New Paleogene mantises from the Oise amber and their evolutionary importance

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Introduction

Mantises (order Mantodea) are one of the smaller orders of insects with only 2500 described species (Patel and Singh 2016). Moreover, fossils of Mantodea are uncommon and the majority of them cannot be accurately associated with extant lineages of Mantodea (Cui et al. 2018). It is linked to two facts. First, the taxonomy of Mantodea is complicated, incongruent with phylogeny (Svenson and Whiting 2009, but see Schwarz and Roy 2019), characters are difficult to observe and often unpreserved in fossils (Ross 2019), while the general morphology of extant Mantodea is highly convergent in several lineages. Second, Mantodea are large insects, with body sizes ranging between 1 to 17 cm. Consequently, amber preserved mantises are almost exclusively nymphs, with some possible ootheca (Li and Huang 2019), while compressed fossils are isolated wings, where taxonomic characters are generally not present or unusable (Wieland 2013: figs. 414–417).

Therefore, even if there are more than ten currently known genera of Mesozoic fossil Mantodea (Delelòs et al. 2016), only one of them (Ambermantis Grimaldi, 2003) is currently associated with the crown group Mantodea but only in one of the many phylogenetic analyses (Grimaldi 2003). Furthermore Delelòs et al. (2016) considered Ambermantis as an incertae sedis. So it is probable that none of them could reliably be associated with the extant lineages of Mantodea (although undescribed material may help to resolve this problem, e.g., Ehrmann 1999; Xia et al. 2015). All ancient Cenozoic mantises from Palaeocene of Menat (France) are also considered by Cui et al. (2018) as doubtful. These authors reassigned Prochaeradodis enigmaticus Piton, 1940 to the Blattodea, while this taxon was previously considered as one of the oldest known species of crown Mantodea (see Legendre et al. 2015). Cui et al. (2018) also criticized the placement of Arvernineura insignis Piton, 1940 into the Chaeteessidae and concluded that: “there are no formally described and well-assigned fossil crown Mantodea suitable for date calibration” (Cui et al. 2018: 361).

Moreover, “stem sister group” of Mantodea (non-mantises Dictyoptera more related to Mantodea than Blattodea) are also not known. Several candidate taxa have been proposed. Most of them are controversial and/or unreliable: Palaeozoic Paoliida have been proposed as “ancestors” of mantises (Béthoux and Wieland 2009), but Prokop et al. (2014) refuted this hypothesis and considered the Paoliida as sister group of the Dictyoptera on the basis of their wing venation; several Palaeozoic “roachoids” have been considered as sister group(s) of Mantodea, such as Raphidiomimidae and Manipulatoridae (Vršansky and Bechly 2015), or Rap-
Mantodea, 1838

Systematic palaeontology

Class Insecta Linnaeus, 1758

Order Mantodea Burmeister, 1838

Material and methods

A summary on the state of the art concerning the lowermost Eocene Oise amber can be found in Brasero et al. (2009). The middle Paleocene Menat fossil site (Menat Basin, Puy-de-Dôme, France) is a volcanic maar containing rather small palaeolake ca. 1 km in diameter, which contains sedimentary rocks (spongo-diatomite) with remains of diverse aquatic and terrestrial flora and fauna. The composition of faunal and floral remains suggests that this lake was surrounded by a forest and the palaeoenvironment was warm and humid (Wedmann et al. 2018). The age of the Menat fauna is currently estimated as 60–61 Ma (Wappler et al. 2009). The fossils in amber are embedded in small clear pieces. They have been prepared using a diamond disk, examined using a Nikon binocular microscope SMZ 1500. Photographs have been taken with a Nikon D800 with Nikon SMZ25 or a AF-S Micro NIKKOR 60mm f/2.8G ED, and adjusted with Adobe Photoshop CS6. We follow the terminology of Brannoch et al. (2017) for the morphology of the Mantodea.

Family Chaeteessidae Handlirsch, 1926

Included genera: Chaeteessa Burmeister, 1838, Louispitonia nom. nov. (new name for Archaeophlebia Piton, 1940), Arvernineura Piton, 1940, Lithophotina Cockerell, 190 and Megaphotina Gratshev and Zherikhin, 1993.

Genus Louispitonia nom. nov.

Holotype: MNHN-F-R06999, an incomplete hindwing (Piton 1940: fig. 12).

Type locality: South-east of the village of Menat, Menat Basin, Puy-de-Dôme, France.

Type horizon: Middle Paleocene.

Remarks.—Piton (1940) described an incomplete hindwing from the Paleocene of Menat that he named Archaeophlebia enigmatica, and attributed to the ephemeropteran family “Protoneuridae” Piton, 1940 (not the damsselfly family Protoneuridae Tillyard, 1917). Nel and Roy (1996) revised it and attributed this fossil to the Chaeteessidae, but they were not aware that the genus name Archaeophlebia was preoccupied by the extant genus Archaeophlebia Ris, 1909 (Odonata: Libellulidae). Thus we propose a new genus name in replacement of Archaeophlebia Piton, 1940.

Stratigraphic and geographic range.—Only type locality. Chaeteessidae indet.

Figs. 1, 2, 4A.

Material.—MNHN-F.A71139, PA 3309, head and fore legs; MNHN-F.A71142, PA 2420, fore legs. From Le Quesnoy, Chevrère, region of Creil, Oise department, France; lowermost Eocene, Sparnacian, level MP7 of the mammal fauna of Dormaal.

Description.—MNHN-F.A71139: head 1.9 mm long, 2.3 mm wide, triangular, 1.2 times as wide as long; compound eyes globular; antennae filiform, long but incomplete; ocelli not visible; fore coxae without visible spines; fore femora 2.8 mm long, of semicircular section, without distinct carina and probably with six posteroventral spines; fore tibia 1.8 mm long, slightly curved toward posteroventral edge; 14 anteroventral spines, with eighth spine larger than ninth, apical spur not clearly define (Fig. 1). MNHN-F.A71142 is similar to MNHN-F.A71139 (Fig. 2).

Genus Arvernineura Piton, 1940

Type species: Arvernineura insignis Piton, 1940; Menat, Menat Basin, Puy-de-Dôme, France; Middle Paleocene.
Arvernineura insignis Piton, 1940

Figs. 3, 4C.

Material.—MNT NEL1656, a pair of forewings, nearly complete, and some foretibial and forefemoral spines. From South-east of the village of Menat, Menat Basin, Puy-de-Dôme, France; middle Paleocene.

Description.—Preserved length of wing 38.0 mm, width 6.0 mm; fore femoral spines hardly visible and difficult to count; the two fore femora superposed and some spines are detached from raw; six preserved spines on one fore tibia, and 10–11 preserved spines on second with seventh or eighth larger than others; bases of these spines disposed into a curved line, indicating a curvature of fore tibiae (Figs. 3, 4C).

Remarks.—These mantises in amber have a character on the fore tibiae typical of the extant Chaeteessa: a large median spine, larger and thicker than those more basal and distal of antero-ventral raw (Fig. 4). It seems to be absent in all other extant and fossils mantises with known fore legs (TS personal observation; for figures of fossil specimens
This is a putative synapomorphy of the Chaeteessidae. Therefore, it refutes Cui et al. (2018) who doubted about the placement of *Arvenineura* into the Chaeteessidae. Also Cui et al. (2018) stated that the shape of the “stigma” is the same as in *Arvenineura insignis* and in *Cretophotina tristriata* Gratshev and Zherikhin, 1993. However, if we follow the wing nomenclature of Béthoux and Wieland (2009) (also used in Brannoch et al. 2017), we notice that the “stigma” of *A. insignis* covers a part of the RP+M and of the CuA areas, as in extant *Chaeteessa* (Nel and Roy 1996), while that of *Cretophotina tristriata* covers only the CuA area (Gratshev and Zherikhin 1993; Grimaldi 2003). This character does not allow to separate *A. insignis* from the extant *Chaeteessa*. Also the term “stigma” is not appropriate to designate this structure. This structure, a highly sclerotized vein, was referred by Grimaldi (2003) as the “pseudovein”, while in mantises, the stigma generally refers to another sclerotized area of fore wing, often white, smaller and present more basally than the “pseudovein”. Furthermore, some mantises have these two structures well separated on their fore wings. The general shape of the wing of *A. insignis* is also closer to that of *Chaeteessa* rather than of *Cretophotina tristriata* (Nel and Roy 1996; Grimaldi 2003). Lastly, *A. insignis* has a relatively broad area between the fore wing veins ScP and R, as in *Chaeteessa*, while all other extant and most fossil mantises (including *Cretophotina tristriata*) have a narrower area between these veins. Based on all these characters of wing and fore leg, we assume the placement of *A. insignis* into the Chaeteessidae, and its suitability as a crown Mantodea for date calibration.

Although the two specimens of Chaeteessidae of Oise amber could be associated without doubt to this family, we are not able to determinate if they are of the same species. Also, we are not able to assign these specimens to a precise genus of Chaeteessidae due to the lack of characters. *Arvenineura* is closer to these fossils spatially and temporally than *Chaeteessa*, but this does not represent an adequate evidence to associate them to this genus. Consequently, more material is needed to formally describe this (or these) species of Chaeteessidae.

**Stratigraphic and geographic range.**—Only type locality.

### Family Mantoididae Giglio-Tos, 1927

**Included genera:** Mantoida Newman, 1838, Paramantoida Agudelo, 2014, Pseudomantoida gen. nov.

### Genus Pseudomantoida nov.

**ZooBank LSID:** urn:lsid:zoobank.org:act:F11EE9A4-6E7B-4970-A9C1-4A25CA4105DE

**Etymology:** Named after its resemblance to *Mantoida*.

**Type species:** Pseudomantoida extendidera sp. nov., by monotypy, see below.

**Diagnosis.**—The same as for the monotypic type species.

**Pseudomantoida extendidera** sp. nov.

**ZooBank LSID:** urn:lsid:zoobank.org:act:365BADE6-6E90-4154-9A7F-C3B3D6396747

**Fig. 5.**
Etymology: From Latin extend, extend and Greek δέρη, neck; in reference to the extension of the pronotum above the forewings.

Holotype: MNHN-F.A71141 (PA 2378 2/2, the number in the working collection), an adult male with preserved head, prothorax, and fore legs in a hyaline piece of amber.

Type locality: Le Quesnoy, Chevrière, region of Creil, Oise department, France.

Type horizon: Lowermost Eocene, Sparnacian, level MP7 of the mammal fauna of Dormaal.

Material.—We tentatively attribute MNHN-F.A71140 to this taxon, but with doubt (Fig. 6).

Diagnosis.—Small mantises with short and bulky pronotum and fore legs, typical of Mantoididae; fore femora with three posteroventral spines (0–2 or 4 in other Mantoididae, see Agudelo 2014). Lateral margin of pronotum rounded with an extension above bases of forewings.

Description.—Head 3.3 mm long, 3.8 mm wide, approximately as long as wide, triangular with vertex rounded; eyes globular protruding; three large ocelli (probably a male); clypeus without a ridge; antennae long and filiform, third antennomere 2.5 times as long as fourth.

Pronotum: 2.8 mm long, short, as long as wide; saddle-shaped, i.e., with lateral parts bent ventrad; metazona (1.8 mm) two times as long as prozona, but with a long extension above wings, measuring half of its length; outer margin of pronotum raised, forming a lateral carina without spines; lateral margin forming a soft angle in prozona and slightly S-shaped in metazona; central carina on all the length; ventral cervix not visible; scutellum not visible.

Legs: Fore coxae without spines; fore femora 3.8 mm long, thick and short, with a big carina on all length of its dorsal margin and a smaller carina close to the posteroventral raw of spines; three posteroventral spines, at least five same sized anteroventral spines and three discoidal spines; fore tibia with one posteroventral spine in apical position, at least seven anteroventral spines and one apical spur (Fig. 5).

Fig. 4. Fore legs of chaeteessid mantodeans. A. Chaeteessidae gen. at sp. incertae sedis, MNHN-F.A71139, Le Quesnoy, France, Eocene; inner (A1) and outer (A2) side of left fore femur. B. Fore legs of Chaeteessa valida (Perty, 1833), Recent, MNHN collection. C. Habitus of Arvernineura insignis Piton, 1940, MNT NEL1656, Menat, France, Paleocene; photograph with low-angle light (C1), in alcohol (C2). Arrows correspond to tibial major median spines. Scale bars 2 mm.

Fig. 5. Mantoidid mantis Pseudomantoida extendidera gen. et sp. nov., holotype MNHN-F.A71141, Le Quesnoy, France, Eocene. General view from above (A1) and left side (A2).
Fig. 6. Reconstruction of mantidid mantis *Pseudomantoida extendidera* gen. et sp. nov., holotype MNHN-F.A71141.

Remarks.—The head shape of *Pseudomantoida* gen. nov. is quite similar to that of *Mantoida*, more globular than in other “more derived” Mantodea (Wieland 2013: fig. 70). The clypeus has no ridge, as in Chaeteessa, Metallyticus, and *Mantoida* (Wieland 2013: 44, 52). Also the very long antennae, probably longer than the body, is a character of *Mantoida* and Chaeteessa. The short prothorax of *Pseudomantoida* is shared by “*Mantoida*, Chaeteessa, Metallyticus, Amorphoscelinae, and Perlantinae, in which the pronotum is almost square”, suggesting that this last character is homoplastic (Wieland 2013: 50). All these characters are currently considered as synapomorphies of the mantodean crown-group.

The pronotum of *Pseudomantoida* is saddle-shaped, as in *Mantoida* and *Paramantoida*, even its lateral parts are more bent ventrad than in these extant taxa (Fig. 5; Agudelo 2014). The fore tibia has only one posteroventral spine, which is a character only present in the Mantoididae, Amorphoscelidae, and few other taxa. Generally the mantises have such spines more numerous. These characters could be putative synapomorphies of the Mantoididae. This fossil has also an apical claw on fore tibiae, and prominent ocelli, characters shared by the Mantoididae but not the Chaeteessidae (Agudelo 2014). Lastly, *Pseudomantoida* differs from *Ambermantis* Grimaldi, 2003 (type genus of the Cretaceous family Ambermantidae Grimaldi, 2003, a taxon sister group of the “Eumantodea” sensu Grimaldi [2003], crown-group of the Mantodea) in the shorter legs, the S-shaped posterior margin metazona, and different number of spines on fore femora. This last genus shares with *Pseudomantoida* a metazona nearly two times as long as prozona, with a long extension above wings, and outer margin of pronotum forming a lateral carina without spines. Notice that the forewing venation of *Ambermantis* is very similar to those of the extant Mantoididae, especially the anterior branch of the vein AA2 posteriorly pectinate with many branches (see Agudelo 2014). This character could represent a potential synapomorphy of the Ambermantidae with the Mantoididae.

*Pseudomantoida extendidera* appears to be very similar in size to the extant *Paramantoida amazonica* Agudelo, 2014 (head width 3.8–3.9 mm, prozona length 0.8–1.0 mm) (Agudelo 2014). Therefore, the total length of *Pseudomantoida extendidera* was probably also similar, i.e., 18–20 mm long.

*Mantoida matthiasglinki* Zompro, 2005 is currently considered to belong to the genus *Mantoida* and therefore to the Mantoididae. However, it has many characters strongly different from those of other Mantoididae (extant *Mantoida* and *Paramantoida*, and *Pseudomantoida*): cursorials legs are longer than those of Mantoididae (e.g., hind femora as long as abdomen while they are clearly shorter in *Mantoida* and *Paramantoida*); fore legs also longer than those of Mantoididae, and with large spines on femoral and tibial posteroventral spines (small or reduced in Mantoididae). These characters would justify the attribution of this species to another family. The long legs and spines, associated with the very short pronotum, are also known in the Cretaceous *Ambermantidae* (*Ambermantis*) (Grimaldi 2003), which are very similar to *M. matthiasglinki* in their general habitus. However, the anal veins seem to be different; *M. matthiasglinki* has discoidal spines while these seem to be absent in *Ambermantis*. The long legs and spines present in both taxa can result from an adaptation to a particular life style, viz. fast-running predation on small preys. Notwithstanding, *M. matthiasglinki* needs a revision. Currently it cannot be accurately assigned to a precise family of mantises. Therefore, *Pseudomantoida extendidera* is the only one known fossil Mantoididae, thus of great interest for date calibration.

Stratigraphic and geographic range.—Only type locality.

Concluding remarks

The new specimen of *Arverninea insignis* confirms the placement of this species in the Chaeteessidae. Therefore, *Arverninea insignis* is the oldest known reliable representative of this family. *Pseudomantoida extendidera* is the only reliable known fossil of the Mantoididae while *Protohierodula crabbii* is the oldest Artimantodea sensu Svenson and Whiting (2009). These three fossils are suitable for date calibration. Chaeteessidae are currently considered as sister group of all other crown Mantodea while Mantoididae are the sister group of the rest (Artimantodea) (Svenson and Whiting 2009). The three fossils allow dating three successive profound nodes in the phylogeny of Mantodea: with *Arverninea insignis* (60 Myr) for the crown group of Mantodea, *Pseudomantoida* (53 Myr) for the crown group of (Mantodea minus Chaeteessidae), and *Protohierodula* (late Eocene) for (Artimantodea + Mantoididae). In our knowledge, these calibration points are the
only accurate known points and are crucial to understand the evolution of the order.

The most diverse extant group among Mantodea is the Cernomantodea that have cyclopean ear detecting the ultrasonic sounds of bats (Yager and Svenson 2008). This group could have "replace" the Mantodea without earing system during the Paleogene. This hypothesis is congruent with the lack of Cernomantodea and the high diversity of Chaeteessidae in the Paleocene and Eocene. A similar phenomenon occurs with the Neuroptera: Chrysopidae, viz. the Chrysopinae have developed tympanic organ allowing them to detect the ultra-sounds of the bats. Their diversity greatly increased during the Neogene while the Nothochrysinae, without tympanic organ, were much more diverse during the Paleogene (Archibald et al. 2014). Insectivorous bats are recorded in the middle Eocene (those from of Messel having insects in their guts) (Habersetzer et al. 1994; Smith et al. 2012). It seems that the bats’ diversification had a significant impact on many insects that are relatively poor flyers. It also could indicate that Mantodea experienced a relatively recent diversification since the last 40 million years. This is congruent with the lack of morphological synapomorphies, typical of evolutionary radiation.

The two extant families Chaeteessidae and Mantoididae are strictly Neotropical; the presence of the former in Europe, North America, and Asia during the Paleocene and Eocene, and of the later in the Eocene of Europe could appear surprising, but several other similar cases of Paleogene faunal exchanges between the Western and Eastern Hemispheres and/or Paleogene widespread (even worldwide) distributions of taxa are known. For instance representatives of the extant African dragonfly subfamily Neophyinae Tillyard and Fraser, 1940 are recorded in the South American Paleogene but also in the latest Eocene of England (Nel and Fleck 2014). The extant Australian termite family Mastotermitidae has a Paleogene distribution in South and North America, Europe, Asia, Africa, and Australia. The Paleocene–Eocene was an exceptional period for Eurasian-North American interchanges (Brikiatis 2014). Nevertheless South America was well-separated from North America during the Paleocene–Eocene. It is possible that the Chaeteessidae and Mantoididae were present in North America and colonized the South America later. The other hypothesis is that both families were much older (Late Jurassic–Early Cretaceous) and colonized the continents before the opening of the Northern part of the Atlantic Ocean. Future molecular dating and new discoveries of fossil Mantodea in North America and Africa could help to solve this question.

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References

