.13   | –      | –      |
| DPC 17813| right| –           | –      | 0.91   | 0.91   | 1.09   | 1.04   | 1.15   | 1.12   | 1.09   | 1.09   |
| DPC 17947| left | 0.99        | 0.76   | –      | –      | 1.04   | 1.0    | 1.18   | 1.12   | –      | –      |
| DPC 20659| right| –           | –      | –      | –      | 1.08   | 1.16   | 1.20   | 1.17   | 0.99   | 1.0    |
| DPC 21818| left | –           | –      | –      | –      | –      | –      | 1.21   | 1.11   | 1.1    | 0.98   |
| No.      |      |             | 2      | 5      | 9      | 11     | 7      |        |        |        |        |
| Mean     |      |             | 1.0    | 0.770  | 0.966  | 0.912  | 1.118  | 1.071  | 1.191  | 1.127  | 1.143  | 1.061  |
are of the same height. On the mesial portion of the tooth the protoconid is distal in position with respect to the metaconid. The protoconid and metaconid are connected via the metalophulid I which is arc-shaped and runs mesiolingually, delimiting the posterior wall of a broad mesial shelf. On this shelf, there is a prominent isolated anteroconid mesiolabial to the metaconid. The lingual wall between the metaconid and the entoconid is low, leaving the wide and deep mesoflexid closed lingually. In DPC 17947 (Fig. 3C), there is an incipient mesostylid near the mesial aspect of the entoconid. The ectolophid is short, low relative to the cusp height, and joins the protoconid to the junction of the anterior arm of the hypoconid and the hypolophid. The anterior arm of the hypoconid is short and connects to a well-developed hypolophid. The latter extends lingually to connect with the entoconid. On the distal portion of the crown, the entoconid is placed mesially with respect to the hypoconid. The hypoconulid is a well-developed cusp on the middle of the posterolophid forming the very distal portion of the tooth. The posterolophid runs from the hypoconid to end and taper distolabial to the base of entoconid, leaving the posterior basin opened lingually.

Five specimens preserve the P₄ (CGM 83743, DPC 8825, DPC 10710, DPC 14393, and DPC 17813). The tooth is relatively shorter and broader when compared with the dP₄. It is roughly rectangular to square in shape, with the talonid slightly wider than the trigonid and bearing four main cusps (metaconid, entoconid, protoconid, and hypoconid). The metaconid and entoconid are placed roughly transverse to the protoconid and hypoconid, respectively. The metalophulid I is complete in DPC 8825 and DPC 17813, but in CGM 83743, DPC 10710, and DPC 14393 the metalophulid I is interrupted by a narrow notch lingual to the protoconid. In DPC 14393 (Fig. 3G) the metaconid and the protoconid are more bulbous and there is an incipient anteroconid mesial to the protoconid. There is no hint of the posterior arm of the protoconid in any of the specimens. In CGM 83743 (Fig. 3F) and DPC 14393 (Fig. 3G), the anterior basin of the P₄ is generally large and open linguallly via a deep notch on the lingual wall. The other specimens have a low lingual wall closing the mesoflexid. The hypoconid has a well developed anterior arm that is connected to a well developed ectolophid. The hypolophid shows considerable variability—in DPC 8825, DPC 14393 and DPC 17813, the hypolophid arcs distolabially to form a direct connection with the posterolophid, delimiting a small fovea, while in DPC 10710 (Fig. 3E) the hypolophid is complete, connecting to the anterior arm of the hypoconid. In CGM 83743 (Fig. 3F), the hypolophid extends from the entoconid toward the anterior arm of the hypoconid but it ends abruptly at the center of the tooth, connecting the posterior basin with the central basin via a notch. On the distal portion of the crown, the hypoconulid varies from being small to distinct and is subsumed into a posterolophid that terminates at the distal aspect of the entoconid, delimiting the posterior margin of the tooth.

The M₁ is roughly rectangular in shape, with almost all specimens being slightly longer than wide. The tooth has five bulbous cusps (protoconid, metaconid, entoconid, hypoconid and hypoconulid) and three transverse cristids in the occlusal pattern (metalophulid I, hypolophid, and posterolophid). The metaconid is placed transverse to the protoconid, while the entoconid is situated mesial to the hypoconid. The metalophulid
II is not present, with only a small knob protruding from the lingual face of the protoconid. The metalophulid I runs labially from the metaconid to reach the mesiolingual side of the protoconid. In DPC 14056 (Fig. 3I), the metalophulid I is interrupted by a small notch. The hypolophid is well developed and attaches to the anterior arm of the hypoconid near that crest's junction with the ectolophid. The ectolophid is well developed and situated near the middle, or just labial to the middle, of the tooth. The posterolophid runs distolingually from the hypoconid and terminates at the base of the distal aspect of the entoconid, delimiting the posterior margin of the tooth. The hypoconulid is well developed and more or less the same size as the hypoconid. The two lingual cusps (metaconid and entoconid) are relatively small compared to the labial cusps. The mesolophid is broader and delimited by a lingual wall formed by the posterior arm of the metaconid that reaches the mesial aspect of the entoconid. The hypoflexid is transversely wide and deep.

The M2 is quadrangular with a rounded posterior portion. The occlusal surface of the M2 is similar to that of the M1 but differs in being slightly larger in size, and having more developed transverse lophids, a broader trigonid relative to the talonid, a taller posterolophid and lingual wall of the trigonid, a less distinct hypoconulid, and a more lingually placed metaconid.

The M3 is preserved in six specimens (DPC 10710, DPC 11345, DPC 14056, CGM 83743, DPC 17813 and DPC 20381). Some specimens are triangular in shape, with a talonid that is much narrower than the trigonid. Most M3s are smaller in size than M2s, however in two individuals (DPC 10710 and DPC 11345) the M3 is similar in size to the M2. Otherwise, the M3 has a very similar occlusal pattern to that of M1 and M2. The metalophulid II varies from being very short to absent. On DPC 11345, the metalophulid II of M3 runs lingually to reach the middle of the mesolophid. The hypoconulid is submerged into the short posterolophid, delimiting the distal lobe of the crown.

The upper incisors are shorter and more highly arched than those of the mandible. They extend posteriorly to terminate just anterior to the dP3. The incisors are covered by smooth enamel which extends labially and medially to cover only one third and one fourth of both sides respectively. In the occlusal surface, in the middle of the dentine layer there is a pulp cavity with a slit shape.

The dP3 is well preserved in three specimens (DPC 16815, DPC 10300 and DPC 20965). It is a very small simple round tooth in occlusal view, with one main cusp that abuts the mesial surface of dP4. On DPC 10300 there is a small accessory cusp.

The dP4 varies from being square to slightly more trapezoidal in outline. The crown bears four major cusps (paracone, metacone, protocone and hypocone), all of which are more or less the same size and height. The anterostyle is small and situated mesial to the protocone, and is connected to that cusp by a short crest that runs from the mesiolingual part of the protocone. The anteroloph extends mesiolabially from the anterostyle and terminates near the mesial base of the paracone. The protoloph is a well-developed transverse crest. There is no hint of a parastyle, mesostyle, or mesolophule. The metaloph is short and turns mesiolingually from the metacone to meet the metaconule. The metaconule is situated near the center of the crown and is connected to
the hypocone by a well-developed anterior arm of the hypocone. A mure is present and meets the protoloph labial to the protocone. The posteroloph is relatively low and runs labially from the hypocone to delimit the posterior margin of the tooth and connects with the distal aspect of the metacone. The labial wall is complete and relatively low as it does not reach the height of the two labial cusps (paracone and metacone).

The M\(^1\) is very similar in occlusal configuration to that of dP\(^4\), but is larger, relatively broad labiolingually, and has relatively well-developed lophs and cusps, including a relatively large anterostyle and a metacone that is relatively small and lingually positioned when compared to the paracone. The metaconule is weakly developed and the connection between the metacone and metaconule is either very faint or absent. The paraflexus is relatively small when compared with that of dP\(^4\). The M\(^2\) is broader than M\(^1\) and has a similar occlusal configuration, but the distolabial corner of the tooth is much different in having a reduced and somewhat crestiform metacone that lacks any hint of a metaloph; together with the anterior arm of the metacone and the posteroloph, the metacone encloses a large fossa comprised of a broadly open mesoflexus + posteroflexus.

The only other hystricognath crania from the Paleogene of Africa are also from Quarry L-41, so the description of the new skull elements of Qatranmys (Figs. 9 and 10) is based on comparison with the sympatric and synchronous Gaudeamus (Sallam, Seiffert & Simons, 2011), Acritophiomys (Sallam, Seiffert & Simons, 2012) and Birkamys (Sallam & Seiffert, 2016). As with most fossils from L-41, the new specimens are compressed and bear numerous surface cracks and displacements due to severe postmortem distortion.

We figure as much as is possible through volume rendering of the skulls, using high-resolution micro-CT scans with minimal physical preparation of these small and fragile specimens. Four crushed cranial specimens have been recovered (DPC 10300, DPC 14324, DPC 16815 and DPC 20956). DPC 10300 is dorsoventrally crushed and includes most of the front of the cranium, including the premaxillae with two upper incisors, the frontals, both maxillae and the entire dentition aside from both M\(^3\); DPC 16815 is mediolaterally crushed and preserves the snout (premaxillae and nasals), the maxillae, the frontals and the parietal in addition to complete dentition aside from both M\(^3\); DPC 14243 is a maxilla with roots of dP\(^4\) and alveoli of dP\(^3\); and DPC 20956 is a maxilla that preserves the third and fourth premolars.

The nasal bones can only be seen clearly in DPC 16815, whereas in DPC 10300 they are highly deformed and extensively damaged. In DPC 16815, the paired nasals' articular surfaces with the frontals extend to the level of the dorsal zygomatic ramus, above the level of the dP\(^4\) (as in Gaudeamus). In DPC 10300, the bones extend backward to articulate with the frontals at the level of M\(^1\)-M\(^2\) posterior to the infraorbital foramen (due to post-mortem displacement). The articular relationships between the nasals and the premaxillae are obscured by distortion.

The premaxillae house two upper incisors, form the upper diastema, and contribute to much of the rostrum. The posterodorsal processes of the premaxilla are preserved in DPC 16815 but missing in DPC 10300. On the lateral side of the posterodorsal process, there is a bony keel protruding dorsally. In lateral view the premaxillae decrease in width anteriorly leading to an arched diastema as in Gaudeamus and Acritophiomys.
The premaxilla is bounded posterolaterally by the maxilla. Despite the postmortem distortion, the borders of the incisive foramen can be clearly seen in DPC 10300 (Fig. 10B), showing that it was large and elongate and likely formed an “anterior palatine fenestra” as in Acritophiomys, Birkamys, Gaudeamus, Mubhammys, and Waslamys (Sallam & Seiffert, 2016). The postorbital processes are present but very small, unlike Gaudeamus which has prominent processes oriented laterally and posteriorly; in Qatranimys, the process has a distinct vascular foramen on its underside, observable on both sides of DPC 10300. The approximate outline of both orbits is preserved on both sides of DPC 16815, but details of the orbital mosaic are impossible to determine due to breakage. The jugal is not preserved in any of the specimens. The suture between the lacrimal bone and the posterodorsal process of the premaxilla, and that with the dorsal zygomatic ramus is preserved on both sides of DPC 16815, however the outline of the lacrimal bone cannot be determined with confidence. The dorsal exposure of the lacrimal bears a small foramen on the right and left sides of DPC 16815 (Fig. 9A). The lacrimal foramen is relatively large and is situated in the middle of the bone.

In lateral view, the maxilla contributes to the anterior wall of the orbit, moreover the facial process of the maxilla joins with the posterior portion of the premaxilla to form the lateral wall of the rostrum, and the medial portion of the infraorbital foramen. All of the crania have an enlarged (hystricomorphous) infraorbital foramen, through which the medial masseter muscle pars anterior extends (Hautier & Saksiri, 2009). The infraorbital foramen shows a ventrolaterally rounded outline, as seen on the left side of DPC 16815 (Fig. 9A). The ventral ramus of the zygomatic process of the maxilla extends laterally from the area in front of the dP³ and then arches posteriorly, delimiting the anteroventral portion of the orbital margin. The anteroventral portion of the ventral zygomatic ramus bears a deep fossa for the insertion of the superficial masseter muscle, and, posteriorly, a relatively shallow fossa for the origin of the lateral masseter, as in Gaudeamus and Acritophiomys. The dorsal zygomatic ramus is oriented dorsoventrally in DPC 16815 (Fig. 9A). As in Gaudeamus, the roots of the ventral and dorsal rami extend anteriorly to roughly the same point. It is difficult to trace the original morphology of the palate due to damage, however it appears to be flat and broad. The parietal bones are poorly preserved in DPC 16815, whereas in DPC 10300 they are completely missing (Fig. 10).

**Comparison of Q. safroutus with other possible phiocricetomyines**

Q. safroutus differs from Birkamys in having a more posteriorly placed mental foramen; smaller metaconids and entoconids relative to protoconids and hypoconids; relatively well developed dP4-M2 hypoconulids; a more distally placed dP4 protoconid; and in lacking M1–2 anterior cingulids. The dP4-M2 of Q. safroutus differ from those of Birkamys in having distinct metaconules (rather than being submerged into the mure) and more robust primary cusps. Q. safroutus differs from Neophiomys minutus in having relatively weak development of the metalophulid II on M1–2; well-developed dP4 metaconules; a relatively large dP4 paraflexus; a relatively well developed posterior arm of the paracune on dP4; and a relatively weak and more labially placed mure on dP4. Q. safroutus differs from
Neophiomys paraphiomyoides in having a distinct anteroconid, a broad mesial shelf, and smaller cusps on dP4. The metaconid of dP4 is anteriorly placed relative to the protoconid, rather than being buccolingually opposed. The two latter cusps are connected by a relatively long metalophulid I rather than being closely positioned. The M1–2 of Q. safrouotus differ from those of N. paraphiomyoides in having no development of metalophulid II or anterocingulid in M1–2 and in having a well-developed hypoconulid on M2. The upper molars differ by exhibiting a distinct central metaconule, a relatively weak mure and no development of a metaloph. Q. safrouotus differs from Neophiomys dawsonae in having a well-developed metaconule, labial and lingual walls on upper and lower molars respectively, no development of the metalophulid II and no anterior cingulid. Q. safrouotus is smaller than Mubhammys vadumensis, and differs in having a more posteriorly placed mental foramen; lingually closed mesoflexids; and mesial and lingual margins of the mesiolingual corner of M1–2 that form an obtuse angle and no development of anterior cingulids. In dP4, Q. safrouotus differs in having a more distally placed protoconid; a broad mesial shelf; a well-developed anteroconid; and a relatively short but robust ectolophid and relatively long metalophulid I. In the upper molars, Q. safrouotus differs in having relatively well developed dP4–M1 metalocunes; labially closed parafossete (parafossete); no mesostyles; and a weak mure. Q. safrouotus differs further by having a metaloph, anterostyle, and more distinct anteroloph on dP4. Q. safrouotus differs substantially from Phenacophiomys occidentalis by being smaller, and in having labial cusps larger in size than the lingual cusps; a more distally placed protoconid and relatively short ectolophid in dP4; and no anterior cingulid or any development of metalophulid II on lower molars. In the upper molars, Q. safrouotus differs in lacking mesostyle. Q. safrouotus differs from “Protophiomys” tunisiensis in having a well-developed metalophulid I connecting the protoconid with metaconid on the dP4 rather than being separated. The lower molars of Q. safrouotus differ further in having a lingual wall; no development of the metalophulid II; and no anterior cingulids. Furthermore, M1 shows no development of a mesostyle.

**DISCUSSION**

We detected significant morphological differences between T. lavocati from Fayum Quarry E and the “T. lavocati” specimens from the Libyan sites DT-LOC-2 and ZR5 by Jaeger et al. (2010) and Coster et al. (2012), respectively. Among other features, the bases of the protoconid and the hypoconid in M1–2 of T. lavocati are basally inflated, forming an incipient cingulid. Furthermore, the mesoflexids are closed lingually via the posterior arm of the metaconid and the anterior arm of the entoconid, and there are no mesostyles or mesolophules in the upper molars. The bivariate plot of M1 and M2 proportions (Fig. 6) shows broad overlap in M1 relative to M2 in T. lavocati from Quarry E, and obvious size differences from both the late Eocene (Dur at-Talah) and early Oligocene (Zallah) “T. lavocati”.

Jaeger et al. (2010) suggested that the rodent layers from Dur at-Talah correlate with Chron 18n (~39 and 38 Ma; Upper Bartonian)—several million years older than the
T. lavocati type locality Quarry E, which is placed near the bottom of Chron C12r (Seiffert, 2006), at ~31–33.2 Ma (i.e. Rupelian). If the date that Jaeger et al. (2010) proposed for the Dur at-Talah sites, and identification of T. lavocati at those sites, were correct, it would require the species duration for T. lavocati to be, at a minimum, 7 million years long. Oddly, Jaeger et al. (2010) never acknowledge this discrepancy and its biochronological implications, and instead focused their discussion entirely on evidence that they considered to support an older rather than a younger age for the sites. Sallam & Seiffert (2016) helped to resolve this paradox by using a Bayesian tip-dating approach to estimate the ages of the Dur at-Talah rodents, and found that the sites more likely date to the late Eocene—i.e. intermediate in age between the Fayum BQ-2 and L-41 localities, and several million years younger than the ages proposed by Jaeger et al. (2010). This late Eocene age for the Dur at-Talah rodents has also been accepted by Marivaux et al. (2017).

Given the morphological and metric grounds provided in this work, in addition to stratigraphic range, it can now be demonstrated that the specimens from the Libyan sites actually do not belong in the species lavocati. Hence, the specimens from Dur at-Talah and Zallah placed in “T. lavocati” should be revised in the future. Moreover, the Fayum Quarry E in the lower sequence of the Jebel Qatrani Formation is considered to be the type and only locality of T. lavocati. Nevertheless, we maintain the genus name Talahphiomys as the generic replacement name for “Phiomys” lavocati as this replacement name has priority.

We note interesting points of similarity between T. lavocati and the enigmatic early Oligocene Phiocricetomys (Wood, 1968) that are not seen in the Libyan “T. lavocati” material, namely the development of an incipient labial cingulid around the M1 protoconid in T. lavocati—likely presaging the large and well-developed M1 labial cingulid seen in P. minutus. They also share a deep sulcus separating the isolated dP4 protoconid from the anteroconid and metaconid. Future phylogenetic analyses that take into account all existing Paleogene Afro-Arabian hystricognaths can test the hypothesis that T. lavocati is the exclusive phiocricetomyine sister taxon of Phiocricetomys.

The tiny species Qatranimys safroutus from Quarry L-41 is one of the most diminutive rodent fossils known. Q. safroutus is presumably more primitive than T. lavocati in having no inflation in the labial part of M1–2, and in retaining metalophulid I in dP4. For the first time, the large sample of Q. safroutus allows for an understanding of natural intraspecific variation within a phiocricetomyine species, and bolsters the case for the Libyan “T. lavocati” not being conspecific with either T. lavocati or Q. safroutus.

The phylogenetic position of phiocricetomyines relative to Phiomorpha remains a matter of uncertainty. Many studies placed them outside of the Phiomorpha-Caviomorpha clade (Sallam et al., 2009; Sallam, Seiffert & Simons, 2011; Sallam & Seiffert, 2016; Marivaux & Boivin, 2019) while others placed them as stem phiomorphs (Sallam & Seiffert, 2016, 2019). Future phylogenetic analyses that aim to test these alternate hypotheses can now take into account cranio-mandibular features in addition to dental characters.
CONCLUSIONS
In summary, our analysis of the available material from the Fayum and Dur at-Talah suggests that the Fayum Quarry E is the type and only locality of *T. lavocati*, and that the specimens from Dur at-Talah and Zallah do not belong to this species. Several features suggest that *T. lavocati* may be the exclusive phiocricetomyine sister taxon of *Phiocricetomys*. The vast material of the late Eocene *Q. safroutus* further supports the exclusion of the Libyan “*T. lavocati*” from both *T. lavocati* and *Q. safroutus*, the latter of which shows some primitive features when compared with *T. lavocati*. The dental features of the diminutive *Q. safroutus* further expand our understanding of the interspecific variation that might be expected among phiocricetomyine species. However, in order to develop a more detailed scenario for the membership of Phiocricetomyinae and relationships among known species, a more extensive phylogenetic analysis with both cranio-mandibular and dental characters will be needed.

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INSTITUTIONAL ABBREVIATIONS
CGM    Egyptian Geological Museum, Cairo, Egypt
DPC    Duke Lemur Center Museum of Natural History, Duke University, Durham, North Carolina, USA
YPM    Yale Peabody Museum of Natural History, Yale University, New Haven, United States

ADDITIONAL INFORMATION AND DECLARATIONS

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**Competing Interests**
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**Author Contributions**
- Shorouq F. Al-Ashqar conceived and designed the experiments, performed the experiments, analyzed the data, prepared figures and/or tables, authored or reviewed drafts of the paper, and approved the final draft.
- Erik R. Seiffert conceived and designed the experiments, performed the experiments, analyzed the data, prepared figures and/or tables, authored or reviewed drafts of the paper, and approved the final draft.
- Dorien de Vries performed the experiments, authored or reviewed drafts of the paper, and approved the final draft.
- Sanaa El-Sayed performed the experiments, authored or reviewed drafts of the paper, and approved the final draft.
- Mohamed S. Antar performed the experiments, authored or reviewed drafts of the paper, and approved the final draft.
- Hesham M. Sallam conceived and designed the experiments, performed the experiments, analyzed the data, authored or reviewed drafts of the paper, and approved the final draft.

**Data Availability**
The following information was supplied regarding data availability:
The specimens are deposited in the Egyptian Geological Museum, Cairo, Egypt (CGM) and the Duke Lemur Center Museum of Natural History, Duke University, Durham, North Carolina, USA (DPC).
The specimen digital models are available at MorphoSource:
CGM 83743, https://doi.org/10.17602/M2/M159418;
DPC 4275, https://doi.org/10.17602/M2/M159418;
DPC 5057, https://doi.org/10.17602/M2/M159377;
DPC 8181, https://doi.org/10.17602/M2/M159381;
DPC 8825, https://doi.org/10.17602/M2/M159383;
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