The long-awaited first instar larva of *Paussus favieri* (Coleoptera: Carabidae: Paussini)

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**Abstract.** *Paussus favieri* Fairmaire is one of only two species of the myrmecophilous carabid tribe Paussini known from Europe. Larvae are known from only 10 of the 580 paussine species. As in many beetles with considerably modified later instar larvae, the first instars represent a valuable source of informative characters for taxonomy and phylogenetic analyses (primary chaetotaxy, egg-bursters, etc.). Therefore, the discovery of the first instar larva of *P. favieri* is particularly important, as it represents only the second species for which this larval stage is known. In this paper we describe the behavior and morphology of the larval first instar of *P. favieri* (subtribe Paussina of Paussini) and compare it with that of *Arthropterus* sp. (subtribe Cerapterina), which is the only other 1st instar described in the Paussini. Most surprisingly, we found that the 1st instar of *P. favieri* lacks a prostheca, which was previously thought to be a synapomorphy of Paussina + Platyrhopalina. Rather, *P. favieri* has a unique mandibular structure that seems to be functionally analogous to the prostheca. It is a long, broadly lanceolate, distinctly flattened structure apparently homologous to the medial mandibular seta (MN2*), which arises from an area behind the cutting edge of mandible. We predict that the function of the prostheca and this similar structure in *P. favieri* are involved in a specialized feeding strategy that may include soliciting trophallaxis from their host ants. We also report some observations of the first instar hatching from the egg, feeding on liquid and a behaviour we interpret as a “calling behavior,” all of which were videotaped and posted on the Tree of Life Web Project.

**INTRODUCTION**

Myrmecophiles (obligate symbionts of ants) are challenging to study because they are rare, they live in concealed environments (ant nests), and they have complex but little understood interactions with their hosts. While it is difficult to study their behavior and life cycle in nature, there are also challenges rearing them in laboratory conditions. Because these difficulties hamper direct observations, the nature of most interactions between myrmecophiles and ants, and the function of many structural adaptations, remain a mystery or a matter of speculation (e.g., Di Giulio & Moore, 2004).

Myrmecophiles possess specialized adaptations in order to be accepted by ants and to survive and develop within their nests. Such adaptations include: chemical and morphological mimicry; specialized behaviors to feed on and/or be fed by ants; and structural and chemical modifications to avoid ant attacks. Holometabolous myrmecophiles are unique in that they have different suites of adaptations for very different life history stages (larva, pupa, adult). For many holometabolous myrmecophiles, adults can freely enter and leave the nests, but the larvae (and pupae) must remain in the nest full-time and they are completely dependent upon the ants for survival.

Most members of the carabid beetle subfamily Paussinae are myrmecophiles and all members of this subfamily have a unique type of larva referred to as “dis-cotelic” (Di Giulio, 1999, 2008) for the presence of a wide, transverse, plate-like structure, called the terminal disk at the end of an up-curved abdomen (Bousquet, 1986; Di Giulio, 1999; Di Giulio et al., 2000). The terminal disk is composed of modified epipleurites VIII and IX, tergite VIII, and the urogomphi. In free-living, non-myrmecophilous larvae the terminal disk is used to trap prey through an ambush feeding strategy, after the prey are attracted to it by chemical attractants (Costa et al., 1988; Di Giulio, 1999; Di Giulio & Vigna Taglianti, 2001; Moore & Di Giulio, 2006). All described species of the monophyletic tribe Paussini are thought to be parasitic myrmecophiles as both larvae and adults (Darlington, 1950; Nagel, 1979, 1997; Di Giulio & Moore, 2004; Geiselhardt et al., 2007). Components of the terminal disk are fused in myrmecophilous Paussini larvae such that they cannot trap prey like their free-living relatives. Substances on the fused terminal disk most likely appease ants and play important roles in establishing and maintaining of their symbiosis with ants (Boving, 1907; Oberprieler, 1985; Bousquet, 1986; Luna de Carvalho, 1989; Di Giulio & Moore, 2004; Di Giulio, 2008).

The first Paussini larva was discovered more than hundred years ago (Boving, 1907; Di Giulio, 2008). Since then our knowledge of the group has grown very slowly. To date, we know larvae of only 10 of approximately 580 described species (for a list, see Di Giulio & Moore, 2004; Moore & Di Giulio, 2006).
and 4 of the 22 genera (Arthropterus W.S. MacLeay, 1838, Platyrhopalopsis Desneux, 1905, Paussus Linnaeus, 1775, Granulopaussus Kolbe, 1938). Two other larval descriptions (Wasmann, 1918; Brauns, 1914; van Emden, 1922) are too vague and superficial to be useful, the identifications having been based solely upon speculations, and the deposition of material is unknown (Di Giulio et al., 2003; Di Giulio & Moore, 2004; Di Giulio, 2008). While first instar larvae are widely acknowledged to be an important source of diagnostic characters (i.e., primary chaetotaxy and egg burster shape) (Bousquet & Goulet, 1984), most of the described Paussini larvae are second or third instars. To date the only first instar larva described in this tribe is that of Arthropterus sp., classified in the subtribe Cerapterina (Di Giulio & Moore, 2004).

The tribe Paussini is mainly tropical and subtropical with very few species known in the Palearctic Region (see Nagel, 2003 for an updated list). Only two species are distributed in Europe, Paussus turcicus I. Frivaldszky von Frivald, 1835 and P. favieri Fairmaire, 1851. Paussus turcicus occurs in West-Balcan, Turkey, S-Caucasus and Turkmenistan (Nagel, 1987, 2003) and P. favieri is an Atlanto-Mediterranean species (Nagel, 1987, 2003) present in southwestern France, Spain, Portugal, Morocco, Algeria and Tunisia (Casale et al., 1982; Nagel, 1987). There are some museum specimens of P. favieri from Sicily and Sardinia (Casale et al., 1982), and one from Corsica (Zerche, 1990), however, no specimens have been collected on these islands during the past 100 years. The closest relatives of P. favieri, are P. biflagellatus Luna de Carvalho, 1973 and P. krelli Kaupp & Rödel, 1997 (Luna de Carvalho, 1980; Kaupp & Rödel, 1997) known only from the sub-Saharan countries of Ghana and the Ivory Coast. For over 150 years P. favieri has attracted the intense interest of researchers and collectors, due to its rarity, its bizarre structural adaptations to a myrmecophilous lifestyle, and the fact that it is one of the few paussine species known from the Mediterranean Region. Through the years, many researchers have tried to learn about the life cycle and preimaginal stages of this species but to no avail (an apodous, physogastic larva was erroneously attributed to P. favieri by Xambeu, 1892).

Recently, on a collecting expedition to the High Atlas Mountains of Morocco we collected a gravid female of P. favieri and in our laboratory in Rome we were able to rear the first instar larva of this remarkable species. These specimens represent the first record ever of first instar specimens of the subtribe Paussina. In this paper we describe P. favieri first instar larval morphology, provide notes on collecting adults, rearing conditions and behavior of the larva in captivity. We also compare this larva to the first instar of Arthropterus, the only other first instar described for the tribe Paussini.

MATERIAL AND METHODS

Several adult specimens of Paussus favieri, collected during an expedition to High Atlas Mountains in Morocco (2009) from nests of Pheidole pallidula (Nylander, 1849), were kept in captivity for about one month under controlled conditions. During this time, their behavior and interactions with their host ants were studied (Di Giulio et al., unpubl.). One of the females laid two eggs on the moist filer paper. Eggs were held at room temperature (about 25°C), inside a vial closed by a moist cotton plug. Two larvae hatched eleven days after isolation. Larvae were observed under Olympus SZX16 stereomicroscope, and interactions among the larvae, adults and host ant brood were recorded by a camera (ColorView II, AnalySIS software Olympus SIS, Cell *D, 2006). Two days after hatching, the small larvae were submerged in boiling distilled water for a few seconds and then transferred to 70% EtOH. Larvae were drawn using Olympus SZX16 equipped with drawing tube (Figs 1–5). Then, one specimen was rehydrated, cleared in 10% KOH, transferred to hot lactic acid, dehydrated through a series of EtOH baths of increasing concentration (10, 20, 50, 70, 95 and 100%), left overnight in a clove oil bath, and mounted on a slide with Canada balsam. This specimen was illustrated by using a light microscope Olympus BX51 equipped with drawing tube. The second specimen was dehydrated through a series of EtOH baths of increasing concentration (70, 80, 90, 95 and 100%), critical point dried (Bal-Tec CPD 030), mounted on a stub (by using self adhesive carbon disks), sputtered with gold (Emitech k550 sputter coater), and observed with Philips XL30 scanning electron microscope and FEI Dualbeam FIB/SEM Helios Nanolab (L.I.M.E. laboratory, University “Roma Tre”, Rome). In this paper, the general terminology of larval structures follows Lawrence (1991). The term “frontoclypeolabrale” is used here instead of “frontale” or “fronal plate” (more often used in larval taxonomy of Carabidae) or “frons” or “frontal plate” (more often used in larval taxonomy of Carabidae) or “frontale” or “frONTAL region” (see Lawrence, 1991), following the recent taxonomic descriptions of Paussinae (Di Giulio & Moore, 2004, 2009; Moore & Di Giulio, 2006) and the available larval key (http://tolweb.org/notes/?note_id=3430). Notation of primary setae and pores follows the system of Bousquet & Goulet (1984), modified for Metrius contractus Eschscholtz, 1829 (Bousquet, 1986). As some of the sensilla of P. favieri larva present on the abdomen and terminal disk are homologous to those recognized by Bousquet (1986) in Metrius contractus (sensilla S-I to S-V), by Di Giulio (1999) and Di Giulio et al. (2000) in several species of Pachyteles Perty, 1830 (sensilla S-I to S-VII), and by Di Giulio & Moore (2004) in Arthropterus sp. (sensilla S-I to S-VIII), we adopted the same nomenclature used by these authors. Notation of microsculpture follows Harris (1979). An asterisk (*) following a coded seta indicates that the homology...
between the structure in the *P. favieri* larva and the corresponding code is questionable.

**RESULTS**

**Behavioral observations on Paussus favieri first instar larva**

Immediately after hatching the terminal disk is closed. That is, the dorsal and lateral plates are stuck to the ventral plates (http://tolweb.org/media/44075). The larva actively moves, twisting its head and thorax up, down and side to side for about one hour. After this period of hyperactivity, it remains still, stretching and inflating its body until the terminal disk is completely opened and sclerotized. Once the body becomes white-yellow and head and claws more sclerotized, the larva becomes active again, opening and closing its mandibles (which it can move independently of one another) with abdomen and terminal disk bent dorsally. It is unable to use its short legs for walking, but rather for support. On several occasions we observed the larva grasping the filter paper that lined the observation chamber with its mandibles, thereby creating an anchoring point. The larva would then flex its body vertically, bringing the terminal disk up and over its head (http://tolweb.org/media/44071). We also observed what we interpret to be a “calling behavior” in which the larva balances on its hind legs and urogomphi as it lifts its head and thorax into the air, moving from side to side while

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**Fig. 2. Paussus favieri** first instar larva: a – head outline, dorsal view; b – head outline, ventral view; c – left side of cephalic capsule with setation, dorsal view; d – right side of cephalic capsule with setation, ventral view.
opening and closing its mandibles, and waving its mid and fore legs (http://tolweb.org/media/44074).

Most often the larva did not show interest toward the host ant brood, but on one occasion we observed an unsuccessful attempt to pierce the brood with its mandibles (http://tolweb.org/media/44076). However, when offered damaged brood the larva began to actively suck the host’s hemolymph, demonstrating an aptitude for liquid feeding (http://tolweb.org/media/44077).

Description of Paussus favieri egg and first instar larva

Egg

Ovoid, whitish, surface reticulation with cell length of 3.5–4.8 µm. Egg-shell assignable to type B of Kaupp et al. (2000). Egg length 0.7 mm; width 0.4 mm.

First instar larva

Diagnosis. The presence of a long, broadly lanceolate, distinctly flattened medial mandibular seta (Figs 3c, 7e–f).
easily distinguishes this larva from all known larvae of Paussini.

Measurements. Body length 2.3 mm (from tip of mandibles to the terminal disk); cephalic capsule maximum width (at base of antennae) 0.45 mm, medial length (from occipital foramen to anterior emargination of fronto-clypeolabrum) 0.18 mm, occipital foramen width 0.3 mm; antennal length 0.19 mm; mandible length (measured along outer margin) 0.22 mm, length of modified seta 0.1 mm; pronotum maximum width 0.52 mm, maximum length (along ecdysial suture) 0.2 mm; leg length 0.35 mm; diameter of terminal disk 0.68 mm, dorsal plates length (measured along medial suture) 0.35 mm, urogomphi (ventral plates) length (measured along medial suture) 0.28 mm.

Habitus and coloration. When newly hatched, the larva appears transparent and very soft, except for apex of mandibles and egg-bursters; after sclerotisation, body soft, weakly sclerotised, not physogastric, up-curved (Figs 1, 6a), with cup-shaped, sclerotised, terminal disk (Figs 1, 5, 8a) held in an elevated position by a flattened, sac-like abdomen (Figs 1, 6a,c). Body white-yellow, semi-transparent; egg-bursters, claws and terminal disk heavily sclerotised and light-brown; mandibles reddish-brown (medially) to dark-brown (at apex).

Microsculpture. Cephalic capsule, mouthparts, thoracic and abdominal tergites, legs and pygidium smooth or irregularly wrinkled to carinate, probably due to incomplete sclerotisation. Basal half of mandibles, dorsally, with irregularly reticulate to scale-like, rugulose microsculpture. Medial region of epipharynx and hypopharynx with multipointed microsculpture, arranged in parallel, transverse rows; lateral margin of frontal emargination and sides of hypopharynx with thin elongate protuberances (Fig. 7f); membranous body areas microgranulate (Fig. 6f); surface of terminal disk densely covered by finely granulate microsculpture, mixed with sparse, conical spines (Figs 8c–f).

Chaetotaxy. Head. Frontoclypeolabrale with many (about 70) setae of different sizes (identification not possible) and not strictly paired (Figs 2c, 7a); primary pores apparently absent. Each parietal plate (Figs 2c–d, 7a–b) with about 45 setae of different sizes, about 20 dorsal and about 25 ventrolateral. Antennae (Figs. 3a–b, 7d): Antennomere I with 8 additional setae, mainly dorsally and laterally, all primary pores present; antennomere II with 3 additional setae; antennomere III with 13 setae and 3 pores, mainly on dorsal side, identification with primary structures not possible. Mandible (Figs 3c, 7e–f) with seta MN1* dorsolateral, elongate, bent inward; seta MNb* modified into a long, broadly lanceolate, distinctly flattened structure, directed mesad; 5 pores present on each mandible: MNa lateral, MNb* and additional pore mesodorsal, MNc* and additional pore more distally, close to base of modified MNb*; Setal group gMX on stipes with 9 setae: 4 longitudinally lined and 5 and on the remnant of lacinia (?); about 9 setae present on lateral and ventral sides of stipes: 5 at the level of the partially fused palpomere I (including MX10) and 4 mesolaterally (2 lateral possibly being MX2 and MX3); apex of galeomere II with MX9* long and small blunt-tipped sensorial papilla; maxillary palpomeres as follows: II with MXe and MXf ventral and 2 additional setae, one lateral and one mesal; III with 4 dorsal setae and 2 pores; IV with 2 additional setae subbasally on ventral side, 3 slender digitiform sensilla subapically and one apical sensorial area with about 9 elongate subequal papillae (sensilla basiconica); cardio with 2 setae (mesal possibly MX1); pores MX3, and MXe not visible. Dorsal and lateral setae on prementum moderately long and hair-like, ventral setae elongate and curved; LAe subapical, much longer than palpomere I; 8
Fig. 6. *Paussus favieri*, SEM images of first instar larva: a – habitus, right dorsolateral view; b – thorax, dorsal view; c – abdomen, dorsal view; d – right legs of pro and mesothorax, anterior view; e – left prothoracic claw; f – right mesothoracic spiracle. Scale bars: a = 500 µm; b = 300 µm; c = 400 µm; d = 100 µm; e = 20 µm; f = 5 µm.
Fig. 7. *Paussus favieri* SEM images of first instar larva: a – head, dorsal view; b – head, right dorsolateral view; c – right egg-burster, right lateral view; d – apex of right antenna, dorsal view; e – left mandible, dorsal view; f – mouthparts, anterodorsal view. Scale bars: a–b = 100 μm; c, e, f = 50 μm; d = 10 μm.
pairs of setae dorsally on prementum, LA3–5 not possible to identify; ventrally LA; basally close to midline, 4 pairs of setae (LA; and 3 additional setae) apically; labial palpomeres as follows: I with 4 pores ventrally (including of setae (LA 2 and 3 additional setae) apically; labial palpomeres as follows: I with 4 pores ventrally (including LAa); II with 3 additional setae: I short, dorsal, and 2 lateral (1 basal spiniform and 1 more distal very short); 2 slender digitiform sensilla subapically and one apical sen- sorial area with about 9 elongate subequal papillae (sen- silla basiconica), similar to those of maxillary palpomere IV. Thorax. Pronotum (Figs 4b, 6b) with about 55 setae on each side of ec dysial suture (primary setae impos- sible to identify); notopleural setae very long and thin; mesonotum and metanotum (Fig. 6b) with about 15 setae on each side of ec dysial line. Distal leg segment (corre- sponding to fused trochanter, femur, tibia and tarsus) with a tuft of about 70–80 long and thin setae, surrounding also the apical claw (Figs 4a, 6d–e). Sternal areas of pro-, meso- and metathorax with about 10 pairs of setae, mesal ones longest. Abdomen. Terga of abdominal segments I–VII (Fig. 4c) with about 20 pairs of setae each (identification not possible) and no pores. Dorsal and lateral plates of terminal disk dorsally (tergal side, Fig. 8b) with many short, spiniform, regularly spaced (every 20–40 µm) sensilla S-VII: about 150 (mostly on subapical area) on each dorsal, about 15 on each lateral plate; each ven- tral plate (urogomphus) with about 50 long setae on ven- tral side; perimeter of disk (margin of dorsal, lateral and ventral plates) with about 55 elongate (120–130 µm) sen- silla S-II (Figs 5, 8c), regularly spaced at margin of plates and urogomphi, radially oriented to the surface of ter- minal disk: 10–13 on each dorsal plate, 5 on each lateral plate and about 11–12 on each urogomphus; each sen- sillum S-II lanceolate, straight or slightly curved and pointed apically, basally tapered and inserted in a sunken socket; dorsal surface longitudinally keeled (9 slightly protruding carinae) and bearing irregular pits (about 2 µm), giving it a spongy appearance (Fig. 8c). Dorsal sur- face of terminal disk with many sensilla S-I (Figs 5, 8d–e): about 55 on each dorsal plate, 5–6 on each lateral plate and about 40 on each ventral plate (urogomphi); each sensillum S-I composed of a dome-like protruding base (diameter 15–18 µm, height 8–10 µm), subapically with a crown of 4–6 multisupplanted lobes and apically with a medial short seta (3–4 µm), expanded and multisupplanted at apex (Fig. 8e); S-I with longer frayed seta and smooth conical base regularly alternating with S-II along entire margin of plates; 4 spiniform sensilla emerging from medial apex of dorsal plates. Dorsal plates and urogomphi with many scattered filiform microstructures 0.4 µm thick (Fig. 8f) emerging from cuticular pores (diameter 2.5 µm). Epipleurites of abdominal segment I with 1 or 2 setae each; epipleurites of abdominal seg- ments II–VII with 4–6 setae each. Hypopleurites of abdominal segments I–VII with 3–5 additional setae each. Sternal areas (Fig. 4c) with 20–30 setae on each abdominal segment (homologisation not possible). Pygidium without setae.

Head. Strongly transverse (Figs 2a–b, 7a), two times as wide as long, prognathous, subparallel-sided, basally nar- rowed, with maximum width at base of antennae; cephalic capsule steeply sloped dorsally from base to apex, not retracted into prothorax (Fig. 7a); base of head capsule and occipital foramen lined with a sclerotized band. Frontoclypeolabral sub-hexagonal (Fig. 2a); ante- rior margin not sclerotized, distinctly conceave, medially very thin and slightly emarginated; surface distinctly convex posteriorly and anterolaterally, deeply conceave anteromedially; transverse frontal keel absent; egg- bursters well developed, consisting of two strongly sclerotized longitudinal keels, posteriorly slightly conver- gent, ending anteriorly in a sharply pointed spine directed anteriorly (Figs 2a, 7a–c); coronal suture absent; fronto- clypeolabral almost fused with parietalia, frontal sutures very fine but still visible in light microscope. Parietalia (Figs 2a–c, 7a–b) subparallel-sided, without stemmata, protruding at base of antennae; ventral walls of parietalia medially fused into a short gular suture. Antennae (Figs 2a, 3a–b, 7a, d) 4-jointed, directed forward and slightly convergent anteriorly, not reaching apices of the mandi- bles, inserted in extensive membranous, soft elevations; antennomeres I–III wide; I asymmetrical, shorter on ental side, slightly longer than broad, more than three times as long as II; II short, three times wider than long; III about six times as long as III; II slightly shorter than IV; IV very small compared to others; sensorial appendage ovoid, positioned ventrolaterally on antennomere III (Fig. 7d), about as long as IV. Mandibles (Figs 3c, 7e) subtriangular, slightly falcate apically, 1.5 times as long as wide at base, with single, ventral cutting edge; dorsal surface deeply excavate along occlusal margin, particularly at base of modified mandibular seta; retinaculum slender, triangular, sharp and pointed, displaced and directed apically, forming, together with the pointed apex, a bidentate mandible; prostheca and plicellus absent. Maxilla (Figs 3f–g) with small, subtriangular cardo and subquadrate stipes; maxillary palpus 4-jointed: I partially fused with stipes, about twice wider than long; II slightly longer than I; III subquadrate, twice as long as II; IV conical and subulate; galea 1-jointed (corresponding to galeomere II of Ozaenini, galeomere I being fused with stipes), digitiform, almost straight, distinctly tapered from base to apex; lacinia small, vestigial; lateral margin of stipes distinctly curved, occlusal margin straight without basal tooth. Labium composed of a membranous mentum, ventrally sclerotised prementum and 2-jointed palps. Prementum (Figs 3d–e) slightly enlarged from base to apex, particularly in ventral view; basal half of prementum dor- sally bulging, distal half with vestigial subapical ligula, represented by prominent bases of strong setae LAc. Labial palpomeres subequal in length: 1 subquadrate, II conical, subulate (Figs 3d–e). Hypopharynx densely cov- ered by transverse parallel rows of pointed papillae (Fig. 7f), closely fitting in the vault of the oral cavity.

Thorax. Thoracic segments (Figs 1, 6b) slightly sclero- tised; tergites, pleurites and sternites poorly delimited, mostly recognizable by their setae and a smooth surface; tergites similar in length, widely transverse with slightly rounded (pronotum) or straight (meso- and metanotum)
Fig. 8. *Paussus favieri* SEM images of first instar larva: a – terminal disk, apical view; b – terminal disk, dorsal view; c – dorsal plates, marginal sensilla S-II; d – dorsal plates, sensilla S-I; e – dorsal plates, sensillum S-I; f – dorsal plates, filiform structure (glandular pore with substance? sensillum?). Scale bars: a = 300 µm; b = 200 µm; c = 30 µm; d = 40 µm; e = 5 µm.
sides; surface of tergites irregularly convex (Fig. 6b); longitudinal ecysial line restricted to pronotum. Pronotum composed of single distinct sclerite, slightly concave anteriorly and convex posteriorly, about three times as wide as long. Mesonotum and metanotum, with tergites longitundinally subdivided into two subtriangular parts, widely separated by membranous integument. Sternal of thoracic segments soft, un sclerotised.

Spiracles. All spiracles annular-uniformous (Fig. 6f); peritreme dome-shaped, convex and with a round hole at top; atrium poorly developed. Mesothoracic spiracles between pro- and mesonotum similar to but slightly larger than abdominal spiracles, with wider atrium; small rudimentary spiracles present in metapleura. Abdominal spiracles I–VIII dorsolateral, partially sunken in bulged membranous areas above epipleura (Fig. 1).

Legs. Short and highly modified, all similar in type and length (Figs 4a, 6d), composed of 2 joints subequal in length: basal joint representing coxa, basally embedded into soft membranous areas, dorsally articulated with medially subdivided pleural sclerite; distal part of coxa obliquely truncate to allow folding of second joint; latter cylindrical, digitiform, slightly tapered to apex, composed of completely fused trochanter, femur, tibia and tarsus; sutures between segments hardly visible. Procoxae more slender and shorter than meso- and metacoxae. Second joint bearing single conspicuous claw (Figs 4a, 6e), apically curved and pointed.

Abdomen. Subparallel-sided, gradually enlarged toward terminal disk, scarcely or not sclerotised, distinctly curved in an upturned position (Figs 1, 6a, c); narrowest at segments III–V; terga flattened or slightly convex, pleura and sterna swelling. Terminal disk (Figs 5, 8a–b) wider than segment VII, regularly round, with perimeter strongly raised, corrugated and markedly sclerotized; disk composed of 6 symmetrical subtriangular plates joined by thin membranous lines; dorsal plates wider than others, medi ally convex and bearing abundant sensilla S-I; lateral plates small; ventral plates corresponding to modified urogomphi, similar in shape to dorsal plates, with surface flattened or slightly concave. Pygidium cy lindrical, dorso-ventrally flattened, ventral to urogomphi.


DISCUSSION

Within the subfamily Paussinae, larvae are known from 4 of the 5 tribes (Metriini, Mystropomini, Ozaenini, Paussini), the larvae of Protopaussini being still unknown. Within these tribes, only one or few species of 11 genera (out of 46; Lorenz, 2005) are known as larvae, often described based on a single specimen of second and/or third instar. Though our knowledge of Paussinae larvae is very limited, the taxonomic sampling is broad enough to provide information about relationships among major clades. However since larvae are rarely collected in the field and difficult to rear in the laboratory, phylogenetic analyses based on larval morphology compared different instars of the taxa included in the analyses (Bousquet, 1986; Beutel, 1992; Vigna Taglianti et al., 1998; Di Giulio et al., 2003; Di Giulio & Moore, 2004). For this reason, only morphological characters related to shape of the head capsule, head appendages, legs and terminal disk have been used. It has not been possible to include first instar larval characters, such as those related to egg bursters or primary chaetotaxy (Bousquet & Goulet, 1984), which are widely acknowledged to be phylogeneti cally informative (Bousquet & Goulet, 1984; Bousquet, 1986; Arndt, 1998; Meier & Lim, 2009). In fact, until now first instar larvae in Paussinae were known for only four species: *Metrius contractus* (Metrini), *Pachyteleae vignai Deuve, 2000* (Ozaenini), *Goniotropis kuntzeni* (Bänninger, 1927) (Ozaenini), and *Arthropterus sp.* (Paussini).

The discovery, description and observations of the 1st instar larva of *Paussus favieri* presented herein are signi ficant in many respects. First the paper provides information about the larval stage, behavior, and life cycle of a rare species that is increasingly endangered due to the destruction of suitable habitats (*P. favieri* is endemic to the Mediterranean Region, and is one of only two species of Paussini occurring in Europe). It also increases the number of described myrmecophilous larvae, thereby broadening our understanding of evolutionary adaptations to a myrmecophilous life in Paussini. In the following discussion, the morphological characters observed in *P. favieri* 1st instar larva are analyzed and discussed in comparison with those of *Arthropterus sp.*, the only other known species of the Paussini.

All the synapomorphic characters of Paussini (Di Giulio & Moore, 2004) are found in the larva of *Paussus favieri*, including: Neck not constricted, head prognathous, coronal suture absent, antenna short and broadly inserted in bulging membranous base, large antennal sensorial appendage, mandibles short and pointed with some long setae, stipes without basal tooth, setal group gMX extremely reduced, prementum bulging, one tarsal claw, urogomphi plate-like, terminal disk perfectly round, pygidium ventral to urogomphi.

Most Paussini larvae have an elongate digitiform prost heca, a soft structure with an anterior ridge that almost reaches the apex and arises near the base of the ental surface of the mandible. Surprisingly, the prostheca is not present in the 1st instar of *P. favieri*, although it has been considered a synapomorphy of the Paussina + Platyrhop alina. While *P. favieri* does not have a prostheca, it does have a similar structure that is apparently homologous to the median mandibular seta (MN2*). It is a long, broadly lanceolate, distinctly flattened structure (Figs 2a, 3e, 7a,e–f), arising from an area behind the cutting edge of the mandible. The functions of the prostheca and of this similar structure in *P. favieri* are unknown, but possibly these larvae have a specialized feeding strategy that may include soliciting trophalaxsis from the host ants.
The reduction of the base of the cephalic capsule (frontal + parietalia), with loss of coronal suture and basal part of frontal sutures up to the base of the egg bursters, is shared by the larvae of *P. favieri* (Figs 2a, 7a) and other known larvae of the Paussini subtribes Paussina and Platyrhopalina, being possibly their synapomorphy (Di Giulio & Moore, 2004). This reduction from the ancestral state found in other Paussinae tribes (e.g., Metriini, Ozaenini) may be related to the secondary acquisition of a prognathous position of mouthparts and head derived from a hyperprognathous ancestral state (Di Giulio et al., 2003). Such a basal degeneration of the head (including the widening of the occipital foramen and a disappearance of the neck) could be an adaptation for living with ants.

As in all known larvae of Paussina, the basal maxillary palpomere and the stipes of *P. favieri* are partially fused (Figs 3f–g); in some other *Paussus* species this fusion is complete. A small remnant lacinia is present in *P. favieri* (Fig. 3f), whereas it is completely absent in other Paussina and Platyrhopalina. *Arthropterus* has a highly derived lacinia with a row of strong hook-like setae.

In contrast to the larva of *Arthropterus*, which has distinct and functional leg articles, the trochanter, femur, tibia, and tarsus of *P. favieri* are fused (Fig. 4a) and remnant sutures are only visible with SEM (Fig. 6d). Such fusion is a synapomorphy of Paussina and Platyrhopalina, though in *Platyrhopalopsis* traces of oblique sutures between the fused articles are clearly visible. The reduction from two (Metriini + Mystropomini + Ozaenini) to one claw seems to be a defining character of Paussini. Most Paussina and Platyrhopalina larvae have a very small and thin tarsal claws, whereas *Arthropterus* and *P. favieri* larvae have conspicuous, robust tarsal claws. It is possible that this is a characteristic of all first instar larvae as compared with latter instars.

As in the other *Paussus* larvae (Paussina) and in *Platyrhopalopsis* (Platyrhopalina), an anterior medial emargination of frontoclypeolabral is present in *P. favieri* (Figs 2a, c, 7a), while this modification is absent in the *Arthropterus* larva. For this reason we consider this character to be synapomorphic of the subtribes Paussina and Platyrhopalina. In *P. favieri*, we observed that the medial anterior margin moves during liquid feeding and we predict that this movement may increase the efficiency of suction.

There are several differences between the terminal disk of the first instar *Arthropterus* and *P. favieri* larvae (Figs 5, 8c, e):

1. The marginal setae of *P. favieri* are lanceolate, thin, with a pointed tip, while in *Arthropterus* they are clavate (sensilla S-VIII of Di Giulio & Moore, 2004, possibly homologous to sensilla S-II). Such a feature has not been observed in any other larva of Paussini and could be a feature only present in the first instars. In *P. favieri* a dense substance emerges from the bases of these setae and runs onto the surface of the disk. A possible glandular function was also reported for the homologous setae of *Ar thropterus* sp. (Di Giulio & Moore, 2004). The differences of these setae could be functionally related to different ways of supplying the substance to the ants, from the apex of the club-shaped sensilla in *Arthropterus*, or from the surface of the cup-shaped terminal disk in *P. favieri*.

2. The disk of *P. favieri* is cup-shaped, as in all species of the subtribes Paussina and Platyrhopalina, with upcurved margin, while in *Arthropterus* the disk is biconvex with a simple margin.

3. The dorsal plates of *P. favieri* are slightly wider than ventral plates, and the lateral plates are small, while in *Arthropterus* the dorsal plates are almost twice as large as the ventral, and the lateral plates are wide.

4. The sensilla S-I located on the terminal disk are complex with multisupinate, dome-like bases and short fringed setae in *P. favieri*, whereas the homologous structures in *Arthropterus* are short, simple and coniform, bearing an elongate, simple, blunt-tipped seta.

Nothing is known about the way of life of Paussini larvae except for several field observations which confirm that they are myrmecophilous. In most cases larvae have been found inside ant nests or carried about by the host ants (Luna de Carvalho, 1959, 1992; Geiselhardt et al., 2007). No data on oviposition, feeding, development or behavior are available, not even for the first instar larvae of *Arthropterus*, hatched in captivity from eggs laid by an unidentified female (Di Giulio & Moore, 2004). Various authors have speculated that Paussini larvae feed on host brood and that the round cup-shaped terminal disk is probably adapted for supplying attractive substance to worker ants (Oberprieler, 1985; Luna de Carvalho, 1992). Based on the observation of several characters, unusual for a predaceous carabid larva (i.e., shortened and somewhat degenerated head capsule, reduced mouthparts, unique presence of a prostheca, partial atrophy of legs), Di Giulio (2008) advanced the possibility that they could at least partially be fed by the ants through trophallaxis. The behavioral observations reported here do not exclude this hypothesis. In fact, the “calling behavior” described above (first paragraph of Results section) is similar to that reported for ant larvae soliciting trophallaxis (Hölldobler & Wilson, 1990).

REFERENCES


