Natural history and morphology of the hoverfly *Pseudomicrodon biluminiferus* and its parasitic relationship with ants nesting in bromeliads

Volker S. Schmid¹,²*, Mirian N. Morales³b, Luciane Marinoni³c, Rafael Kamke⁴d, Josefina Steiner⁴, Anne Zillikens²,⁴e

¹Biologie I, Universität Regensburg, 93040 Regensburg, Germany
²Med.-Naturwissenschaftliches Forschungszentrum, Ob dem Himmelreich 7, Universität Tübingen, 72074 Tübingen, Germany
³Universidade Federal do Paraná – UFPR, Dept. de Zoologia, Pós-graduação em Entomologia, Cx. Postal 19020, 81531-900 Curitiba, Paraná, Brazil
⁴Universidade Federal de Santa Catarina, Laboratório de Abelhas Nativas (LANUFSC), Centro de Ciências Biológicas, Campus Universitário Trindade, 88040-900 Florianópolis, Santa Catarina, Brazil

Abstract

The syrphid subfamily Microdontinae is characterized by myrmecophily of their immature stages, i.e., they develop in ant nests. Data on natural history of microdontines are scarce, especially in the Neotropics. Based on fieldwork in southern Brazil, this study provided new data on development and ecology of the hoverfly *Pseudomicrodon biluminiferus* (Hull) (Diptera: Syrphidae) as well as the first morphological descriptions of male genitalia, larvae, and pupa. Immature specimens were specifically found in colonies of the ant species *Crematogaster limata* Smith (Hymenoptera: Formicidae) found in rosettes of the bromeliad species *Aechmea lindenii* (E. Moren) Baker (Poales: Bromeliaceae) and *A. nudicaulis* (L.) Grisebach. Third instar larvae were observed preying on ant larvae, revealing the parasitic nature of *P. biluminiferus*. In this and several other aspects, the natural history of *P. biluminiferus* is similar to that of Holarctic microdontine species. Exceptions include: (i) indications that adults of *P. biluminiferus* outlast the winter months (in contrast to 3rd instar larvae in Holarctic species) and (ii) *P. biluminiferus’* relationship with bromeliads. The importance of bromeliads for this host-parasite system is evaluated in this paper. The single occurrence of another, unidentified microdontine species’ pupae in a nest of the ant species *Camponotus melanoticus* Emery (Hymenoptera: Formicidae) is reported.

Keywords: Brazil, *Camponotus*, *Crematogaster*, Formicidae, host record, Microdontinae, myrmecophily, Neotropics, Syrphidae

Correspondence: volker.schmid@biologie.uni-regensburg.de, mirian_nm@yahoo.com.br, lmarinoni@ufpr.br, rafael_kamke@yahoo.com.br, anne.zillikens@uni-tuebingen.de. *Corresponding author*

Editor: Robert Jeanne was editor of this paper.

Received: 29 May 2012 Accepted: 12 October 2012 Published: 12 March 2014

Copyright: This is an open access paper. We use the Creative Commons Attribution 3.0 license that permits unrestricted use, provided that the paper is properly attributed.

ISSN: 1536-2442

**Introduction**

Colonies of social insects like ants, bees, wasps, and termites provide a very beneficial environment for their brood that includes good nutrition, shelter, favorable climatic conditions, and protection against predators. A highly diverse range of species (called myrmecophiles if associated with ants) has evolved to utilize these benefits for their own development by living inside social insect colonies as either mutualists, commensals, or parasites (Wilson 1971; Kistner 1982; Hölldobler and Wilson 1990; Lachaud et al. 2012). Among myrmecophilous species, those of the syrphid subfamily Microdontinae stand out due to the strange slug- or coccid-like shape and movements of their larvae that caused taxonomic confusion until the early 20th century (Wheeler 1908). Moreover, 454 species of Microdontinae have been described worldwide (Reemer 2013a), but biology and host relationships have been studied for only a few, which are mostly Holarctic species.

All three larval stages of most microdontine species occur in nests of ants (Wheeler 1908; Andries 1912; Akre et al. 1973; Duffield 1981). Field populations of host ants have been shown to be infested by microdontine brood at rates that range from 16% (Akre et al. 1973) to 33–50% of nests (van Pelt and van Pelt 1972). The reported number of microdontine brood items per ant nest varied greatly, from two (van Pelt and van Pelt 1972) to more than 240 (Akre et al. 1973), and reported means ranged from three to six brood items (Duffield 1981; Schönrogge et al. 2002). Most microdontine species developed one brood generation per year (Akre et al. 1988); the species *Microdon fuscipennis* Macquart (Diptera: Syrphidae) develops at least one generation yearly, and some species develop more than one (Duffield 1981). In North American and European species, 3rd instar larvae have been observed to overwinter in their hosts’ nests (Andries 1912; Garnett et al. 1985; Akre et al. 1990).

Larvae of several microdontine species have been reported to feed on their hosts’ brood (Hocking 1970; van Pelt and van Pelt 1972; Duffield 1981; Garnett et al. 1985; Barr 1995). In this context, the larvae can be regarded as either predators of the ants’ brood or parasites of an infested colony as a whole; as a “superorganism” (Hölldobler and Wilson 2009), the colony may be subject to group-level selection. This is the manner in which the terms predation and parasitism will be used throughout this article. A distinct case is primary parasitism of Microdontinae as reported for the species *Hypselosyrphus trigonus*, which infests colonies of *Pachycondyla villosa* (Pérez-Lachaud et al. 2014).

Pupation of microdontine larvae takes place near the surface, where entrances of the nests are found (Wheeler 1908; Duffield 1981; Garnett et al. 1985; Akre et al. 1988; but see Andries 1912). The dorsal surface of the immature stages of development is convexly curved, a characteristic that is most pronounced in 3rd instar larvae and pupae, and covered with a distinct pattern of tubercles, reticulations, or similar structures. The larvae display a posterodorsal stigmatic scar, and pupae are additionally characterized by two small anterodorsal stigmatic horns (Andries 1912; Garnett et al. 1990).

The microdontine pupal stage lasts 11–28 days (Andries 1912; Greene 1923b; Jordan 1968; van Pelt and van Pelt 1972; Akre et al. 1973). Emergence of adults occurs in the early morning or at night. The mere process of emergence requires one to a few minutes, whereas wing expansion takes at least 5–10...
minutes and up to several hours (Wheeler 1908; Andries 1912; van Pelt and van Pelt 1972; Akre et al. 1973, 1988; Duffield 1981; Forti et al. 2007). Sex ratios near 1:1 have been reported (Akre et al. 1973, 1988; Duffield 1981).

Microdontine brood has been reported to be associated with various ant species, predominantly of the genera *Formica* and *Camponotus* (subfamily Formicinae) (Duffield and Thompson 1981; Reemer 2013a). Microdontine larvae and pupae are usually treated indifferently by their hosts (Wheeler 1908, 1910; Jordan 1968; Akre et al. 1973; Garnett et al. 1985, but see van Pelt and van Pelt 1972, who reported that ants investigated and killed some microdontine larvae) or even transported like ant brood by worker ants (Garnett et al. 1985). Similarly indifferent ant behavior toward microdontine imagines immediately after emergence was described by Wheeler (1908). On the other hand, Andries (1912) and Akre et al. (1973) observed that adult microdontines were immediately attacked by ants upon emergence.

At present, few publications about microdontine-ant associations in South America are available, and in most cases the syrphid species was not identified, such as in Paraguay (Sharp 1899), Guyana (Wheeler 1924), and Brazil (Borgmeier 1923; Luederwaldt 1926). Only Forti et al. (2007) reported a fully identified association in Brazil, in which *Microdon tigrinus* Curran (Diptera: Syrphidae) lived in nests of leaf-cutter ants, *Acromyrmex coronatus* (F.) (Hymenoptera: Formicidae). Reports from Central America are also scarce, and include the countries of Panama (Wheeler 1924; Mann 1928) and Costa Rica (Longino 2003a). There are few studies of microdontine-ant myrmecophily in Africa (Wasmann 1894; Speiser 1913; Hocking 1970), Australia (Buschinger 1998), and Asia (Hironaga and Maruyama 2004). A comprehensive overview of the worldwide distribution of reports on microdontine-ant associations was recently provided by Reemer (2013a).

An important condition for the study of microdontine-ant relationships is the ability to identify larvae and pupae, because it is not always feasible to rear adults (Garnett et al. 1990). Detailed knowledge of the morphology of immature developmental stages might also be useful for analyzing the phylogeny of Microdontinae. While descriptions and identification keys to immature stages are available for some Holarctic species (Wheeler 1908, 1910; Greene 1923a, 1923b, 1955; Dixon 1960; Novak et al. 1977; Garnett et al. 1990), no keys and only one general description for a Neotropical species (Wheeler 1924) were available.

During fieldwork in secondary forests of coastal southern Brazil, microdontine pupae were found in bromeliads (Bromeliaceae) inhabited by ants. Bromeliaceae are a monocot plant family mainly distributed in the Neotropics and neighboring subtropical regions (Benzing 2000). Within the microcosms of their leaf rosettes, they harbor a highly diverse range of aquatic and terrestrial animals that are frequently associated with ants (Dejean and Olmsted 1997; Frank and Lounibos 2008; Camargo and Oliveira 2012). In spite of this diversity, no immature specimens of microdontine species had been found or identified in bromeliads prior to the current study.

In order to obtain more data about immature stages of the hoverfly *Pseudomicrodon biluminiferus* (Hull) (Diptera: Syrphidae) and to analyze the animal-bromeliad relationship, there were two aims of the present study. The first was to describe the biology, ecology, and
host-parasite interactions of the Neotropical species *P. biluminiferus* for comparison with species of temperate regions. The second aim was to provide detailed morphological descriptions of immature stages and adults. During the authors’ research on *P. biluminiferus*, immature specimens of another microdontine species were found. Data on this finding are presented and briefly discussed. Aside from morphology, most results about microdontine larvae concern the 3rd instar because 1st and 2nd instar larvae were rarely found.

**Materials and Methods**

**Sample acquisition and studies on developmental biology and ecology**

Fieldwork was carried out between December 2005 and April 2010 on a mountainside covered with secondary forest in Santo Antônio de Lisboa (27° 30’ S, 48° 30’ W) within the district of Florianópolis on Santa Catarina Island in southern Brazil. The first microdontine specimens were found in ant nests within terrestrial bromeliad rosettes (Figure 1A, B, D). To find more specimens, about 300–400 bromeliad rosettes were examined in the field along trails from August to February, and once during April. Bromeliad rosettes were mainly of the species *Aechmea lindenii* (E. Morren) Baker (Poales: Bromeliaceae) (Schmid et al. 2010) and *A. nudicaulis* (L.) Grisebach (Figure 1A, B), and were examined for the presence of ants and immature microdontine specimens (Figure 1D, E). To assess the infestation rate of ant colonies by Microdontinae, 33 ant nests within *Aechmea* spp. were examined for microdontine brood on 5 November (\(N = 20\)), 15 November (\(N = 8\)), and 4 December 2008 (\(N = 5\)), in a different section of the forest each time. To estimate the number of brood items per colony, five ant nests were thoroughly searched for microdontine larvae, pupae, and fresh puparia by breaking the bromeliad rosettes apart leaf by leaf.

To obtain adults, three bromeliads containing ant colonies with *P. biluminiferus* brood were taken into the laboratory, and each was placed into a bucket and enclosed with a fine gauze net. Every day the nets were checked for adult flies. Twenty pupae and one 3rd instar larva collected from bromeliads in the field or in the laboratory were individually placed in transparent vials with a piece of wet paper to provide humidity.

A distinct change of the larval body toward the slightly higher and more slender pupal shape (Figure 1F, I) was interpreted as pupation, marking the beginning of the pupal stage; a similar change has been described in other microdontine species (Wheeler 1908; Andries 1912). This definition might include the prepupal phase (van Pelt and van Pelt 1972), since other authors have defined pupation as beginning “only when the anterior spiracles appeared” (Akre et al. 1973) (“psc” in Figures 1, 5). Because the time of appearance of those spiracles was not noted in our study, only the first pupal stage definition given above could be applied in the current study, which is perhaps an overestimation of the true pupal stage by 1–3 days. Twenty-six adults were obtained altogether, 19 of which could be associated either with an exact emergence time (\(N = 4\)), a reasonable estimate of emergence time (e.g., discovery before wing expansion; \(N = 3\)), or the time of first sighting (\(N = 12\)). Exact emergence time was obtained by serial photography at a rate of one picture every 10 seconds using a Caplio R5 digital camera (Ricoh, [www.ricoh.com](http://www.ricoh.com)), or by video with a Handycam HDR-SR10E camcorder (Sony, [www.sony.com](http://www.sony.com)).
The identity of *P. biluminiferus*, which was initially named *Microdon biluminiferus*, was confirmed by F. Christian Thompson, who compared adult specimens with the holotype deposited in the Naturhistorisches Museum Wien, Vienna, Austria. Ants were identified by comparison with samples in the collection of the Native Bee Laboratory of the Federal
Behavioral studies
To observe the behavior of *P. biluminiferus* larvae and ants toward one another, two artificial laboratory nests consisting of a depression in plaster covered with glass plates were constructed (Figure 1G). Laboratory nests and vials were stored in rooms without climatic and light period control, but were always close to windows so that light conditions were presumably similar to moderately open forest.

First, one colony of the ant host species, *Crematogaster limata* Smith (Hymenoptera: Formicidae), was transferred into an artificial nest (Figure 1G, H, I). The colony contained about 200–300 workers, several winged sexuals, a few dozen ant brood items of different developmental stages, and three 3rd instar *Pseudomicrodon* larvae. To examine whether *Pseudomicrodon* larvae are specifically integrated into their host nests or whether they are generally ignored by their hosts, two 3rd instar *Pseudomicrodon* larvae from another *C. limata colony were placed into the first nest (as described above) after the three initially-present *Pseudomicrodon* larvae had pupated. To obtain an additional (though not completely independent) replicate, the two transferred larvae were moved two weeks later into the second artificial nest, which contained yet another *C. limata colony. Behavior of ants and *Pseudomicrodon* larvae was observed for approximately 3–5 minutes immediately after transfer as well as occasionally during the following two weeks. Larval behavior was recorded with a Handycam HDR-SR10E camcorder (Sony) five times prior to manipulation and twice post-transfer; total duration of video recording was 37 min. Ant and *Pseudomicrodon* larvae behavior was also examined for ant-syrphid interactions, as was the emergence of one adult in a laboratory ant nest.

To test whether lack of aggression by host ants is a general trait of colonies in the study area, a field experiment was conducted. From each of two bromeliad-inhabiting *Cr. limata* colonies (donors, both containing *Pseudomicrodon* larvae), five worker ants were transferred into two other ant nests (receivers, one with and one without *Pseudomicrodon* larvae). Donor colonies were 3 m apart from receiver colonies and situated in different groups of rosettes than the receiver colonies. In addition, two *Pseudomicrodon* larvae were moved between nests during this aggression test. The authors observed whether the transferred ants were treated aggressively by the host ants in each nest.

Morphological studies
Larvae were fixed in Kahle’s solution and preserved in 70% ethanol. Voucher specimens of microdontine larvae and adults, as well as associated ants, were deposited in the collection of the Native Bee Laboratory of the Federal University of Santa Catarina and Father Jesus S. Moure Entomological Collection, Department of Zoology, Federal University of Paraná, Curitiba. The morphology of the immature specimens and adults was examined with an MZ 75 stereomicroscope (Leica, [www.leica.com](http://www.leica.com)). Male genitalia were cleared in a 10% KOH solution for 36 hours, neutralized with glacial acetic acid, washed with 70% ethanol and then distilled water, and stored in glycerol.
Light micrographs were obtained with a DFC 500 digital camera attached to an MZ 16 stereo microscope (Leica). Images were captured using IM 50 software (Leica), then mounted using Automontage software (Syncroscopy, www.syncroscopy.com). Scanning electron microscopy images were obtained with a JSM-6360 LV microscope (JEOL, www.jeol.com) in order to study details of the 3rd instar larval cuticle, posterior respiratory process, marginal band, ventral surface, and pupal spiracle. Terminology is derived from Roberts (1970) and Rotheray and Gilbert (1999) for larvae, and from Thompson (1999) for adults.

Results

Discovery of two Microdontinae species
Immature specimens of two Microdontinae species were found within ant colonies in bromeliad rosettes (Figure 1D–F, J, K) on Santa Catarina Island in southern Brazil. At least 30 larvae and 21 pupae of *P. biluminiferus* were collected, 26 of which were reared to adults. Two puparia and one pupa of a second species of Microdontinae (Microdon- tinae sp. 1) were found in a queenless nest of *Camponotus melanoticus* Emery (Hymenop- tera: Formicidae) within a rosette of *A. nudicaulis*, but could not be reared to imago. The same rosette was also inhabited by a colony of *Crematogaster limata* infested with *P. biluminiferus*. Pupae of the two microdontine species could be readily distinguished (Figure 1F, K).

Biology of *Pseudomicrodon biluminiferus*
Both in the field and in the laboratory, pupae and puparia were frequently located near the nest entrance and/or with their anterior ends directed toward the opening (Figure 1I, Videos 1, 2), while larvae were additionally scattered throughout other nest parts (Figure 1D). When dates were compiled across all years (comprising April and the months from August to February), larvae were found between early November and mid-December as well as on 16 April 2010. From the field colonies taken to the laboratory, seven *Pseudomicrodon* imagines emerged. Out of the 21 immature specimens placed in vials, 16 imagines were obtained. Two failed to expand their wings, which appears to be a laboratory effect that has been reported elsewhere (Jord- dan 1968; Akre et al. 1973, 1988). Three individuals pupated and emerged within laboratory ant nests. For those three specimens and one larva that pupated in a vial, the duration of the pupal stage was determined to last 18 days ($N = 1$) or 19 days ($N = 3$).

Imagines emerged between 20 November and mid-January (Figure 2). Live pupae were not found outside of this period. Occasional searches for microdontine brood in *Cremato- gaster* nests in the field from August to October as well as late January and February
resulted in no findings. Thirteen of 19 adults (68%) emerged before 08:00. Most remaining individuals were found later in the day (09:40 to 12:38), but since they were no longer in wing expansion posture, they might also have emerged in the early morning. One exception was an individual found at 17:00, whose pupa had been noted as being closed at 10:00. For seven individuals, the exact time of emergence could be determined or estimated as being between 05:57 and 08:25.

Emergence took between 0.5 and 1.5 min from the first appearance of the head to the first steps of the adult outside the puparium (Videos 1, 2). First signs of imminent eclosion were visible starting approximately 20 min before actual emergence (Video 2) and consisted of the following: (1) a darkening of the anterior end of the pupa, probably due to the inner surface getting wet, (2) the three anterior plates (surrounded by lines of weakness) being pushed outward several times without the head emerging clearly, and (3) movements of the adult within the puparium. The single adult that was observed eclosing within an ant nest walked straight toward the nest entrance and quickly left the nest (Video 1). After emergence, adults moved to an elevated place (a wet paper towel or a bromeliad leaf) and expanded their wings within about half an hour (Figure 1L, Video 3). Sex ratio was 8:5 (female:male) and did not differ significantly from a 1:1 distribution ($\chi^2 = .69, p = .4054, N_{total} = 13$).

**Descriptions of *Pseudomicrodon biluminiferus***

**Third instar larva.** (Figures 3A–G, 4C). Length: 6.06–6.84 mm, maximum width: 5.25–5.27 mm ($N = 2$). Hemispherical, buff in color. Cephalic segment retracted, mouthparts reduced and internal (as shown for prepupa, Figure 4F). Dorsally from metathorax to posterior end strongly convex, with a reticulate pattern formed by line arrangements of granulation radiating from numerous wart-like processes of different sizes (Figure 3A, B). Bigger processes with eight to nine rays, smaller ones with four to six. Posterior respiratory process sessile, trapezoidal in anterior
and posterior view (Figures 3C, 4C), circular basally (dorsal view). Apex of posterior respiratory process constituted by four flat emarginations, ecdysial scars oval (Figure 3C), spiracular openings indistinct (Figure 3C), cuticle rough (Figure 3D). Ventral surface flattened, ventrolateral pubescence consisting of fine setae (Figure 3G). Prolegs and crochets absent. Marginal band (Figure 3E, F): distal portion with two rows of multi-branched flattened setae, proximal portion with three to four lines of papilliform protuberances.

**Instar identification.** *Pseudomicrodon biluminiferus* has three larval instars, which can be distinguished by differences in dorsal reticulation process patterns and developmental degree of the posterior respiratory process. In the first instar, dorsal reticulation processes are scant, inconspicuous, and composed of little brown rounded shapes (Figure 4A); apex of posterior respiratory process is not differentiated in emarginations. In the 2nd instar, dorsal reticulation processes are larger and partially sclerotized (Figure 4B). Apex of posterior respiratory process is differentiated in four emarginations, which persist in the 3rd instar. In the 3rd instar, the cuticle is strongly sclerotized, having numerous wart-like processes of different sizes surrounded by line arrangements of granulation (Figure 4C). In 3rd instar larva and prepupa, the area from which each pupal spiracle emerges forms a
slightly convex disc without reticulation (Figure 4C–E).

**Pupa** (Figures 3H–I, 5A, B, D, E). Length: 6.95–9.33 mm; maximum width: 4.05–6.16 mm (N = 3). Brownish. Differing from 3rd instar larva by being firmly attached to substrate by a pair of pupal spiracles and solid sclerotisation on ventral side (Figure 5A, B, D, E). Pupal spiracles papilliform, dark brown (Figure 5A, B), tuberculate around tips, and reticulated around basal half (Figure 3H). Spiracular openings simple and situated on tubercles at the tip of that structure (Figure 3I). Opercular opening of puparium notched dorsally (Figure 5C).

**Adult** (Figures 6A–F, 7A–D). The holotype male from Espírito Santo state, Brazil, was deposited in Naturhistorisches Museum Wien. Hull (1944) described the specimen well, but omitted descriptions and illustrations of the male genitalia. The sexes are similar; both are dichoptic (Figure 6A–F), and they differ only in their genital abdominal segments. Male genitalia (N = 5): hypandrium membranous; aedeagus elongated and thin, basally globose (Figure 7D); surstylus longer than wide, concave on internal surface (Figure 7A, C), arcuate in lateral view (Figure 7B); cercus wider than long, arcuate in dorsal view (Figure 7A–C).

**Material examined.** Three 1st instar larvae, two 2nd instar larvae, two 3rd instar larvae, one prepupa, three pupae, and five puparia. Adults: eight females and five males from Santo Antônio de Lisboa, municipality of Florianópolis, Santa Catarina, Brazil.
Host identity and infestation rate

All *P. biluminiferus* larvae, pupae, and adults were obtained from nests of ants belonging to the genus *Crematogaster*. All collected ants were identified as *C. limata* ($N_{\text{colonies}} = 13$). Their nests were usually found in bromeliad rosettes with broad, short, erect leaves (Figure 1A) that formed cavities between one another (Figure 1B, D). Frequently, carton-like sheaths closed the rosettes’ upper openings (remains can be seen in Video 2 at the left edge of the video frame). Sometimes, rosettes with slender, far-projecting leaves also formed such cavities at their bases, making those plants suitable nest sites, too.

Twelve of 36 systematically examined *Crematogaster* nests contained brood of *P. biluminiferus*. Twenty-four colonies of other ant genera were found in bromeliad rosettes: 2x *Brachymyrmex*, 2x *Camponotus*, 1x *Pachycondyla*, 3x *Pheidole*, 1x probably *Azteca*; remaining records not identified because no samples were collected. Two rosettes contained signs of microdontine infestation: one *Brachymyrmex coactus* Mayr (Hymenoptera: Formicidae) nest with remains of three microdontine puparia, and *Microdontinae* sp. 1, as reported above. Almost all ant nests were found in rosettes of *A. nudicaulis* and *A. lindenii*. Only two *C. limata* nests (1–2 m distant from each other, therefore probably the same colony) with *Pseudomicrodon* pupae and larvae were detected in rosettes and dead infructescence stems of the bromeliad *Vriesea friburgensis* Mez var. *paludosa* (L. B. Smith) L. B. Smith (Poales: Bromeliaceae). Five infested ant nests were searched thoroughly for microdontine brood and were found to contain 2 (5) (numbers in parentheses include puparia), 2 (2), 0 (4), 13 (13), and 5 (5) brood items, with a mean of 4.4 (5.8).

Behavioral interactions

In the laboratory colonies, *P. biluminiferus* larvae and pupae were almost completely ignored by the ants (Videos 1, 4–6). Ants were rarely observed inspecting the *Pseudomicrodon* larvae with their mandibles and antennae, and they never behaved aggressively. At times, the ants placed their own brood upon the *Pseudomicrodon* larvae and pupae or walked over them as if they were normal nest ground. When transferred to another ant nest, *Pseudomicrodon* larvae were not attacked by resident ants, whereas foreign (but conspecific) worker ants were attacked (Video 6, Figure 1M). The single *Pseudomicrodon* imago that was observed upon emergence was not clearly attacked by ants while it left the nest. Upon emergence of the fly’s head, two workers pointed their abdomens toward it, probably depositing poison or alarm pheromone. Even when ants were walked over by the fly, they showed at most undirected alarming behavior by raising their gasters (Video 1). Twice a 3rd instar *Pseudomicrodon* larva was observed drawing an ant larva beneath its body (Videos 4, 5). Aside from this, no interactions between *Pseudomicrodon* brood and ants were observed.

Discussion

At present, the current study is the only study on pre-adult life history and host associations of Neotropical microdontine species, aside from that of Borgmeier (1923), Forti et al.
(2007), and a few mere host records where the microdontine and/or ant species mostly remained unidentified. We provide new data on morphology, development, and ecology of the myrmecophilous syrphid *P. biluminiferus* that supplement the original description (Hull 1944), which only gives morphological details of an adult male for one locality. To the authors’ knowledge, the current study is the second record of microdontine-ant associations involving bromeliads (after Pérez-Lachaud et al. 2014), and the first of microdontine larvae developing in nests of the ant species *Camponotus melanoticus* and *Crematogaster limata*.

**Biology of *Pseudomicrodon biluminiferus***

Because only relatively large 1st instar larvae were found, freshly-hatched ones were probably missed, which might have resulted in an underestimation of the period of larval occurrence throughout the year. Unfortunately, the small data set did not allow for estimation of the duration of each larval stage.

Holarctic microdontine species usually have only one generation per year, i.e., a univoltine life cycle (Akre et al. 1988). There is at least one exception (*M. fuscipennis*) with two or more generations (Duffield 1981), and several species have been assumed to also have two or more generations (Duffield and Thompson 1981). The lack of *P. biluminiferus* larvae found from mid-January to the end of February suggests either that the occasional searches during that period were not sufficient or that the adults made a pause in reproduction. If the latter is true, the life cycle of *P. biluminiferus* must be at least bivoltine. The number of generations of *P. biluminiferus* that develop per year cannot be decided without additional field data. It is unknown which developmental stages are present during the winter in southern Brazil; the lack of larvae and pupae within ant nests in the months before November suggests that it is the adults (or, less likely, eggs or 1st instar larvae) rather than large larvae as in Holarctic species (Andries 1912; Garnett et al. 1985; Akre et al. 1990). In *M. tigrinus*, the only other South American microdontine species whose biology has been studied, adults were reported to be present in the winter months May and/or June, and larvae and pupae were found during the whole year in São Paulo State, southeastern Brazil (Forti et al. 2007). This suggests that in tropical and subtropical regions, adults and/or larvae of Microdontinae might be the developmental stages that outlast climatically unfavorable times.

The observations concerning pupation and emergence concur with observations of North American and European microdontine species (Wheeler 1908; Andries 1912; van Pelt and van Pelt 1972; Akre et al. 1973, 1988; Duffield 1981; Garnett et al. 1985). Established observations of the number of brood items per nest (Duffield 1981; Schönrogge et al. 2002) and of the sex ratio being near 1:1 (Akre et al. 1973, 1988; Duffield 1981) are also reinforced by the current study. The orientation of pupae toward the nest entrance (mostly in the direction of the rosette leaf tips in the field) might be an adaptation to ease leaving the nest and reaching an elevated place for wing extension. The same seems to be the case in Microdontinae sp. 1 (Figure 1J). In microdontine species whose adults are attacked by the host ants (Andries 1912; Akre et al. 1973), this trait probably serves mainly to minimize contact with the ants.

Females of *P. biluminiferus* might search specifically for ant nests in bromeliad rosettes to oviposit. Herbivorous insects have long been known to use visual stimuli in host plant detection (Prokopy and Owens 1983; Bernays
and Chapman 1994), and it is well established that parasitoids may be attracted by their hosts’ preferred microhabitats even in absence of their hosts (Godfray 1994). In this sense, bromeliads can be considered microhabitats with high probability of microdontine hosts occurring. Assuming that bromeliads (especially their visually conspicuous inflorescences, see below) can be recognized by the syrphid’s eyes more easily than other nest sites (e.g. dead sticks, soil, or inconspicuous cavities within other plants), searching for these plant structures could be regarded as a highly effective strategy. Nevertheless, the flies might have to apply further mechanisms to find suitable host nests within large groups of bromeliad clones, e.g. using chemical host recognition cues, as reported for the hoverfly *Microdon mutabilis* (Schönrogge et al. 2008).

**Ramifications of new taxonomic placement of *Pseudomicrodon biluminiferus***

Recently, the species *Microdon biluminiferus* was transferred to the genus *Pseudomicrodon* (Reemer and Ståhls 2013). Due to this new taxonomic placement of *P. biluminiferus*, the present study constitutes another novelty. All previous records of microdontine larvae preying on immature stages of their host ants apply to species of the genera *Microdon s.s.* and *Omegasyrphus* (Reemer 2013a). *Pseudomicrodon* is the third known genus with occurrence of ant brood predation.

**Identity and occurrence of host ants***

Although not all *Crematogaster* nests containing *Pseudomicrodon* brood were identified at species level, all are considered to belong to *C. limata*. This is because no other *Crematogaster* species was found inhabiting bromeliad rosettes throughout the duration of the study, and because *C. limata* appears to be one of the most common ground-dwelling ant species at the study site (Rosumek et al. 2008, referred to as *Crematogaster* sp. 1), frequently visiting inflorescences of terrestrial bromeliads (Schmid et al. 2010).

The mere presence of microdontine brood in an ant colony does not necessarily imply that those ants are natural hosts, because ant colonies may abandon their nests (leaving microdontine brood behind), which can then be recolonized by other species (Wheeler 1906, 1910; Akre et al. 1990; Schönrogge et al. 2000, 2002). This possibility for recolonization renders many records based on single or rare findings of microdontine hosts doubtful. The current study shows that *C. limata* is indeed a valid host record for *P. biluminiferus* because: (1) this syrphid was repeatedly found in nests of *C. limata*, (2) no aggression on behalf of the ants toward the *Pseudomicrodon* larvae was observed, and (3) observations strongly indicate that the *Pseudomicrodon* larvae are predatory myrmecophiles that feed on brood of *C. limata*.

Longino’s (2003b) statement that large colonies of *C. limata* may be distributed over several small cavities within a small area concurs with our observation that frequently two to four adjacent rosettes within a group of bromeliads were occupied by *C. limata*. Queens were never found (except in one small founding colony) in the present study, so there might indeed be only one ant colony scattered over several rosettes. This should be favorable to the *Pseudomicrodon* females in case they try to relocate their maternal host colony for ovipositing, as females of *M. mutabilis* (Elmes et al. 1999) are known to do.

The association between *P. biluminiferus* and *C. limata* described here is not restricted to the bromeliad genus *Aechmea*. Considering the high density of bromeliad rosettes on the forest floor (up to 200 plants/ha, Müller and
and in the canopy, Santa Catarina Island presumably houses a large host population for local *P. biluminiferus* flies. This assumption is strengthened by the unpublished finding that 11% of *Aechmea* rosettes contained ant nests, of which 63% belonged to the genus *Crematogaster*.

Few microdontine hosts have been identified in Neotropical regions so far, and only four of them belong to the genus *Crematogaster* (Wheeler 1924; Mann 1928; Longino 2003a; present study). Regarding the high number of microdontine species in the Neotropics (202; Reemer and Ståhls 2013; Reemer 2013b), there are certainly many more microdontine-ant associations waiting to be discovered, probably many involving *Crematogaster* (100 Neotropical species; Fernández and Sendoya 2004).

**Host-parasite relationship**

The lack of *P. biluminiferus* brood in nests of other ant species suggests that the myrmecophile is specifically adapted to *C. limata*. The exception, remains of puparia in a *Brachymyrmex* nest, may be explained by colony turnover as described above. Desertion of *Pseudomicrodon* larvae by a previously-disturbed *Crematogaster* colony was once recorded in the field, which supports the assumption of colony turnover.

The indifferent behavior of *C. limata* ants toward the *Pseudomicrodon* larvae, together with the weak aggression toward *Pseudomicrodon* larvae from another ant nest, indicates that the *Pseudomicrodon* brood is well-adapted to live with this ant species. However, preliminary examinations of non-polar cuticular substances of *P. biluminiferus* larvae and their hosts suggested that there is no mimicry or camouflage of the ants’ chemical profiles by the parasites (unpublished data), contrary to what was reported for two North American microdontine species (Garnett et al. 1985; Howard et al. 1990a, b; Stanley-Samuelson et al. 1990). Instead, *Pseudomicrodon* larvae, pupae, and newly-emerged adults might employ a strategy called “chemical insignificance” (Lenoir et al. 2001, 2012), meaning that the animals are not detectable as aliens due to the absence (or very low amounts) of “suspicious” substances. If this is true, chemical recognition of *P. biluminiferus* by the ants might not have been the selective force that drove this syrphid into specialization with *C. limata* as host. Instead, other elements might have been involved, e.g. host localization. However, the cuticular chemistry of this *Pseudomicrodon* species must be analyzed more thoroughly before confident conclusions can be drawn.

Garnett et al. (1985) reported 1st and 2nd instar larvae of North American microdontine species being carried by host ants when disturbed and described a specific “cocoon mimicry” actively performed by microdontine larvae, revealing a behavioral adaptation of the parasites to their hosts. There were no observations of ants carrying *Pseudomicrodon* brood in the present study, even though colonies containing those parasites were frequently disturbed in the field. The large 3rd instar was mainly recognized, and occasionally 2nd instar larvae. The fact that small *Pseudomicrodon* larvae were never found might be explained not only by absence of these stages but alternatively by the existence of similar brood mimicry, causing 1st instar larvae to be overlooked upon inspection of ant colonies. The former case seems to be more likely because numbers of *Pseudomicrodon* larvae did not increase in colonies kept in the laboratory for weeks. Either way, occurrence and behavior of the small stages of *P. biluminiferus* deserve closer examination.
The single observation of a newly-emerged adult leaving a laboratory nest undisturbed by the surrounding ants corresponds to a similar case described by Wheeler (1908). However, Andries (1912) and Akre et al. (1973) observed that microdontine adults were killed immediately after emergence, and Wheeler (1908) reported that they were attacked later during expansion of their wings. Whether this is the case in *P. biluminiferus* could not be determined because no contact between adults and host ants was observed outside the nests.

Our observations corroborate several reports of microdontine larvae feeding on their hosts’ brood (e.g., van Pelt and van Pelt 1972; Dufﬁeld 1981; Garnett et al. 1985; Barr 1995; Pérez-Lachaud et al. 2014). Nevertheless, this is not necessarily true for all microdontine species that have also been observed or assumed to feed on coccids (Borgmeier 1923, 1953), infrabuccal pellets of ants (Wheeler 1908; Donisthorpe 1927), fungi and tree sap (Garnett et al. 1985), or detritus (Forti et al. 2007). The infestation rate of *C. limata* colonies by *P. biluminiferus* (33%) lies well within those reported for Holarctic microdontine species, which are 33–50% (van Pelt and van Pelt 1972) and 16% (Akre et al. 1973).

**Single discovery of Microdontinae sp. 1**

Throughout the study period, Microdontinae sp. 1 was recorded only once. The same might be true for its putative host species, *Camponotus melanoticus*, although not all ants found in bromeliads were identiﬁed. Concordantly, in a prior extensive census of ants within bromeliads in the study area, *C. melanoticus* had been found only once in one plant of *A. nudicaulis* (Rosumek et al. 2008; A. Zillikens, J. Steiner, unpublished data). This implies either that the host and probably also the parasite occur only rarely in the study area, or that *C. melanoticus* colonies infrequently live in bromeliads. The discovery of *C. melanoticus* and *Crematogaster limata* nesting in the same bromeliad rosette, infested by different microdontine species, suggests species speciﬁcity of the microdontine-ant associations. Furthermore, it might be a case of facultative parabiosis (Weber 1943), in which two ant species coexist without being dependent on each other. Hopefully, in future studies, adults of Microdontinae sp. 1 will be reared for identifying the species and exploring this unknown microdontine-ant relationship further.

**The role of the bromeliads**

Colonies of *C. limata* have been reported as not showing speciﬁc preferences for nesting sites (Longino 2003b). So, as long as thorough examinations do not reveal the opposite for the local population on Santa Catarina Island, it must be assumed that these ants occur frequently outside of bromeliads, providing even more potential host nests for *P. biluminiferus* than the current study indicates. Depending on cues the adult ﬂies use for host location (Godfray 1994), ant nests in bromeliads might be easier to detect from a distance because of visual stimuli provided by the rosettes, frequently enhanced by the conspicuously-colored inflorescences (Figure 1A–C) that appear during the syrphid’s reproductive season (Schmid et al. 2010; Dorneles et al. 2011). Moreover, the vase-like shape of the bromeliad rosettes might concentrate olfactory signals emitted by the host colonies between the leaves, thereby impeding long-range diffusion of chemical cues while simultaneously facilitating short-range host recognition (see Schönrogge et al. 2008 for use of chemical signals by *M. mutabilis*). The possibility of bromeliads playing a major role in this microdontine-ant association surely warrants further investigation.
Conclusions
The biology of *P. biluminiferus* is largely similar to that of species in Holarctic regions, from general biology to predatory behavior and host specificity. However, two traits of this syrphid may differ from established knowledge: the overwintering developmental stage (presumably adults) and the (probably specific) association with bromeliads. The latter, together with the discovery of Microdontinae sp. 1, adds two more cases to the small list of microdontine species associated with ants that reliably nest in certain types of plants (Wheeler 1924; Hocking 1970). Due to the high number of known microdontine species, especially from the Neotropics, and the low number of studies on microdontine-ant associations in this region, there is a high potential for discovery of other relationships (e.g., with parasitoids of Microdontinae, as reported recently by Hansson et al. 2011). This indicates that the ecologies of the inconspicuous microdontine flies and the well-studied ant family Formicidae have not been sufficiently investigated to obtain a deep understanding of this interesting host-parasite system. We hope that the data provided in the current study will encourage future research in this area.

Acknowledgements

We thank F. Christian Thompson for identifying *Pseudomicrodon biluminiferus* and Menno Reemer for providing holotype pictures; Rosângela Borges Freitas of Centro de Microscopia Eletrônica (CME) of Universidade Federal do Paraná for helping with the S.E.M. photographs; Vitor Antonio Nardino and project *Taxon line* – Network of Biological Collections from Paraná State – for helping with the automontage of images; Jacques Delabie and Eduardo Cereto for help with ant identification; late Maike Hering de Queiroz and Diomário de Queiroz for access to the study site; Erhard Strohm for access to GCMS; Simone Langner for indispensable help; G. W. Elmes and two anonymous reviewers for their fruitful comments on the manuscript. Research was authorized by IBAMA, permits 090/2005 and 260/2006, and by SISBIO, permits 12486-1, 12826-1, and 12826-2. This study is part of the projects “Internal dynamics of rain forests: specificity of animal-plant interactions” (BMBF, Germany, 01LB0205A1) and “Importância das bromélias para a manutenção da diversidade da fauna associada na Mata Atlântica” (CNPq, Brazil, 690143/01-1) within the Brazilian-German program “Mata Atlântica”. Volker S. Schmid, Anne Zillikens, Rafael Kamke, and Josefina Steiner acknowledge financial support by BMBF and CNPq within this program. Mírian N. Morales was supported by "Programa de Capacitação em Taxonomia" (PROTAX-CAPES, Brazil, 562257/2010-1).

References


Elmes GW, Barr B, Thomas JA, Clarke RT. 1999. Extreme host specificity by Microdon mutabilis (Diptera: Syrphidae), a social parasite of ants. Proceedings of the Royal


Reemer M. 2013a. Review and phylogenetic evaluation of associations between Microdontinae (Diptera: Syrphidae) and ants (Hymenoptera: Formicidae). *Psyche* 2013, Article ID 538316. DOI: 10.1155/2013/538316

Reemer M. 2013b. Taxonomic exploration of Neotropical Microdontinae (Diptera:


