Sexual communication in day-flying Lepidoptera with special reference to castniids or ‘butterfly-moths’

Víctor Sarto i Monteys\textsuperscript{2,3,*}, Carmen Quero\textsuperscript{1}, María Carmen Santa-Cruz\textsuperscript{4}, Glòria Rosell\textsuperscript{5}, and Angel Guerrero\textsuperscript{1,*}

\textsuperscript{1}Department of Biological Chemistry and Molecular Modelling, IQAC (CSIC), Jordi Girona 18. 08034 Barcelona, Spain. E-mail: agpqob@cid.csic.es; cqlqob@cid.csic.es
\textsuperscript{2}Institute of Environmental Science and Technology (ICTA), Entomology, Plants and Health - Room Z/141, Building Z – ICTA-ICP, 08193 Bellaterra (Barcelona), Spain. E-mail: victor.sarto@uab.cat
\textsuperscript{3}Department of Agriculture, Livestock, Fisheries, Food and Environment (DAAM), Catalonian Government - Service of Plant Health, Avda. Meridiana, 38, 08018 Barcelona, Spain.
\textsuperscript{4}Department of Cell Biology, Physiology and Immunology, Universitat Autònoma de Barcelona. 08193 Bellaterra (Barcelona), Spain. E-mail: Carmen.Santa-Cruz@uab.cat
\textsuperscript{5}Department of Pharmacology and Therapeutic Chemistry (Unit Associated to CSIC), Faculty of Pharmacy, University of Barcelona, Av. Diagonal s/n, 08028 Barcelona, Spain. E-mail: grpqob@cid.csic.es

Running head: Sexual communication in day-flying Lepidoptera

*Correspondence should be addressed to either author: E-mail: agpqob@cid.csic.es. Tel.: +34-93-4006120; Fax: +34-93-2045904; E-mail: victor.sarto@uab.cat. Tel.: +34-93-5868647 (ext. 8647); Fax: +34-93-5813331
Abstract

Butterflies and moths are subject to different evolutionary pressures that affect several aspects of their behaviour and physiology, particularly sexual communication. Butterflies are day-flying insects (excluding hedylids) whose partner-finding strategy is mainly based on visual cues and female butterflies having apparently lost the typical sex pheromone glands. Moths, in contrast, are mostly night-flyers and use female-released long-range pheromones for partner-finding. However, some moth families are exclusively day-flyers, and therefore subject to evolutionary pressures similar to those endured by butterflies. Among them, the Castniidae, also called ‘butterfly-moths’ or ‘sun-moths’, behave like butterflies and, thus, castniid females appear to have also lost their pheromone glands, an unparallel attribute in the world of moths. In this paper, we review the sexual communication strategy in day-flying Lepidoptera, mainly butterflies (superfamily Papilionoidea), Zygaenidae and Castniidae moths, and compare their mating behaviour with that of moth families of nocturnal habits, paying particular attention to the recently discovered butterfly-like partner-finding strategy of castniids and the fascinating facts and debates that led to its discovery.

Keywords: Castniidae; *Paysandisia archon*; butterflies; Zygaenidae; chemical communication; mating behaviour.
Introduction

Lepidoptera is one of the most diverse insect groups with currently about 160,000 described species (Kristensen et al., 2007; van Nieukerken et al., 2011), although the total number of extant species is estimated to be around half a million (Kristensen et al., 2007). Within this vast group of insects and until the late 1980s, only two basic partner-finding strategies pertaining to ‘butterflies’ and ‘moths’ were known. In short, male butterflies used their vision to detect conspecific females at some distance and to pursue them. Female butterflies, in turn, had no sex pheromone glands in their ovipositors and therefore did not release any long-range pheromone to attract males. In contrast, male moths used their olfaction system to detect females at some distance because the latter release long-range pheromones* from their pheromone glands. Once together and in close courtship interactions, males (butterflies and moths), and in some cases also females, released close range pheromones or ‘scents’ that facilitated or prevented the last courtship steps leading to copulation. The butterflies, all diurnal except the moth-like hedylids, simply used vision to find mates in their sunlit environment with no need to produce long-range sex pheromones. The mostly nocturnal moths, in turn, kept the so-called “female calling plus male seduction” strategy, which implied the production of long-range sex pheromones. Table 1 summarises the partner-finding strategies of nocturnal and diurnal lepidopteran groups.

It must be mentioned, however, that three other partner-finding strategies have been described in night-flying moths (Hallberg & Poppy, 2003), although their occurrence is rare: (1) mutual calling in the noctuid Trichoplusia ni (Hübner) where both sexes “call” (Landolt & Heath, 1989), (2) reverse calling in the Pyralid rice moth Corcyra cephalonica (Stainton), where the

*Note: Long-range sex pheromones are generally emitted from epidermal glands in membranous areas of the ovipositor, the ancestral state in ditrysian Lepidoptera (Hallberg & Poppy, 2003).
male emits a pheromone at a distance, and the female responds by releasing a pheromone at close-range and induces the male to copulate (Zagatti et al., 1987; Hall et al., 1987), and (3) “lekking”, where several males gather together in a group (the lek) to which females are attracted by the male-produced pheromones, and mating takes place within the lek. Lekking behaviour has been reported in Hepialidae, Pyralidae and Arctiidae (Hallberg & Poppy, 2003). A good understanding of the above-mentioned strategies is important in natural resource management, not only for Lepidoptera of economic importance but also for endangered species and for those living in threatened habitats. The case of the Gondwanan family Castniidae, also called ‘butterfly-moths’ or ‘sun-moths’, is particularly exemplary in this respect. In the Neotropics, many of them live in threatened habitats since their boring larvae depend on tree-dwelling forest plants; however, a few species have adapted to boring into crop plants introduced by man, such as sugarcane, banana and African oil palm, subsequently becoming important pests of such crops (Sarto i Monteys & Aguilar, 2005). One of them, *Paysandisia archon* (Burmeister) was introduced into Europe (Spain) in the mid-1990s to spread eastwards to Bulgaria, Greece and Cyprus and become a serious pest of many palm species (Sarto i Monteys & Aguilar, 2005). In Australia, less than 50 castniid species occur, all included in the genus *Synemon* Doubleday (Edwards et al., 1998). *Synemon* larvae feed underground on the roots and rhizomes of grasses and sedges, and suffer a drastic reduction in their populations because of the clearing or modification of vast areas of native grasslands, woodlands and heathlands across Australia, therefore requiring urgent protection measures (Douglas, 2004). In this respect, knowing in detail how a species communicates sexually may give resource managers significant clues to either control or protect any specific endangered population. This review deals with the sexual communication in day-flying Lepidoptera, either butterflies or moths with diurnal habits, paying particular attention to castniids and to their recently suggested butterfly-like partner-finding strategy. We have also
included Zygaenidae moths (genus *Zygaena* Fabricius) because they display a dual partner-finding strategy between the castniids/butterflies and the other day-flying moths.

**The Butterflies (Superfamily Papilionoidea) and their reproductive behaviour**

Butterflies comprise 11.9% (ca. 18,800 species) of all described Lepidoptera. They are currently grouped within the taxonomic superfamily Papilionoidea, with seven families, namely Papilionidae, Pieridae, Riodinidae, Lycaenidae, Nymphalidae, Hesperiidae (skippers) and Hedyliidae (van Nieukerken *et al.*, 2011). Two recent molecular studies (Regier *et al.*, 2009; Mutanen *et al.*, 2010) strongly supported this grouping, although formerly skippers and hedyliids were placed in separate superfamilies and the other five families were grouped into only one superfamily. Skippers are more closely related to hedyliids than to the other butterflies (Regier *et al.*, 2009; Mutanen *et al.*, 2010), although hedyliids are mainly nocturnal and the available data (Scoble, 1986; Scoble & Aiello, 1990) suggest that their reproductive behaviour resembles that of moths. It has been suggested that the reproductive behaviour of skippers and other butterflies (but not hedyliids) may have evolved independently as an adaptation to diurnal habits (Sarto i Monteys *et al.*, 2012).

Butterflies had always been thought to be unique in their partner-finding, at first based on visual cues as mentioned above. After the pursuit flight, when the two sexes get together (i.e. in close-range interactions), males release short-range pheromones and there is mounting evidence that females may also do the same (Wiklund, 2003). However, females lack conspicuous scent organs, such as the typical sex pheromone glands, which makes the study of their chemical signals for male recognition and mating particularly difficult (for reviews see Boppré (1984), Hallberg & Poppy (2003)).

Male butterflies use basically two mating strategies, namely perching and patrolling (Scott, 1974; Wiklund, 2003). Perching males (Fig. 1) sit and wait for flying females, which actively assume the role of searching for males. Perchers are territorial, typically faithful to their
perching sites and readily willing to expel other males from their territories, largely by non-contact aerial interactions. The two ‘fighting’ males circle or hover near each other for a period of time before one of them flies away from the site. In contrast, patrolling males do not sit waiting for females, but actively search for them in places where they can be expected with a certain probability (Davies, 1978; Wickman & Wiklund, 1983; Wiklund, 2003; Kemp & Wiklund, 2004). Perching and patrolling may not be mutually exclusive and some species can perform both. Thus, in the speckled wood butterfly, *Pararge aegeria* (Linnaeus), males fight over sunspot territories on the forest ground; winners gain sole residency of a sunspot and behave as perchers, whereas losers patrol the forest in search for females (Bergman *et al.*, 2007). In other cases, a male butterfly which usually patrols might behave as a percher, e.g. on windy or overcast days. After female detection, perchers and patrollers pursue the female at close range, assessing her size, shape and wing pattern to be certain she is suitable for mating (Rutowski, 2003; Warrant *et al.*, 2003; Wiklund, 2003). At this close range male butterflies release pheromones that convey information to the females, inducing them to respond (mate or reject). Such male scents are produced and/or disseminated in special structures, the most common being alar androconia, i.e. specialised male scales located on the forewings, hindwings or both, and ‘hairpencils’, modified scales present on wings or the abdomen.

Male sex pheromones (MSPs) in butterflies have long-been thought to be vital in courtship, mate-choice or acceptance by females (sexual selection), species isolation and/or recognition (Boppré, 1984; Costanzo & Monteiro, 2007). In this respect, it is noteworthy that in the nymphalid butterfly *Bicyclus anynana* (Butler) the MSP composition changes along the insect life span, a signal which may be used by the insect for male identity and male age (females prefer to mate with middle-aged rather than younger males) (Nieberding *et al.*, 2012).
Burnets and Forester moths (Family Zygaenidae) and their reproductive behaviour

The zygaenids comprise four subfamilies and about 1,000 described species worldwide (Tarmann, 2004; van Nieukerken et al., 2011). With few exceptions, e.g. the nocturnal *Zygaena nocturna* Ebert and some related species, they include typically day-flying moths with a slow, fluttering flight. Their partner-finding strategy corresponds to the typical pattern for moths, with females calling males by releasing long-range sex pheromones (Subchev, 2014). Their sex glands are located at the tip of the abdomen (between segments 8-9, as usual in moths) (Fig. 2A-B), or on the anterior parts of tergites 3-5 of the abdomen, as found widespread in the subfamily Procridinae (Hallberg & Subchev, 1997).

Visual cues are also important in the mating behaviour of zygaenids, although only in the short-range phase of the courtship. Thus, in the six-spot burnet, *Zygaena filipendulae* (Linnaeus), the long-range attraction of males is mediated by female-released pheromones, but when the flying male is within ca. 50 cm range, then visual cues determine the rest of the courtship (Zagatti & Renou, 1984). Also, in the vine bud moth *Theressimina ampelophaga* (Bayle-Barelle) (Procridinae) males attracted to a synthetic sex pheromone dispenser displayed more copulation attempts when a female model (visual stimulus) was attached to the dispenser (chemical stimulus) (Toshova et al., 2007). It is uncertain whether optical cues play a significant role in the rare nocturnal zygaenids, like *Z. nocturna*, since males were found to reach calling females in the dark, mostly between 21 and 23 h (A. Hofmann pers. com.).

More surprising is the dual partner-finding strategy shown by the five-spot burnet, *Zygaena trifolii* (Esper) (Naumann, 1988; Prinz & Naumann, 1988). The females have typical sex pheromone glands that release pheromone to attract males in late afternoon. In the morning, however, they rest atop grasses close to where their cocoons were spun, do not release pheromones and can be found by males using optical cues exclusively (female wing pattern, spot colouration and specimen size). In late afternoon, the females move down into the
vegetation, where they would not be easily spotted by flying males, and release the pheromone.

The likely evolutionary advantages of the dual partner-finding strategy have been reported (Naumann et al., 1999). Although it is likely that the dual strategy may be present in other species of the subgenus Zygaena, it is not well-established how widespread this strategy is among other European Zygaeninae. In this context, Hofmann & Kia-Hofmann (2010) noted that the optical cues used by males of Z. trifolii during the morning and occasionally leading to ‘morning copulae’, cannot be considered as a general strategy and may vary from species to species depending on ecological circumstances (e.g. altitude, semi-desert, woodland). In this respect, behavioural studies carried out on Z. niphona Butler (Koshio, 2000) and Z. fausta (Linnaeus) (Friedrich & Friedrich-Polo, 2005) revealed that these species did not show the dual partner-finding strategy but only the widespread combined chemical and optical afternoon strategy, as described above for Z. filipendulae.

Notwithstanding, the discovery of the above-mentioned dual strategy in Z. trifolii is very significant from an evolutionary point of view because it was the first documented case in which a day-flying moth was not using long-range pheromones for partner-finding, at least in the morning.

The Castniids or ‘Butterfly-moths’ (Family Castniidae) and their reproductive behaviour

The Castniidae are day-flying, brightly coloured and median/large-sized moths, occurring in the Neotropics, SE Asia and Australia, with only about 110 species described (Edwards et al., 1998). They are currently grouped within the superfamily Cossoidea, with seven families, namely Brachodidae (little bear moths), Cossidae (cossid millers or carpenter millers), Dudgeoneidae, Metarbelidae, Ratardidae (Oriental parnassian moths), Sesiidae (clearwing moths) and Castniidae (van Nieukerken et al., 2011). Initially, Minet (1991) had placed the
Castniidae in the superfamily Sesioidea together with Sesiidae and Brachodidae, but recent molecular studies grouped the Sesioidea with some Cossoidea in a large, near-monophyletic (but internally unresolved) assemblage that included Cossoidea, Sesioidea and Zygaenoidea (Regier et al., 2009; Mutanen et al., 2010). Many species in this heterogeneous group are diurnal.

Castniids are interesting Lepidoptera in the following respects:

(1) The Neotropical species of castniids remarkably mimic many butterflies living in the same area in form, colours and habits, and form a truly Batesian mimicry association (Miller, 1986). The levels of mimicry between butterflies and castniids, two groups of phylogenetically-distant lepidopterans, are unparalleled in the order Lepidoptera and this has granted to castniids the term ‘butterfly-moths’.

(2) Castniid males are territorial and display perching behaviour as butterfly males (Sarto i Monteys et al., 2012), and references therein), which is also an unparalleled trait in moths.

(3) Most importantly, and in contrast to other known moths including day-flying moths, castniid females appear to have lost their abdominal pheromone glands, so that they do not release long-range pheromones to attract conspecific males. This evolutionary breakthrough was first hypothesised by Sarto i Monteys and Aguilar (2005), based on numerous field observations of the behaviour of *Paysandisia archon*, a large castniid moth (Fig. 3) which had been introduced into Europe from Argentina, as cited above, becoming a pest of palm trees (Sarto i Monteys, 2002). Experimental evidence brought forward to confirm the hypothesis that *P. archon* females do not release long-range pheromones to attract conspecific males was provided by Sarto i Monteys et al. (2012) and Riolo et al. (2014), although not without debate (Delle-Vedove et al., 2014) (see below). The fact that castniids mostly rely on visual cues for partner-finding, as most butterflies do (see above), was already noticed in the early 1900s by the German naturalist Adalbert Seitz (Seitz & Strand, 1913).
Territoriality and perching/patrolling behaviour in castniids

*P. archon* males usually perch on palm leaves or cut rachises around the trunk close to the crown (Fig. 3) (Sarto i Monteys & Aguilar, 2005). When a perching male watches another male approaching his territorial spot, he immediately takes off towards the intruder and a pursuit begins. The pursuit flight is very powerful and rapid, and the flight path is generally straight although right/left shifts may also occur (Sarto i Monteys & Aguilar, 2005). If the flying pair cross the territory of another male, this third one would immediately join the pursuit so that the flying group would now be constituted by three individuals, and so on. These pursuit flights are not long-lasting and males soon fly back to their perching spots.

Most males behave like perchers, i.e. they are faithful to a territory or spot they ‘defend’. These spots are located within palm-infested plots, where females would be flying around after emergence and detected by perching males. In Catalonia, NE of Spain, the areas of these plots are not large (usually less than 3,000 m²), and it is unclear how the territorial spots are shared by competing males, especially when infestation is high. It is likely that males with no ‘territory’ move away in search of new plots to colonise and females to mate. In this case, they would behave as patrollers, as supported by our occasional observations of lone-flying males. As in other castniids, the territoriality of *P. archon* is poorly understood, and thus several questions remain unanswered, such as: who wins the territorial spot?; how large are the territorial spots?; or what drives the likely migration of males and females to other palm plots?. Based in our observations, the mating behaviour of *P. archon* cannot be properly performed in nature unless large areas are available to the moths, and so studies carried out only in small insectaries or cages are not suitable for fully understanding the behaviour of these insects and may lead to wrong conclusions.

Do female castniids have pheromone glands in their ovipositors?
Several morphological, chemical and ethological facts combined appear to demonstrate that *P. archon* females have apparently lost their pheromone glands. These facts are the following:

(1) The territorial male behaviour described above does not support that female castniids use long-range pheromones for partner-finding, with vision playing a determinant role in this task.

(2) Hexane extracts of *P. archon* female ovipositors and other female body parts have yielded no compounds with putative pheromone activity (Acín, 2009; Sarto i Monteys *et al.*, 2012).

Also, analysis of ovipositor extracts of 1- and 24-h virgin females of *P. archon* (N=10) in hexane resulted in the identification of 24 different compounds but none of them elicited any significant GC-EAD (gas chromatography-electroantennographic detector) responses on male antennae (Riolo *et al.*, 2014). The GC-EAD system allows determination of the electrophysiological activity of every compound eluting from the capillary column when the outlet of the column is split in a specific ratio (usually 1:1) between the GC detector and the male antenna.

(3) In most Lepidoptera, when female moths adopt the ‘calling’ position, the glandular area containing the sex pheromone gland is exposed and the pheromone is released (Percy-Cunningham & MacDonald, 1987; Hallberg & Poppy, 2003). A well-defined periodicity for calling is widespread in nocturnal and diurnal moths that use long-range chemical communication (e.g. (Haynes *et al.*, 1983) and references therein). For instance, females of the nocturnal tobacco budworm *Heliothis virescens* (Fabricius) call during the period 23:30-02:30 h (Sparks *et al.*, 1979), whereas those of the artichoke plume moth *Platyptilia carduidactyla* (Riley) call mainly between 2 and 6.5 h after the onset of the scotophase (Haynes *et al.*, 1983). In the diurnal gypsy moth *Lymantria dispar* (Linnaeus), females call continuously from 10:00 to 22:00 h but some females may continue calling at night during the scotophase and early photophase (Charlton & Cardé, 1982). In diurnal burnet moths of the genus *Zygaena*, most females may call for 5 to 10 h per day (A. Hofmann pers. com.). Therefore, in diurnal moths the periodicity of pheromone release and calling appear to be not
as discrete as in the nocturnal moths, but in all cases, females expose their glandular area
during several hours to release the pheromone. Nothing similar has been observed in *P.
archon* females. We have frequently noticed that females quickly extrude/retract their
ovipositors for some seconds, but never adopt a typical ‘calling’ position that implies keeping
ovipositors extruded for a long period of time. Riolo et al. (2014) have also reported that
females perform the extrude/retract action very often throughout their lives, but it appears not
to be related to calling behaviour. These authors concluded that ovipositor extrusion might be
involved in the female physiological state (i.e. egg load) or in thermoregulation activity, as
observed in the hawk moth *Eumorpha achemon* (Drury).

(4) The antennae of castniids and butterflies are strikingly similar, with no apparent sexual
dimorphism. The antennae are the ‘noses’ of moths and butterflies and their morphology and
sensilla are suited to their needs (Hansson, 1995; Hallberg & Poppy, 2003). Moth antennae
are generally sexually dimorphic, and those of males contain a certain population of sensilla
housing olfactory receptor neurons (ORNs) sensitive to the pheromone components.
Butterflies, in turn, possess thin and clubbed antennae and display no sexual dimorphism.
They use sex pheromones only for close-range communication and therefore lack the highly
sensitive detection system found in male moths. In a Scanning Electron Microscopy (SEM)
study of male and female antennal sensilla of several day-flying Lepidoptera, namely sesiids,
butterflies (pierids and skippers) and castniids (*P. archon*), Sarto i Monteys *et al.* (2012)
concluded that *P. archon* male antennae were unsuited to detect long-range pheromones.

(5) The abdominal tip (segments 8 and 9-10) of female Lepidoptera forms a telescope-type
oviscapt, commonly called ‘ovipositor’. In most Cossoidea, the intersegmental cuticle
connecting segments 8 and 9 is long when the ovipositor is fully extended. Below that cuticle
are located the glandular epithelial cells that produce pheromones. In sesiids, which are very
closely related to castniids, such cuticle shows many buds, each topped with one thin and
curved ‘hair’ (Fig. 4) that is supposed to help release the pheromone (Tatjanskaitė, 1995).
However, SEM studies on *P. archon* ovipositors showed that the 8-9 intersegmental cuticle was devoid of such structures, and instead multiple longitudinal smooth folds could be seen, simply allowing for ovipositor expansion, as if there were no pheromone glands underneath (Figs. 5, 6) (Sarto i Monteys *et al.*, 2012). More recent histological studies confirmed this assumption as there was no evidence of pheromone gland tissues below the intersegmental cuticle of the *P. archon* ovipositor (Riolo *et al.*, 2014).

The latter five facts combined appear to clearly indicate that, as in female butterflies, *P. archon* females do not possess any abdominal gland to release a volatile pheromone to attract conspecific males, and this may likely be widespread in Castniidae. However, against this assumption, Delle-Vedove *et al.* (2014) claimed that *P. archon* females ‘call’ males using a pheromone identified as (E,Z)-2,13-octadecadien-1-yl acetate from ovipositor extracts of sexually mature females but no further details were given. They also concluded that the insect displays a “moth-butterfly hybrid” strategy relying on both chemical and visual clues. The chemical thought to be the female sex pheromone of *P. archon* had been identified in females of a number of Sesiidae, especially of the genus *Synanthedon* Hübner, and in females of the leopard moth *Zeuzera pyrina* (Linnaeus) (Cossidae) (El-Sayed, 2014). In this respect, it should be noticed that this pheromone was used in one-day field tests carried out at two sites in Catalonia to check a possible attractant effect on *P. archon* males. The tests took place in sunny days of mid-July and observations lasted continuously from 12 to 15 h, when *P. archon* males are particularly active. Three filter papers and three paper dummies depicting an adult of *P. archon* were impregnated with 1 μg of *Z. pyrina* pheromone dissolved in hexane. Such gadgets were set spaced 8 m apart on palm trunks (*Trachycarpus fortunei* (Hook.) H. Wendl. and *Chamaerops humilis* Linnaeus) within commercial gardens heavily infested by *P. archon*. At both sites not a single *P. archon* male approached to either lure suggesting that this pheromone does not attract males of this castniid (Vassiliou & Sarto i Monteys, 2014).
Mating behaviour of *P. archon* at close range

The courtship behavioural sequence of *P. archon* was first described in detail by Sarto i Monteys *et al.* (2005, 2012) as the following six consecutive steps: 1. **Localization/pursuit.** A territorial perching (or maybe patrolling) male locates a flying female that has entered his territory and pursues her. The pair fly together along the palm rows close to each other (about 10-15 cm) and at heights near the palm crowns. 2. **Alighting.** Then, the pair alight, led by the female, facing up on upright surfaces (a palm leaf or crown, the sides of a mesh tent, etc.). The female may walk shortly until reaching a spot where she can rest comfortably, folding her wings in the common noctuoid position, and if the male is accepted, she will remain still for the rest of the courtship. 3. **Orientation.** The male, which alighted a few cm below the female and has been closely following her movements, moves up and approaches to her with his wings folded. There is no male flickering. 4. **Thrusting.** While approaching the female, the male usually touches the edges of her wings with his head/antennae, sometimes inserting the antennae briefly under her wings. Also, his antennae and/or legs may also make contact with the side of the female. Both sexes keep their wings fully folded. 5. **Attempting.** The male curls his abdomen and opens up his clasping genital valvae in order to contact and grasp the female copulatory orifice to accomplish the copula. 6. **Copulation.** While in copula, both sexes stay motionless, facing up side by side, and with the male in a lower position than the female.

Recently, the courtship behaviour of *P. archon* has received further attention (Delle-Vedove *et al.* 2012, 2014; Riolo *et al.*, 2014) with both research groups providing a deep quantitative analysis of the behaviours involved (up to 14 defined by the former authors and 20 by the latter). Both groups also provided kinetic diagrams of courtship behaviour indicating, for each behavioural step, the frequency of transitions to other courtship steps. They basically confirmed the main six behavioural steps described above, including in the sequence analysis all types of behaviours displayed by both sexes during courtship. One of such behaviours was
the ovipositor extrusion. According to Delle-Vedove et al. (2014) the extrusion (1-10 times during periods of 13-48 s each before displaying another behaviour type) was synonymous to ‘calling’, i.e. females emitting a sex pheromone to attract males. In contrast, according to Riolo et al. (2014), extrusion of the ovipositor was not related to calling but possibly to the female physiological state or to thermoregulation activity, as cited above.

Other behaviours during *P. archon* courtship which deserve special mention are **antenna cleaning** (in both sexes) and **male ‘scratching’**. Females clean their antennae about 3 times more often than males, regardless of courtship outcome (Riolo et al., 2014), and because females have a higher olfactory sensory surface area in their antennae than males (Sarto i Monteys et al., 2012), this suggests that the perception of volatiles is highly important to *P. archon* females – probably more than it is for males whose antennae are unsuited for detecting long-range pheromones (see above).

Male ‘scratching’ is an interesting behaviour introduced by Frérot et al. (2013) and Delle-Vedove et al. (2014). When performed, the male walks and scratches/rubs its midlegs rapidly on the substrate, supposedly helping the release of a male pheromone produced and/or held in the midlegs (see below) and inducing the female to take-off and initiate a hovering flight. The authors, however, do not provide any evidence that such ‘scratching’ implies releasing pheromone from the male midlegs nor its unambiguous association to some kind of response by the female.

**Castniids androconia and likely role of *P. archon* male putative pheromones**

The structures presumed to be involved in the reproductive behaviour of castniid males have been poorly studied, although four types have been so far identified in the Neotropical species of the subfamily Castniinae: (1) a complex and very large abdominal (sternal) androconial organ with a brush in the hindlegs, formed by long, soft, pale scaling on the inner surface of femur, tibia and basitarsus, which supposedly helps distribute the gland secretion over the sternites in the abdomen; (2) large paronychia (i.e. bristle-like structures) on the pulvilli of
midlegs pretarsi; (3) notably enlarged midlegs basitarsi, generally (but not exclusively) in combination with large midlegs pretarsal paronychia (see Fig. 7A-D); (4) alar androconial organs located either on the underside of the forewings or the upperside of the hindwings (Jordan, 1923; Le Cerf, 1936). Whereas structures 3 and 4 seem to be common to most castniids, those individuals bearing structure 1 lack structure 2, and vice versa (Jordan, 1923); *P. archon* for instance holds structures 2, 3, and 4.

Very few reports have been found in the literature about the possible presence of sex pheromones in the Castniidae family and only concern those of females (Rebouças et al., 1999). It was not until 2012 that three putative male pheromones were reported for the first time from *P. archon* male wings (Sarto i Monteys et al., 2012). The compounds were identified as (Z,E)-3,7,11-trimethyl-2,6,10-dodecatrienal ((Z,E)-farnesal), the corresponding E,E isomer ((E,E)-farnesal), and (E,Z)-2,13-octadecadienol, which elicited significant GC-EAD responses on female antennae. Farnesals were found in the forewings and hindwings of males only (Fig. 8), although the relative amount detected in both types of wings was highly variable. The biological significance of farnesals in the male wings of *P. archon* is unknown, but it is noteworthy that both isomers of the chemical were identified in male glands located in the forewings of the rice moth *Corcyra cephalonica* and elicited walking attractancy on females (Zagatti et al., 1987). We could hypothesise that these chemicals may be used by *P. archon* females for sexual selection, as occurs in the nymphalid butterfly *Bicyclus anynana* whose females use n-hexadecanal, one specific component of the male sex pheromone, for that purpose (Nieberding et al., 2012).

In female castniids, sexual selection may also be likely influenced by their monandrous condition. Thus, it is known that most *P. archon* females behave monandrously with only a few of them (6%) mating twice, always before laying their first eggs (Delle-Vedove et al., 2012). This is probably due to their low fecundity: *P. archon* and other castniid females only lay about 110-130 eggs in their lifetime (Sarto i Monteys & Aguilar, 2005). Therefore, the
monandrous female must choose which type of males can help her reproduce successfully, and she will likely prefer virgin to non-virgin males, since the former are likely to provide bigger spermatophores with higher amounts of sperm, proteins and lipids to be used in egg production (Lauwers & Van Dyck, 2006). In the speckled wood butterfly, *Pararge aegeria*, also a territorial species, copulations with non-virgin males lasted on average 5 times longer than with virgin males, resulting in a 3 times smaller spermatophore (Lauwers & Van Dyck, 2006). The number of eggs laid and the female life span were not affected by the mating status, but there was a significant effect on the number of living caterpillars as copulations with virgin males resulted in higher larval offspring.

It is known that males from several lepidopteran families, either moths (e.g. Arctiinae-Noctuidae) or butterflies (Danainae and Ithomiinae-Nymphalidae), accumulate substances from the host plant at the larval stage as a defence mechanism against predators (Schulz *et al.*, 2004). Many of these chemicals can be subsequently used as pheromone precursors (Eisner & Meinwald, 1987; Trigo *et al.*, 1994). Farnesals are present in plants of the families Araceae, Orchidaceae, Cactaceae, Rubiaceae and others, but have not yet been found as such in palm trees (Areaceae). The latter, which are the only food plants of *P. archon* larvae, contain, however, relatively large amounts of (Z) and (E)-β-farnesenes (see f.i. Knudsen, 2002), and these compounds could be biosynthetic precursors of the farnesals found in *P. archon* through the corresponding intermediate farnesols.

The presence of (E,Z)-2,13-octadecadienol in *P. archon* was also noticed by Frérot *et al.*, (2013) in surprisingly huge amounts (µg) from male midlegs. This compound was identified by its NMR spectrum and GC-MS, and the authors suggested that the midlegs basitarsi were probably the sites of emission. This dienol is a component of the female sex pheromone or attractant of some Lepidoptera, namely some species of the family Tineidae, such as the common clothes moth, *Tineola bisselliella* (Hummel), some species of *Prochoreutis Diakonoff* & Heppner (Choreutidae), and several clearwing moths (Sesiidae), this latter
family closely related to that of castniids (El-Sayed, 2014). The role of the dienol in the chemical communication of *P. archon* is likely different to that of the farnesals. Because the alcohol triggers significant responses in male and female antennae (Sarto i Monteys *et al.*, 2012), it might act as a ‘territorial’ pheromone, i.e. males could use it to let other males know about its presence, either around where they are perching and/or when they alight close to the female in the close-range phase of the courtship.

Females, in turn, may perceive the dienol on flight during the pursuit phase of the courtship or while approaching a male territory. Since male and female *P. archon* antennae are not suited to detect long-range pheromones, as cited above, the latter option would apply only at rather short distances.

In summary, we have reviewed the partner-finding strategies of three day-flying lepidopteran groups, namely butterflies (superfamily Papilionoidea) and the moth families Zygaenidae and Castniidae, and compared their mating behaviour with that of other typical diurnal and nocturnal moth families. Day-flying moths have been subject to analogous evolutionary pressures than those of butterflies, and consequently, at least in some of them, females behave as if they had lost their pheromone glands, not releasing long-range pheromones to attract conspecific males. In fact, as in butterflies, female castniids appear to have lost their pheromone glands, an attribute with no parallel in the world of moths, and this certainly represents an evolutionary breakthrough to what has been known about sexual communication in Lepidoptera. However, as pointed out, we are still far from fully understand the chemical communication of day-flyers, particularly of castniids, and more work should be devoted to unveiling the function of the diverse structures allegedly involved in their reproductive behaviour and the specific role of their sex pheromones. Knowledge of the chemical communication of day-flying Lepidoptera is also important in natural resource management, both for control of new invasive species, like *P. archon*, or to protect specific endangered populations.
Acknowledgments

We thank CSIC for a postdoctoral contract to C.Q. We thank J.-B. Peltier for help with literature and *P. archon* cocoons, L. Aguilar for providing information on *P. archon*’s infested plots in Catalonia. We also acknowledge A. Hofmann and G. Tarmann for helpful literature and/or comments provided on Zygaenidae reproductive behaviour. This work was partially supported by MINECO (AGL2012-39869-C02-01) with assistance from the European Regional Development Fund.
Figure captions

**Fig. 1.** Perching specimen of the large skipper butterfly *Ochlodes sylvanus* (Esper) (Hesperiidae). (Photo V. Sarto i Monteys).

**Fig. 2.** *Zygaena escalerai* Poujade (Zygaenidae, Zygaeninae): (A) Calling female, (B) Closeup of ovipositor at calling, showing expanded intersegmental membrane between segments 8 and 9. (Photos A. Hofmann).

**Fig. 3.** Perching male of *Paysandisia archon* (Castniidae). (Photo V. Sarto i Monteys).

**Fig. 4.** Clearwing moth *Synanthedon* sp. ovipositor (Sesiidae): (A) 20 µm cross-section (seen with light microscope) of intersegmental membrane between abdominal segments 8 and 9 of the ovipositor (top is dorsal). In, integument; Pr, proctodeum; Mu, muscles; Ap, posterior apodema or posterior ‘apophysis’; Tr, tracheae. Scale bar: 100 µm. Many cuticular buds cover the whole intersegmental cuticle, each topped with one thin and curved spinelike process, supposed to help release the pheromone. (B) closeup showing the cuticular buds (arrowheads point to some of them) of the integument. Scale bar: 30 µm. (Photos M.C. Santa-Cruz).

**Fig. 5.** *Paysandisia archon* ovipositor. (A) ventral view of partly retracted ovipositor (treated with potassium hydroxide 10%). (B) side view of fully everted ovipositor (segment 9+10, intersegment 8-9, segment 8) plus intersegment 7-8 and part of segment 7 (in ethanol 70%). Left side is dorsal; right side is ventral. Black arrows show from top to bottom the 9 (+10), 8 and 7 abdominal segments; blue arrows show the intersegmental membranes between segments 8-9 (top) and 7-8 (bottom). Left posterior and anterior ‘apophysis’ or apodemas are also indicated. Scale bars for A, B are 1 and 2 mm, respectively. (Photos (A) M.C. Santa-Cruz; (B) V. Sarto i Monteys).

**Fig. 6.** SEM images of *Paysandisia archon* ovipositor. (A) intersegmental membrane between segments 8 and 9 showing a smooth surface (x 35). (B) closeup, 700x showing multiple longitudinal smooth folds, allowing for extra ovipositor expansion. (C) closeup, 5000x.
Unlike sesiids, the 8-9 intersegmental membrane of *P. archon* ovipositor is devoid of any cuticular buds. Scale bars for A, B, C are 500, 25, and 4 µm, respectively. (Photos V. Sarto i Monteys).

**Fig. 7.** Midleg of *Paysandisia archon* female (A) and male (B). Side view of full midleg (excluding coxa and trochanter), tibia and tarsus are seen lateroventrally. The 1st tarsomere (basitarsus) is not enlarged and appears smaller than the tibia in females (A), while in males the 1st tarsomere is notably enlarged (B). Closeup side view of pretarsal segment showing the two pulvilli with no paronychia in females (C) and forming large paronychia in males (D). Scale bars for A, B are 2 mm and for C, D 0.4 mm. (Photos V. Sarto i Monteys).

**Fig. 8.** GC-MS analysis (chromatogram A,B) of an extract of forewings (A) and hindwings (B) of *Paysandisia archon* males showing the presence of Z,E-farnesal (retention time 13.95 min) and E,E-farnesal (14.26 min) and the corresponding mass spectra (C, D respectively). Peak at retention time 12.66 min of the chromatogram corresponds to the internal standard (IS) (Z)-9-tetradecenol.


Miller, J.Y. (1986) The taxonomy, phylogeny, and zoogeography of the Neotropical moth subfamily Castniinae (Lepidoptera: Castnioida: Castniidae), University of Florida U.M.I.


Table 1. Generalized comparison of partner-finding strategies, pheromone uses and other related traits in nocturnal and diurnal lepidopteran groups

<table>
<thead>
<tr>
<th>Lepidoptera group</th>
<th>FLIGHT PERIOD</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>NOCTURNAL^{1,2}</td>
</tr>
<tr>
<td></td>
<td>Most moth families</td>
</tr>
<tr>
<td>Partner-finding strategy</td>
<td>Moth-like</td>
</tr>
<tr>
<td>Female-released long distance pheromones to attract males</td>
<td>Yes</td>
</tr>
<tr>
<td>Presence of female pheromone glands in the ovipositor^{4}</td>
<td>Yes</td>
</tr>
<tr>
<td>Perching and/or patrolling behaviour in males</td>
<td>No</td>
</tr>
<tr>
<td>Main sense used by males to detect females at long distance</td>
<td>Olfaction</td>
</tr>
<tr>
<td>Clubbed antennae + very reduced olfactory sensory surface on antennae</td>
<td>No</td>
</tr>
<tr>
<td>Male-released pheromones in close range courtship</td>
<td>Yes</td>
</tr>
</tbody>
</table>

^{1}Some species within a 'nocturnal' or 'diurnal' family have adapted to fly in the twilight or prefer to fly in shaded environments
^{2}Some species within a typical 'nocturnal' family have adapted to fly in day time
^{3}Except *Zygaena nocturna* and some related species
^{4}Exceptionally some groups have ovipositor-like pheromone glands in other parts of the abdomen or thorax
Figure

A

5 (pretarsus)

4

3

2

1 (basitarsus)

Tarsus (includes 5 tarsomeres)

Femur

Tibia

P. archon
female Midleg

B

5 (pretarsus)

4

3

2

1 (basitarsus)

Tarsus (includes 5 tarsomeres)

Femur

Tibia

P. archon Male
Midleg

C

1 Arolium

2 Pulvilli (with no paronychia)

2 Claws (-Ungues)

P. archon Female
Midleg pretarsus

dorsal side

ventral side

D

2 Pulvilli (with large paronychia)

1 Arolium

2 Claws (-Ungues)

P. archon Male
Midleg pretarsus

dorsal side

ventral side