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## Antennal sensillar equipment in closely related predatory wasp species (Hymenoptera: Philanthinae) hunting for different prey types

*Équipement sensoriel des antennes dans des espèces proches de guêpes prédateurs (Hymenoptera: Philanthinae), qui chassent des proies différentes*Carlo Polidori<sup>a,\*</sup>, Alberto Jorge García<sup>b</sup>, José L. Nieves-Aldrey<sup>a</sup><sup>a</sup> Departamento de Biodiversidad y Biología Evolutiva, Museo Nacional de Ciencias Naturales, C/José Gutiérrez Abascal 2, 28006 Madrid, Spain<sup>b</sup> Laboratorio de Microscopía, Museo Nacional de Ciencias Naturales, C/José Gutiérrez Abascal 2, 28006 Madrid, Spain

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## ABSTRACT

Despite its potential value in phylogenetic and ecological studies, the morphology of antennal sensilla has rarely been compared quantitatively within the Apoidea. Here, through a scanning electron microscopy analysis, we provide an inventory of different types of antennal sensilla and compare their morphology across 10 species of predatory wasps (Crabronidae: Philanthinae) including species that hunt exclusively either on beetles or on bees to feed their larvae. A sensilla-free area was found on the apical flagellomer of all but two species, and its shape and size appear to be useful for separating Philanthini from Cercerini within the subfamily. A total of eight types of sensilla (sensilla placoidea, sensilla basiconica, two types of pit organs, sensilla coelocapitula and three types of sensilla trichoidea) were found in all species, and an additional rarer type (grooved peg sensilla) was found only in three bee-hunting species and for first time in the genus *Cerceris*. Certain morphological features confirmed the separation of the apoid wasps from the rest of the Apoidea (i.e., bees). A cluster analysis based on the sizes of the different types of sensilla suggested that, overall, sensilla morphology is not a useful taxonomic tool, and thus, other factors likely determine interspecific variability. One candidate factor is the prey type, given some differences in the presence, density, size and distribution of certain types of olfactory sensilla between beetle-hunters and bee-hunters. This hypothesis needs to be further tested quantitatively using a larger species set, more individuals *per* species, additional sensilla features, and a correction for phylogeny.

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## 1. Introduction

The antennae of insects play an important role in food selection, oviposition site selection, and intra- and interspecific communication [1–4]. This sensory capacity of the antennae is achieved through the antennal sensilla, which are the most important receptor organs. Among the Hymenoptera (bees, wasps and ants), these sensilla show

considerable diversity in terms of their morphology, density, and function (olfactory, gustatory, CO<sub>2</sub> sensing, hydrothermal and tactile) [5–7].

There is important variability in the incidence, density, and distribution of different types of sensilla not only among species but also, to different extents, between sexes within a species [8–10]. In addition to phylogenetic effects on such variability, the morphology, numbers and distribution patterns of sensilla may be the product of interacting selection pressures related to feeding and foraging habits, habitats, mating systems, and sociality [11–16].

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Within the Hymenoptera, very few studies have been performed on the comparative morphology of the antennal sensilla in Apoidea, and most of these have concerned bees (s.s. i.e., pollen-foraging Apoidea) [8,15,17–19]. This broad group also includes one large family of wasps, the Crabronidae, which have been the focus of a great number of behavioural and ecological studies in the last 100 years [20–22]. However, the diversity and morphology of the antennal sensilla have been investigated in detail in only five species in this family thus far [23,24], though ultrastructural features of a few olfactory sensilla types have been published for 10 additional species [25,26]. Briefly, the antennae of Crabronidae bear at least eight types of sensilla on their flagellomers, being particularly abundant on the more distal flagellomers, whose general morphology does not differ greatly from that found in bees, i.e., their closest relatives within the Aculeata [21]. In all species, olfactory sensilla, such as the sensilla placoidea and large sensilla basiconica, pit organs (potentially including CO<sub>2</sub> and hygrothermal receptors), and three-four types of sensilla trichoidea (possibly with a gustatory or tactile function) are the most abundant. Their density and distribution appear to differ to some extent both among species and between sexes [23,24]. However, the available data on sensilla morphology are scattered among subfamilies of Crabronidae, so we still do not know the extent of the variability between closely related species (within subfamilies, tribes and genera), particularly when these species differ in terms of important behavioural and ecological traits, such as prey selection.

In fact, one of the main characteristics of apoid wasps is their relatively high specialisation regarding prey use at the species and often at genus level. A given species of wasp hunts only for a restricted group of prey species (rarely only one) belonging to a single order (and often to a single family) of insects or spiders [21]. In some cases, entire crabronid subfamilies are composed of species hunting for the same insect order, while in other subfamilies, different genera hunt for prey belonging to different orders [20,21].

From this point of view, the digger wasp subfamily Philanthinae appears to be an adequate group in which to carry out comparative morphological analyses, as it includes species hunting exclusively either for adult beetles or adult bees to feed their larvae, and this

resource-use diversity is even retained within one genus (*Cerceris*) [27].

The aim of this study is to provide an inventory of the antennal sensilla of 10 species of philanthine wasps spanning three genera: *Cerceris* (including both bee-hunters and beetle-hunters), *Philanthus* (all bee-hunters) and *Trachypus* (all bee-hunters). The genera *Philanthus* and *Trachypus* are more closely related (tribe Philanthini) and are believed to be phylogenetically more primitive, while the genus *Cerceris* is considered more derived (tribe Cercerini) [28]. Hymenoptera are believed to represent the basal prey for this subfamily and, in turn, most likely for the genus *Cerceris* [27]. To date, the antennal sensillar equipment of philanthine wasps has been studied in detail in only two species (*Cerceris rybiensis* L. and *Philanthus triangulum* F.) [23,24], both of which are bee-hunting species. We compared the external morphology of the antennal sensilla of eight additional species (and re-analysed *C. rybiensis* and *P. triangulum*), further searching for potential differences between beetle-hunters and bee-hunters.

## 2. Materials and methods

### 2.1. Study organisms and examined material

The subfamily Philanthinae (Hymenoptera: Crabronidae) is one of the richest taxa (including approximately 1140 species worldwide, distributed in eight genera) among the so-called digger wasps [29]. These species are solitary or primitively social predatory Hymenoptera that nest in the ground, digging multi-chambered nests where prey are placed and eggs are laid [21]. The developing offspring feed upon such prey, while adult wasps are nectar feeders on flowers [21,22].

For the present study, we selected females of 10 species: two from the genus *Philanthus*, one from the genus *Trachypus* and seven from the genus *Cerceris* (two bee-hunters and five beetle-hunters) (Table 1). Specialisation in terms of prey use varies among these species, ranging from the species-specific hunter *P. triangulum* and the family-specific hunters *C. arenaria* and *C. bupresticida* to more generalist species hunting for three or more families, such as *C. rubida*, *P. pulchellus* and *T. denticollis* (Table 1). All of the species analysed are solitary, with the exception of

**Table 1**

List of studied wasp species, with associated head width, prey, references for prey records and origin of samples.

Species	Head width (mm) ( <i>n</i> = 2)	Prey	References for prey	Sample
<i>Cerceris arenaria</i> (L.)	3.5 ± 0.1	Coleoptera (Curculionidae)	[45]	Field (Italy)
<i>Cerceris bupresticida</i> Dufour	3.4 ± 0.2	Coleoptera (Buprestidae)	[54]	MNCN
<i>Cerceris quinquefasciata</i> (Rossi)	2.7 ± 0.1	Coleoptera (Curculionidae, Chrysomelidae)	[55]	Field (Italy)
<i>Cerceris rubida</i> (Jurine)	2 ± 0.1	Coleoptera (Chrysomelidae, Curculionidae, Nitidulidae, Phalacridae, Bruchidae, Scolytidae)	[46]	Field (Italy)
<i>Cerceris rybiensis</i> (L.)	2.8 ± 0	Hymenoptera (Halictidae, Apidae, Andrenidae)	[55]	MNCN
<i>Cerceris sabulosa</i> (Panzer)	2.8 ± 0.2	Hymenoptera (Halictidae, Apidae, Andrenidae, Colletidae)	[55]	Field (Germany)
<i>Cerceris tuberculata</i> (Villers)	5 ± 0.1	Coleoptera (Curculionidae)	[55]	MNCN
<i>Philanthus pulchellus</i> Spinola	2.7 ± 0.1	Hymenoptera (Halictidae, Apidae, Crabronidae)	[56]	Field (Spain)
<i>Philanthus triangulum</i> Fabr.	3.4 ± 0.2	Hymenoptera (Apidae: <i>Apis mellifera</i> )	[20]	Field (Italy)
<i>Trachypus denticollis</i> Spinola	4 ± 0.2	Hymenoptera (Halictidae, Apidae, Colletidae, Crabronidae)	[57]	Field (Chile)

*C. rubida*, which is one of the rare cases of social Philanthinae worldwide and the only case reported for Europe [30].

The examined material comes either from individuals collected in the field (preserved in 95% ethanol) from a variety of European localities in recent years (2005–2010), or from pinned specimens from the entomology collection at the Museo Nacional de Ciencias Naturales (MNCN) in Madrid (Spain). Antennae of ethanol-preserved individuals were dried and then mounted on conventional microscopy stubs in either a dorsal, ventral or lateral orientation, then gold-coated. Whole pinned individuals from the museum collection were subjected to scanning electron microscopy without further preparation. Both the right and left antennae of two females *per* species were analysed.

## 2.2. Scanning electron microscopy (SEM)

SEM images were obtained using a ESEM QUANTA 200 microscope from the FEI Company (Oregon, USA) located at MNCN. We operated with three levels of vacuum (high vacuum and low vacuum 0.08 to 0.60 torr and environmental 1 to 20 torr). The low-vacuum mode (resolution: 3.0 nm at 30 kV [SE], 4.0 nm at 30 kV [BSE], and < 12 nm at 3 kV [SE]) allows inspection at a high resolution and supports the analysis of non-conductive hydrated samples in their original condition with both the large field detector (LFD), as it is close to the sample and will avoid loss of electrons, and the backscatter detector (BSED, backscatter electron detector). This method was used for the specimens obtained from the museum collection. High vacuum conditions (resolution: 3.0 nm at 30 kV [SE], 10 nm at 3 kV [SE], and 4.0 nm at 30 kV [BSE]) were used for the field samples covered with sputtered gold, providing a better resolution in the obtained electronic images. The accelerating voltage was 26 kV; the high and low vacuum were 0.40–0.50 torr; and the working distance was 10 mm.

## 2.3. Sensilla classification and morphometric data

For the inventory of the sensilla, we primarily followed the sensillar classification of Esslen and Kaissling [31] and Ågren [17], based on morphological characters. This classification should be considered as preliminary because the internal structure and the function of the different types of sensilla are still not fully known [5]. The flagellomeres were designated Fl<sup>1</sup> to Fl<sup>10</sup> in a proximal to distal direction.

Based on the SEM images taken of each entire antennal flagellomer, we qualitatively studied the distribution of sensilla types. We did not attempt to quantify the total number of sensilla because in some cases, it was difficult to photograph all flagellomeres in a perfect dorsal, ventral or lateral view for each species (due to the often twisted and bent shape of the antennae). However, for the sensilla placoidea, the most abundant olfactory organs (see also Results), we estimated the density [23]. We counted all of the sensilla placoidea within a square of 100  $\mu\text{m}$   $\times$  100  $\mu\text{m}$  situated approximately in the central zone of the dorsal

side of Fl<sup>9</sup> (a single measurement was performed on two antennae per species).

From pictures taken at higher magnifications (up to 3000 $\times$ ), we calculated the size of the sensilla as follows. We superposed transparent sheets onto printed images showing the sensilla placoidea (SP), sensilla basiconica (SB), pit organs (PO), and sensilla coelocapitula (SC) from above (so that the bases/pits were easily visible), and a manual drawing of the pits where the pore plates lay (SP), the peg base (SB) or the pit (PO) was produced. Using pictures where the SB, sensilla trichoidea (ST: type A, B and C/D) and grooved peg sensilla (GPS) were clearly visible laterally (i.e., the peg length was readily appreciable), we measured their length. Once all of the images were completed, we scanned and imported them into ImageJ software (National Institutes of Health, USA), in which the lengths and areas (for pits and peg bases) of the investigated structures were calculated. The sample of sensilla used for size calculations came from one or both antennae from each species, depending on their visibility/definition in the SEM images. Setae were not considered in the present work because although they are very abundant, these structures are non-innervated and have no known sensory role [32].

Finally, images obtained at higher magnifications (up to 10,000 $\times$ ) were used to further analyse the morphology of the various sensilla types or particular areas of the flagellomeres qualitatively.

As a measure of wasp body size, we recorded the head width [33] (in two females per species), which has been shown to be positively correlated with antennae length in Hymenoptera [15].

## 2.4. Statistical analysis

For each type of sensilla, we calculated the minimum, maximum, means and standard deviation of the mean (SD) for lengths and/or areas (and density for SP). The small numbers of analysed individuals and, consequently, the lack of information regarding intraspecific variability did not permit any analysis of variance to be performed for interspecific comparisons. Instead, to preliminarily explore the relationships among species based on sensilla size, we carried out a hierarchical cluster analysis, which finds relatively homogeneous clusters of cases based on measured variables. The variables used for each species were the mean measures for all types of sensilla, except those for which the sample size (number of measurements taken) was small ( $n < 10$  per species) (a total of eight variables were used). The cluster analysis was performed using Ward's method based on Euclidean distances (dissimilarity) between pairs of objects; this analysis also reported the dissimilarity value (truncation), which likely determines how many clusters best suit the data.

In the text and tables, mean values are shown  $\pm$  standard deviation. To better demonstrate possible differences in sensilla size and density between beetle-hunters and bee-hunters, we also presented the data grouped by prey type (i.e., averaging the values of the single species and then calculating the means and SD across these 5 points [species] per category).

### 3. Results

The head width ranged from 2 to 5 mm on average among the investigated species (Table 1).

The antennae of all of the species are typically filiform. They include a long scape at the proximal end, a rounded pedicel, and a flagellum (composed of 10 flagellomeres in female Apoidea) (Fig. 1). The Fl<sup>1</sup> is longer than all of the other flagellomeres in all species.

In all species except for two (*C. rubida* and *C. bupresticida*) (Fig. 2A), there is a small to conspicuous smooth area almost free of, or with very few, sensilla or setae on the tip of Fl<sup>10</sup> (Fig. 2B–F). In *C. quinquefasciata*, this area is small and shield-shaped and is located on the tip (Fig. 2B). The same shape and position were recorded for *C. tuberculata*, but the size was much larger (almost covering the entire tip). In *C. sabulosa*, this area is also on the tip and is small and roughly circular (Fig. 2C). In *C. arenaria*, it is medium-sized and roughly oval and is also located on the tip (Fig. 2D). In *C. rybyensis*, the shape and position of the area resemble those in *C. arenaria*, but the oval is narrower and smaller. *P. pulchellus* and *P. triangulum* present a large and kidney-shaped area on the ventral side (larger in *P. pulchellus*) (Fig. 2E). Finally, the bare area in *T. denticollis* did not differ in its shape and position from that of *Philanthus* spp., although is slightly wider, making it the largest sensilla-free area recorded among the investigated species (Fig. 2F).

We recognised nine different types of sensilla, which are described in detail below.

#### 3.1. Pit organs (PO)

These sensilla appear on the ventral and sometimes the lateroventral side of the 2nd to 10th flagellomeres (often forming fields) as cuticular openings, and they can include two types: the larger sensilla coeloconica (Scoel) and the smaller sensilla ampullecea (Samp) (Fig. 3A–B). In fact, despite Dietz and Humphreys [34] claiming the small pits

to be Samp and the wider ones to be Scoel in *A. mellifera*, the two types cannot be clearly differentiated based on external morphology using SEM; thus, we conservatively decided to suggest the presence of both only when differences in the area of the pit were clearly visible when the two types of sensilla were found close together (Fig. 3B). Among the investigated species, this difference was clear only in *C. rybyensis*, *C. sabulosa* and *C. bupresticida* (Fig. 3A and B). Thus, we preferred to group Samp and Scoel together as pit organs in the analysis of their distribution and size. The density of PO increases towards the tip of the antenna. The area of the PO pit ranged from 0.9 to 2.2  $\mu\text{m}^2$  across species (Table 2), and it averaged  $1.5 \pm 0.6 \mu\text{m}^2$  for beetle-hunters and  $1.4 \pm 0.4 \mu\text{m}^2$  for bee-hunters.

#### 3.2. Sensilla placoidea (SP)

These olfactory discs or pore plates are oval/elliptical-shaped, and the pits where they lay have an area of  $29.6 \pm 3.5 \mu\text{m}^2$  (average among species) (Table 2), with the longitudinal axis being parallel to the longitudinal axis of the antenna (Fig. 3C). The distal end of the plate rises higher above the cuticle than the proximal end in all species (Fig. 3C–D), although in *C. tuberculata* and *T. denticollis*, this rise appears to be only slightly above the cuticle. A joint-like membrane connects the surrounding cuticle of the plate (Fig. 3D–E). The plate bears many pores. Pore plates are present in all flagellomeres but are rarer on Fl<sup>1</sup> and much more abundant from Fl<sup>2</sup> to Fl<sup>10</sup>, with abundance increasing from proximate to distal flagellomeres. They are found, in particular, on the dorsal-lateral sides of flagellomeres (Fig. 4A), but they are also abundant ventrally. The area of the SP was  $27.7 \pm 3.0 \mu\text{m}^2$  in beetle-hunters and  $31.6 \pm 2.3 \mu\text{m}^2$  in bee-hunters.

The SP density at Fl<sup>9</sup> ranged from 24 to 49 per species in the  $100 \times 100 \mu\text{m}^2$  observation squares (Table 2), and it appeared to be somewhat higher in beetle-hunters ( $41.7 \pm 8.3$  vs.  $27.5 \pm 2.5$ ).

#### 3.3. Sensilla basiconica (SB)

The sensilla basiconica are the largest sensilla, with a mean length of  $8.8 \pm 1.8 \mu\text{m}$  and a mean area of  $65.5 \pm 25.2 \mu\text{m}^2$  in the investigated species (Table 2). The surface of the large sensilla peg is sculptured longitudinally, with many thin furrows, and ends in a perforated tip (Fig. 3F–I). The SB increase in number from the proximal to the distal end of the antenna and are present on the dorsal (and sometimes also the lateral) side (Fig. 4A–B) of the Fl<sup>2</sup>–Fl<sup>10</sup> flagellomeres in all *Cerceris* spp., except *C. sabulosa* and *C. rybyensis*, and from Fl<sup>3</sup> to Fl<sup>10</sup> in these two species and in *Philanthus* spp. and *T. denticollis*. Beetle-hunters seem to exhibit a larger SB area ( $81.4 \pm 11.0 \mu\text{m}^2$ ) than bee-hunters ( $49.6 \pm 22.7 \mu\text{m}^2$ ). Additionally, the length of the SB was  $9.6 \pm 0.9 \mu\text{m}$  for beetle-hunters and  $7.9 \pm 1.9 \mu\text{m}$  for bee-hunters.

When both the SB base area and SB length were considered together, some differences in the general shapes of the SB emerged. In particular, *C. tuberculata* appears to possess a fairly short, but large-based sensilla (length/area ratio: 0.46) (Fig. 3I), while *P. pulchellus* seems

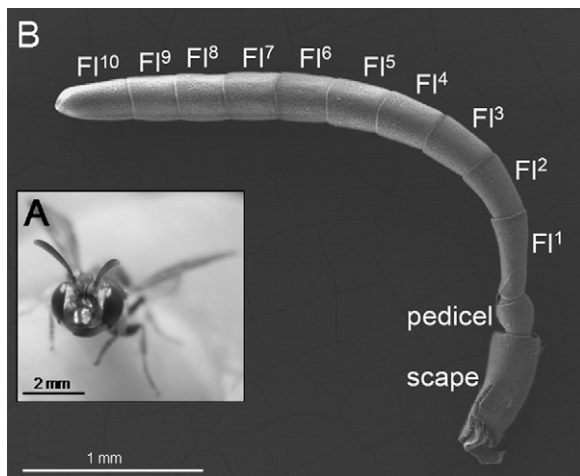


Fig. 1. A. A female philanthine wasp (*C. rubida*). B. Typical structure of the antenna in female Philanthinae (pictured: *C. arenaria*), composed of a scape, a pedicel and ten flagellomeres (Fl).



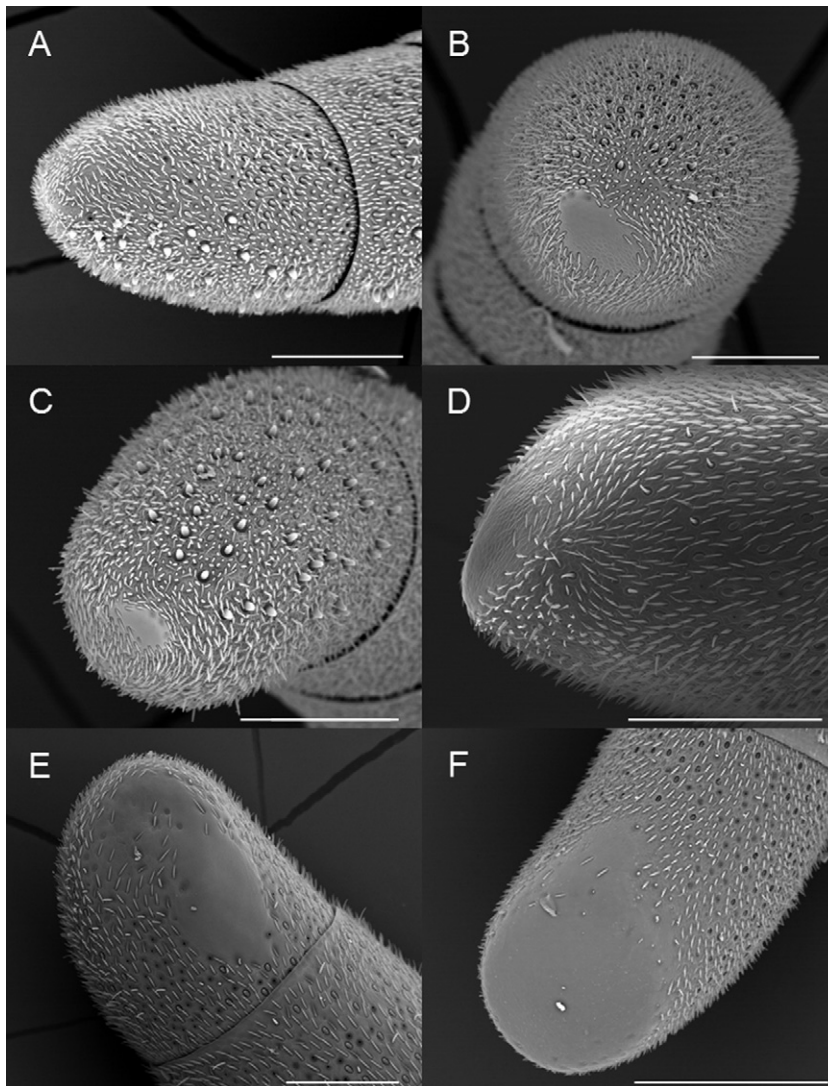


Fig. 2. Variability in the shape, position and size of the sensilla-free area on the apical portion of Fl<sup>10</sup>. A. *C. rubida* (absent). B. *C. quinquefasciata* (small and shield-shaped, tip). C. *C. sabulosa* (small and roughly circular, tip). D. *C. arenaria* (medium-sized and roughly oval, tip). E. *E. P. pulchellus* (large and kidney-shaped, ventral side). F. *T. denticollis* (large and kidney-shaped, ventral side). Bar indicates 100  $\mu\text{m}$  in all pictures except (F) in which it is 200  $\mu\text{m}$ .

to possess a more elongated sensilla (length/area ratio: 0.54–0.57). The values for the other species appear to fall within these extremes (Fig. 3F–H).

### 3.4. *Sensilla trichoidea A (ST-A)*

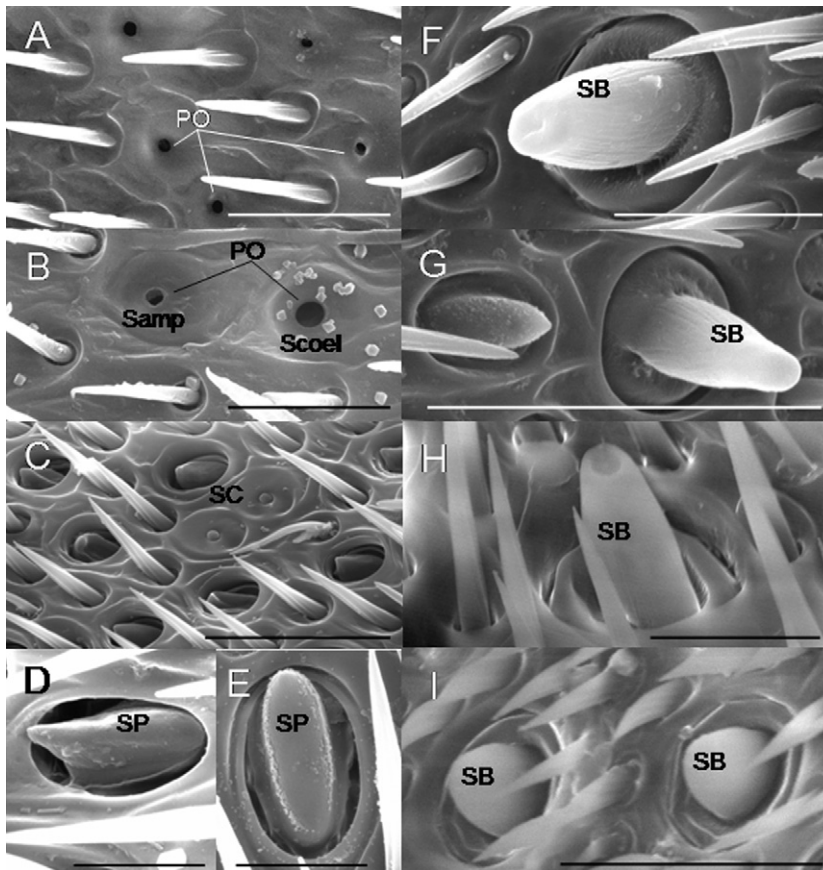
This type of trichoid sensilla exhibits a smooth peg and a tuberculoid-like base connected to the surrounding cuticle through a joint-like membrane (Fig. 5A, C, D). ST-A were present on the dorsal or, more rarely, lateral side of the antennae from the 2nd to 10th flagellomere in *Philanthus* spp. and from the 1st (very rare) to 10th flagellomere in the other species. The highest density of ST-A was found at the apex of the antenna. The length of the ST-A was  $12.8 \pm 4.8 \mu\text{m}$  on average among the investigated species (Table 2); ST-A length was  $15 \pm 0.4 \mu\text{m}$  in beetle-hunters and  $10.4 \pm 5.6 \mu\text{m}$  in bee-hunters.

### 3.5. *Sensilla trichoidea B (ST-B)*

These trichoid sensilla, which are slightly shorter than ST-A ( $11.2 \pm 1.6 \mu\text{m}$  average among all species, Table 2), are sharp, thin and curved against the surface and exhibit longitudinal furrows (Fig. 5A). ST-B are found from Fl<sup>2</sup> to Fl<sup>10</sup> on the dorsal side and from Fl<sup>1</sup> to Fl<sup>10</sup> on the ventral side, with their number appearing to increase distally. The lengths of the ST-B was  $10.8 \pm 2.1 \mu\text{m}$  for beetle-hunters and  $11.6 \pm 1.4 \mu\text{m}$  for bee-hunters.

### 3.6. *Sensilla trichoidea C/D (ST-C/D)*

This group includes various sub-types of trichoid sensilla [23]. ST-C/D are characterised by longitudinal furrows and a wide socket (Fig. 5B) and are found on all flagellomeres both dorsally and ventrally. The length of ST-



**Fig. 3.** Sensilla types on the flagella of philanthine wasps. A. *P. triangulum* (bar: 20  $\mu\text{m}$ ). B. *C. sabulosa* (bar: 10  $\mu\text{m}$ ). C. *C. quinquefasciata* (bar: 20  $\mu\text{m}$ ). D. *C. arenaria* (lateral view) (bar: 5  $\mu\text{m}$ ). E. *C. rubida* (dorsal view) (bar: 5  $\mu\text{m}$ ). F. *C. rubida* (bar: 10  $\mu\text{m}$ ). G. *P. triangulum* (bar: 20  $\mu\text{m}$ ). H. *C. bupresticida* (bar: 10  $\mu\text{m}$ ). I. *C. tuberculata* (bar: 20  $\mu\text{m}$ ). PO: pit organs; Samp: sensilla ampulleacea; Scoel: sensilla coeloconica; SC: sensilla coelocapitula; SP: sensilla placoidea; SB: sensilla basiconica.

C/D was variable, being  $10.4 \pm 3.2 \mu\text{m}$  on average among all species (Table 2), and it appears to be similar in beetle-hunters ( $11.4 \pm 3.5 \mu\text{m}$ ) and bee-hunters ( $9.3 \pm 1.9 \mu\text{m}$ ).

### 3.7. Grooved peg sensilla (GPS)

We found these peg-like organs only in *P. triangulum*, *P. pulchellus* and in *C. sabulosa*, whereas they were apparently lacking in all the other *Cerceris* spp. and in *Trachypus*. The base of the peg is sunk below the surrounding cuticular surface, and it contains deep longitudinal furrows (Fig. 5D). It was rarely observed on the dorsal side from Fl<sup>6</sup> to Fl<sup>9</sup>. The length of the GPS was approximately 11  $\mu\text{m}$  on average in *Philanthus* spp. and approximately 6  $\mu\text{m}$  in *C. sabulosa* (Table 2).

### 3.8. Sensilla coelocapitula (SC)

This type of sensilla is characterised by a bulbous or button-like knob with an irregular surface emerging from an opening in the centre of a quite irregular oval concavity. SC are found ventrally and, notably, even in nearly bare areas where the other sensilla are types generally lacking (Fig. 3C and Fig. 6). For example, in some cases, they were

found in close proximity to the sensilla-free area on the antennal apex (Fig. 6). SC were regularly observed to form compact clusters of 2–3 sensilla (Fig. 6C), and SC are often located in the vicinity of pit organs. Although SC are very rare compared to the other sensilla types, they were found scattered across all flagellomeres. The SC bulb area was  $1.3 \pm 0.3 \mu\text{m}^2$  on average among all investigated species (Table 2) and 0.9–1.2  $\mu\text{m}^2$  for beetle-hunters and 1.3–1.9 for bee-hunters.

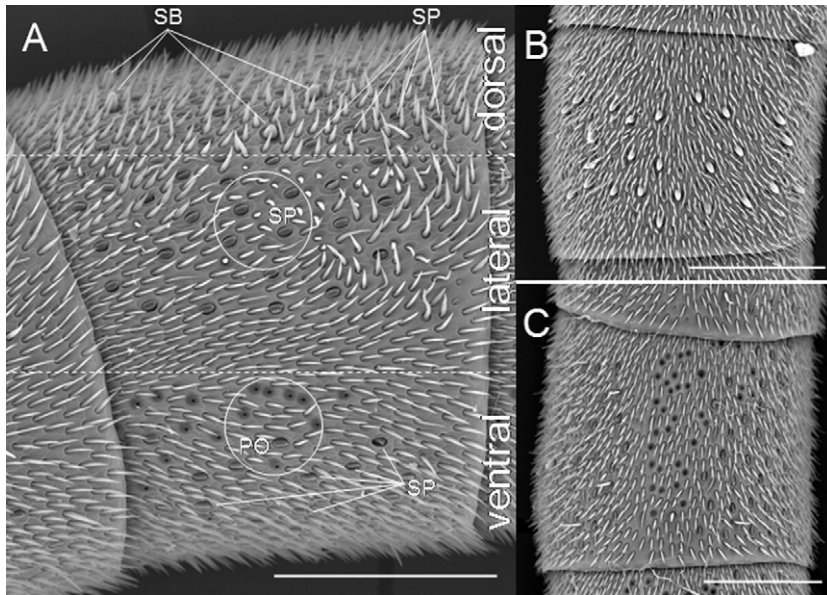
### 3.9. Cluster analysis

The dendrogram obtained through the cluster analysis indicated three main clusters (Fig. 7) (dissimilarity value of truncation = 0.93). Cluster 1 (the first cluster shown from top to bottom in the dendrogram) included two beetle-hunters. Cluster 1 was closer to cluster 2 (distance between group centroid = 0.94), which included two bee-hunters and three beetle-hunters, compared to cluster 3, which was the most dissimilar group from the others (distances to group 1 centroid = 1 and to group 2 = 1.43), contained three bee-hunters. In the “mixed group” (cluster 2), the two bee-hunting species were not highly similar. These similarity patterns show that the sensilla morphology and

**Table 2**  
Mean area and/or length ( $\pm$ SD) of the antennal sensilla in the studied species, and SP density on the FI<sup>9</sup>.

Variable	<i>C. arenaria</i>	<i>C. bupresticida</i>	<i>C. quinquefasciata</i>	<i>C. rubida</i>	<i>C. rybyensis</i>	<i>C. sabulosa</i>	<i>C. tuberculata</i>	<i>P. pulchellus</i>	<i>P. triangulum</i>	<i>T. denticollis</i>
PO (area) ( $\mu\text{m}^2$ )	0.9 $\pm$ 0.3 (10)	2.2 $\pm$ 1.1 (12)	0.9 $\pm$ 0.4 (10)	1.5 $\pm$ 0.3 (13)	1.1 $\pm$ 0.4 (11)	1.3 $\pm$ 0.7 (11)	2.2 $\pm$ 0.3 (14)	2.1 $\pm$ 0.4 (10)	1.1 $\pm$ 0.2 (11)	1.4 $\pm$ 0.2 (9)
SP (area) ( $\mu\text{m}^2$ )	28.8 $\pm$ 1.5 (18)	32.6 $\pm$ 3.2 (37)	28.1 $\pm$ 3.9 (18)	23.8 $\pm$ 0.6 (18)	29.7 $\pm$ 1.5 (18)	27.9 $\pm$ 1.2 (24)	25.2 $\pm$ 2.0 (40)	33.2 $\pm$ 3.9 (24)	33.5 $\pm$ 3.3 (18)	33.7 $\pm$ 3.8 (18)
SP density at FI <sup>9</sup>	46 $\pm$ 7 (2)	34 $\pm$ 2 (2)	29.5 $\pm$ 0.5 (2)	49.5 $\pm$ 2.5 (2)	31.5 $\pm$ 4.5 (2)	26.5 $\pm$ 4.5 (2)	49.5 $\pm$ 3.5 (2)	28 $\pm$ 1 (2)	24 $\pm$ 4 (2)	27.5 $\pm$ 7.5 (2)
SB (area) ( $\mu\text{m}^2$ )	79.3 $\pm$ 6.4 (10)	86.4 $\pm$ 4.2 (8)	78.0 $\pm$ 7.4 (16)	64.8 $\pm$ 4.6 (11)	18.8 $\pm$ 2.6 (15)	73.3 $\pm$ 10.8 (11)	98.6 $\pm$ 12.3 (8)	43.6 $\pm$ 3.4 (20)	78.1 $\pm$ 10.5 (9)	34.4 $\pm$ 3.0 (7)
SB (length) ( $\mu\text{m}$ )	8.8 $\pm$ 1.2 (9)	11.0 $\pm$ 0.7 (8)	9.9 $\pm$ 0.4 (8)	10.1 $\pm$ 0.9 (8)	4.9 $\pm$ 0.9 (11)	8.3 $\pm$ 0.8 (8)	8.5 $\pm$ 0.8 (8)	8.8 $\pm$ 0.9 (8)	10.7 $\pm$ 0.6 (8)	6.9 $\pm$ 0.8 (8)
ST-A (length) ( $\mu\text{m}$ )	11.3 $\pm$ 0.8 (10)	24.9 $\pm$ 2.5 (10)	16.4 $\pm$ 1.6 (10)	8.3 $\pm$ 1.4 (10)	9.8 $\pm$ 2.1 (10)	10.8 $\pm$ 2.0 (10)	14.3 $\pm$ 4.4 (10)	11.9 $\pm$ 1.1 (21)	9.6 $\pm$ 0.4 (10)	10.1 $\pm$ 1.1 (12)
ST-B (length) ( $\mu\text{m}$ )	11.1 $\pm$ 1.9 (11)	7.3 $\pm$ 0.6 (12)	13.9 $\pm$ 1.3 (10)	10.2 $\pm$ 1.5 (11)	13.7 $\pm$ 4.6 (10)	9.5 $\pm$ 1.0 (10)	11.6 $\pm$ 0.9 (10)	10.8 $\pm$ 0.7 (10)	11.6 $\pm$ 0.8 (12)	12.5 $\pm$ 1.8 (9)
ST-C/D (length) ( $\mu\text{m}$ )	8.8 $\pm$ 2.0 (10)	16.8 $\pm$ 1.8 (10)	8.2 $\pm$ 1.8 (10)	9.0 $\pm$ 2.0 (10)	10.0 $\pm$ 1.7 (10)	10.7 $\pm$ 2.4 (10)	14.3 $\pm$ 2.1 (10)	7.5 $\pm$ 0.5 (10)	6.5 $\pm$ 0.8 (10)	11.7 $\pm$ 1.6 (10)
GPS (length) ( $\mu\text{m}$ )	–	–	–	–	–	6.2 $\pm$ 1.0 (2)	–	11.2 $\pm$ 1.5 (3)	10.9 $\pm$ 1.9 (3)	–
SC (area) ( $\mu\text{m}^2$ )	1.0 $\pm$ 0.0 (2)	1.0 $\pm$ 0.0 (2)	0.9 $\pm$ 0.0 (2)	1.2 $\pm$ 0.1 (3)	1.3 $\pm$ 0.0 (2)	1.6 $\pm$ 0.4 (3)	1.1 $\pm$ 0.0 (2)	1.5 $\pm$ 0.1 (3)	1.8 $\pm$ 0.1 (3)	1.9 $\pm$ 0.1 (2)

In brackets there are the sample sizes (number of sensilla measured) (number of antennae for SP density). PO: pit organ; SP: sensilla placoidea; SB: sensilla basiconica; ST: sensilla trichoidea; GPS: grooved peg sensilla; SC: sensilla coelocapitula.



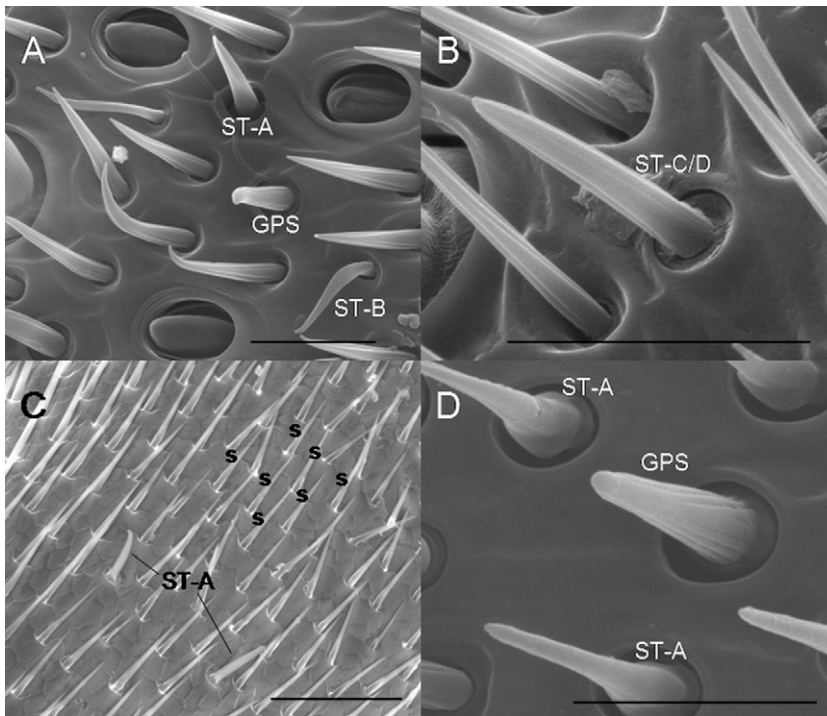
**Fig. 4.** Typical distribution of SP, SB and PO on a flagellomer. A. Inner lateral side of FI<sup>7</sup> in *C. quinquefasciata*. Note that SB are abundant on the dorsal side, PO on the ventral side, and SP on dorsal, ventral and lateral sides. B. Dorsal side of FI<sup>9</sup> in *C. rubida* with abundant SB. C. Ventral side of FI<sup>8</sup> of *C. sabulosa* with a field of PO. Bar is 100 μm.

density did not mirror the known taxonomic relationships among the studied species.

**4. Discussion**

This study contributes to current knowledge regarding the sensilla equipment in female apoid wasps, increasing

the number of species studied in detail from five to 13. Despite the fact that this study provides a very general view and the sensilla types were found to be morphologically similar across these 13 species and similar to other species of Apoidea (i.e., bees) [8,17,19,34,35], our results demonstrated some degree of variability in the size and, at least in one case (SB), the shape of different types of



**Fig. 5.** Sensilla types on the flagella of philanthine wasps. A. *C. sabulosa* (bar: 10 μm). B. *C. sabulosa* (bar: 10 μm). C. *P. coronatus* (bar: 50 μm). D. *P. pulchellus* (bar: 20 μm). ST-A: sensilla trichoidea A; ST-B: sensilla trichoidea B; ST-C/D: sensilla trichoidea C/D; GPS: grooved peg sensilla; s: setae.



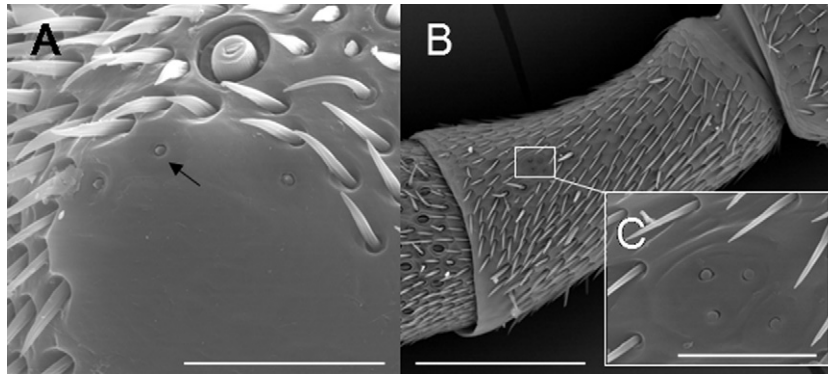


Fig. 6. Presence of SC (one is indicated by an arrow) in areas where all of the other types of sensilla were very rare or absent. A. Bare area at the apex of FI<sup>10</sup> in *C. quinquefasciata* (bar: 20  $\mu\text{m}$ ). B. Ventral side of FI<sup>1</sup> bearing SC in *C. sabulosa* (bar: 100  $\mu\text{m}$ ). C. Details of three fused and one separated SC (bar: 20  $\mu\text{m}$ ).

sensilla, even between closely related species (within genera of Philanthinae). Below, we discuss our findings from a comparative perspective using available information on females of other studied species of Apoidea (and in particular cases, also other types of insects) and previous observations on the two species of wasps investigated for the second time in this study (*C. rybyensis* and *P. triangulum*). We conclude with a discussion regarding the possible effect of wasp ecology on their sensillar equipment.

#### 4.1. General patterns of sensilla distribution

The general distribution patterns of the different sensilla types in the studied species do not differ markedly from those observed in other apoid wasps and in bees [8,17,23,24,34,35]. Most sensilla are mainly located dorsally and sometimes laterally and show increasing abundance distally. Two exceptions to this pattern are the PO, which appear to mainly aggregate in fields on the ventral side of flagellomeres (though in *Bembix rostrata* (L.), they are abundant on the lateral antennal surface [23]), and the SC, which were also found ventrally.

As in many bees and other studied apoid wasps [23,35], there is a bare, often ventral, zone on FI<sup>10</sup> in philanthine

wasps. In this group, the shape and size of this sensilla-free area is highly variable: it is generally small and oval-circular in most *Cerceris* species (and even lacking in two of these species), while it is quite large and kidney-shaped in *Philanthus* + *Trachypus*. Thus, it appears that the tribe Philanthini can be separated from the tribe Cercerini based on the presence, size and shape of this area on the antennal apex. However, in three apoid wasps of the subfamily Bembecinae (two species of *Argogorytes* and *B. rostrata*), there was also no sensilla-free area observed [23], while bees in the families Colletidae, Andrenidae and Halictidae presented a sensilla-free area with an oval shape and a medium-to-large size [8,17,35]. Thus, discrimination of tribes based on the characteristics of this area could work within Philanthinae, but not within Crabronidae or Apoidea. The function of this sensilla-free area is not known.

#### 4.2. Morphology of the sensilla

In general, all of the sensilla types observed in the 10 philanthine wasp species closely resemble those observed in other apoid wasps [23,24] and have homologous counterparts in bees [8,17,19,34,35].

The pit organs (PO) in the investigated species (and in apoid wasps in general) have a remarkably similar external morphology to those observed in bees [8,17,35], with smaller and larger pores, which according to the study by Dietz and Humphreys [34] on *Apis mellifera* L., should correspond to sensilla ampullacea (Samp) and sensilla coeloconica (Scoel), respectively. However, as also observed in some bee and wasp species, distinction of these structures is difficult based solely on SEM analysis, and histological preparation of sensilla sections is necessary to ascertain which pores belong to each of the two types of pit organs. In *Sphecodes* bees and the halictid bee *Augochlora pura* (Say) [15], the distinction was more accurate because in Samp, no inner pit structure was detected by SEM, while in Scoel a low furrowed peg is often observed [35]. In *Sphecodes*, the pit areas were approximately 0.4  $\mu\text{m}^2$  for Samp and 3.6  $\mu\text{m}^2$  for Scoel [35], while in Bembecinae, the Scoel appeared to be much larger (7–28  $\mu\text{m}^2$ , depending on the species) [23] (areas calculated from data on diameters). Among Bembecinae wasps, Ågren [23]

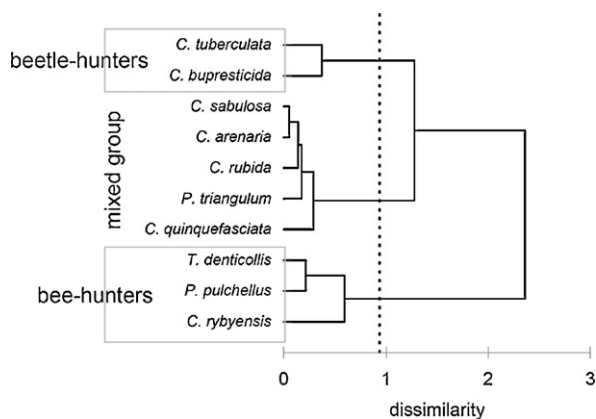


Fig. 7. Graphic representation of species dissimilarity based on agglomerative hierarchical clustering using measures of six types of sensilla (SP, SB, PO, ST-A, ST-B, ST-C/D).

assessed the presence or absence of Samp and Scoel based on the visibility of this peg. These structures were not as clear in the species studied here. Nevertheless, we were at least able to demonstrate that clear size dimorphism was observed in *C. rybyensis*, *C. sabulosa* and *C. rubida* through direct observations performed on closely situated pits. We cannot exclude the possibility that some species lack one of the two types of PO, as this situation has been reported for other apoïd wasps [23]. Samp act as CO<sub>2</sub>, temperature and/or humidity receptors in *A. mellifera* [36] and CO<sub>2</sub>-receptors in ants [37]. Scoel are assumed to have similar functions in *A. mellifera* [38].

The sensilla placoïda (SP) in the studied species did not differ compared to other bees and wasps described previously in their general appearance (roughly circular or oval plates connected within the pit through a membrane). They are approximately 20–40 μm<sup>2</sup> in size, which does not differ considerably from the size observed in bembecine wasps (6–8 μm along the long axis, which would roughly correspond to an area of approximately 20–40 μm<sup>2</sup> for an oval shape with a short axis = 2/3 × long axis) [23] but is markedly smaller than what is observed in some bees (Colletidae) (11–16 μm along the long axis, i.e., approximately 70–150 μm<sup>2</sup> for an oval shape with a short axis = 2/3 × long axis) [17].

The SP in apoïd wasps also show other visible morphological differences compared to bees. In fact, while in bees, the SP exhibit plates that are either level with, or slightly above, the surrounding cuticle [8,17,19,31,35], in Bembecinae [23], Philanthinae [23,24] (this study) and other apoïd wasps [26], the SP are characterised by pore plates elevated from the surrounding cuticle (though weakly in *C. tuberculata* and *T. denticollis*). As SP are multiporous single-walled sensilla [26] that are known to respond to volatiles [11] (and references therein), these structures were assessed as olfactory organs in Apoïdea.

The sensilla basiconica (SB) are large sensilla with a typical stout, cylindrical, bulbous morphology in apoïd wasps [23–25] (this study). In contrast, bees exhibit typically slender, long SB with small sockets [8,17,35]. In addition, in this study, we demonstrated that the peg shape also shows variability within Philanthinae: some species present more elongated SB, while other SB exhibit a large base and a shorter peg. The sizes of the SB, in general, do not appear to be particularly dissimilar from the sizes in apoïd wasps of other subfamilies (approximately 8–10 μm in length [23], compared to approximately 5–11 μm in the species studied here). The large SB were suggested to be olfactory or gustatory organs [23,25,39].

The sensilla trichoïda A (ST-A) of philanthine wasps exhibit a very similar appearance to those of other apoïd wasps [23] and bees [8,17,35]. These sensilla have been described as single-walled hairs with wall pores, suggesting that they most likely have an olfactory function [32].

The sensilla trichoïda B (ST-B) are just slightly smaller than the ST-A in Philanthinae (approximately 11 μm vs. 13 μm in the studied species); their length is similar to that observed in colletid bees (approximately 10–11 μm) [17]. Following Lacher [36] and as discussed by Agren [17] in *A. mellifera*, ST-B may act as mechanoreceptors.

The sensilla trichoïda C/D (ST-C/D) of the philanthine wasps did not differ from those found in bembecine wasps [23] and presumably include a mixture of trichoid types that are not easily distinguishable, in particular, the ST-C and ST-D of bees [8,17,34]. ST-C/D include the longest trichoid sensilla in most Apoïdea, but in general, their length is extremely variable, such that on average, they do not appear to be very dissimilar from the ST-A and SB (*Argogorytes fergei* (Shuckard): 22 μm, *Andrena* spp.: 15 μm, *Sphecodes* spp.: 13–30 μm, *P. triangulum*: 8–16 μm) [8,23,24,35]. The same large size variability was found in the philanthine wasps studied here (maximum length/minimum length ratio ranging from 1.2 to 2.3 across the 10 species). A mechanoreceptor-gustatory function has been suggested for ST-C/D [31,32,36].

Grooved peg sensilla (GPS) were first found in Apoïdea in *P. triangulum* by Herzner et al. [24], and we now show that these structures are present in at least one other species of *Philanthus*, *P. pulchellus*, and in the genus *Cerceris* (*C. sabulosa*). They have not been found in any of the other studied species of apoïd wasps to date [23,24,26] (this study). However, caution should be taken in claiming that GPS are absent because this type of sensilla is rare, and we could have missed certain areas, particularly in twisted and bent antennae, where they may have been present. The length of the peg was found to be similar in our *P. triangulum* sample, in *P. pulchellus* and in the *P. triangulum* sample analysed by Herzner et al. [24] (approximately 9–10 μm), though it was somewhat shorter in *C. sabulosa*. The GPS of philanthine wasps resemble those found in other non-Hymenoptera insects [40,41], and because they are thought to be double-walled-pore sensilla, they are believed to have an olfactory function [42].

Similar to other types of sensilla, such as ST-A, ST-B and PO, the morphology of the sensilla coelocapitula (SC) observed in this study did not deviate from that observed in bees and in other apoïd wasps. The distribution of these structures is also similar across species, with SC often forming small and dispersed groups along the 10 flagellomeres, commonly on the most proximal annuli and also on the very tip of the antenna, i.e., where the other types of sensilla are normally almost absent [8,17,23,24,31,35]. Notably, SC appear to be lacking in females of several species of bees (*Hylaeus communis* Nylander) [17] and bembecinae wasps (*A. fergei*) [23], whereas they were always found in Philanthinae [24] (this study). SC are most likely mechanoreceptors [31].

#### 4.3. Conclusions and a hypothesis to be tested in the future

Overall, the variability in the morphology, size and distribution of sensilla types found in this study confirms the hypothesis of Ågren [23] that antennal sensilla features are not valuable as taxonomic tools in apoïd wasps. With the sole exception of the shape and size of the sensilla-free area on Fl<sup>10</sup>, which appears to separate Philanthini and Cercerini, none of the features of these structures appear to be valid for discrimination among species, genera or tribes. The cluster analysis based on sensilla sizes confirms this observation. A weak impact of taxonomic relationships in

determining sensillar morphology is also observed in some parasitoid hymenopterans, in which gustatory sensilla are similar among species, even in those belonging to different families, suggesting the existence of ecological factors shaping the forms of these structures [43]. Furthermore, the species grouping obtained with the cluster analysis appears to be independent of species size differences. For example, the three species in the tribe Philanthini have SP virtually of the same size, which are larger compared to those of the largest species in the sample (the beetle-hunter *C. tuberculata*); furthermore, *T. denticollis*, one of the largest species, had the second smallest SB.

Thus, it is reasonable to hypothesise that variability in sensillar equipment arose from different ecological demands in different apoid wasps. Here, we preliminary suggest one ecological factor, prey type, as potentially shaping this process because it is a main ecological/behavioural trait differing within species of the studied group. In fact, other factors, such as nesting habits and mating strategies, seem to be largely conservative in the subfamily [20].

Given the observed results, despite their mainly qualitative output, it appears that if prey type has shaped the form of the sensillar equipment, it would presumably act on olfactory sensilla because the differences between beetle-hunters and bee-hunters appeared to be best appreciable regarding both the size and distribution of SB and the density of SP. In particular, the overall, though tentative, suggestion is that beetle-hunters may achieve an increased olfactory sensitivity through larger SB (approximately 1.7-fold larger than bee-hunters), a greater abundance of SB (which are present on nine flagellomeres, compared to eight in bee-hunters), and a higher SP density (approximately 1.6-fold denser than bee-hunters).

However, why would beetle-hunters require enhanced olfaction compared to bee-hunters?

The general behaviour of bee and beetle prey may suggest one possible hypothesis. Wasp species that hunt for flying prey species on flowers, such as bees, certainly rely on olfaction, but mainly after an initial discrimination of the prey's size and shape using vision [44]. In contrast, wasp species hunting for beetle prey, which are often cryptic, exhibit poor mobility and live hidden in vegetation [45,46], would rely more strongly on olfaction while searching and localising prey, while visual recognition, if any, takes place once the prey is found. Thus, bee-hunters would essentially use visual and short-range chemical stimuli to find and recognise their prey, while in addition to these factors, beetle-hunters would also heavily rely on long-range chemical stimuli to find prey. Thus, successful foraging might require increased olfactory sensitivity in beetle-hunters, and larger pegs of SB could represent more efficient olfactory organs, similar to what has been observed in other insects. For example, the demand for extreme sensitivity in moth pheromone communication supported the evolution of long sensilla with a high efficiency in capturing odour molecules [47]. The distribution of SB also appears to cover more flagellomeres in beetle-hunters, but this result should be further confirmed because one bee-hunter (*C. rybyensis*) was shown to have SB on eight flagellomeres in the present study and on nine

flagellomeres previously [23], and thus, we cannot exclude important intraspecific variation.

A higher SP density was related to olfactory sensitivity in previous studies on bees. For example, sensory allometry is associated with foraging task preferences and resource exploitation in *A. mellifera*, with the number of SP being greater in pollen and water foragers, which are known to exhibit higher sensory sensitivity compared to nectar foragers [48]. Additionally, *A. mellifera* males have a greater number of SP than workers (7.36 times more), possibly conferring the high olfactory sensitivity necessary to find the queen during the mating flight [49].

In further agreement with this hypothesis, Galvani et al. [19] estimated that in bees of the tribe Emphorini (Apidae), 85–90% of sensilla in males correspond to olfactory receptors compared to 60–75% in females, whereas the number of contact receptors was greater in females. In sweat bees (Halictidae), parasitic females, which are potentially mainly dependent on longer-range olfactory cues, have fewer olfactory sensilla than non-parasitic females, which also use contact or short-range olfactory cues [15]. The antennae of females the parasitoid species *Tamarixia radiata* (Waterston) (Hymenoptera: Eulophidae) have a greater number of SP than male antennae, suggesting a possible function in the perception of host-related volatile cues [50].

The role of resource (host, prey or pollen) specialisation (in terms of niche width) in determining the density/number of SP is still debated in Hymenoptera. For example, the generalist braconid parasitoids *Cotesia glomerata* (L.) and *C. marginiventris* (Cresson) exhibit less SP than their specialist relatives *C. rubecula* (Marshall) and *Microplitis croceipes* (Cresson) [51,52]. In contrast, across bee families, oligolectic species do not have more SP than generalist species [15]. In the present study, a role of prey niche width also seems improbable, as the most specialised species (*P. triangulum*) shows one of the lowest SP densities, while more generalist species, such as *C. rubida* and *C. sabulosa*, showed widely ranging SP densities.

A relationship between foraging ecology and sensillar equipment, as preliminarily proposed here for digger wasps, is also supported by previous studies on non-aculeate Hymenoptera and non-hymenopteran insects. For example, females of the mymarid parasitoids *Anaphes victus* (Huber) and *A. listronoti* (Huber), despite possessing very similar antennal sensillar equipment, show differences in the number of one type of sensilla (sensilla chaetica type 4), which is lower in the solitary vs. the gregarious species [10]. In six species of Triatominae (Heteroptera), the number and distribution of four sensilla types was found to vary according more to habitat type (e.g., palm trees vs. human houses) than to how species are taxonomically related, supporting the idea that the patterns of antennal sensilla are sensitive indicators of adaptive process in these insects [53].

In conclusion, the sensillar equipment of philanthine wasps is complex, including many types of sensilla, and no evidence of a link between interspecific variability and taxonomic relationships among species is supported; instead, a candidate factor to be considered while searching for the causes of the observed variability may

be prey type. A larger species dataset, an increased number of individuals *per* species, and a statistical analysis of species differences incorporating a robust correction for phylogeny are needed to confirm this hypothesis.

### Disclosure of interest

The authors declare that they have no conflicts of interest concerning this article.

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