

Sting microsculpture in the digger wasp *Bembix rostrata* (Hymenoptera, Crabronidae)

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Abstract

The sting microsculpture of the digger wasp *Bembix rostrata* (Fabricius, 1781) (Hymenoptera, Crabronidae) is studied with the scanning electron microscope (SEM) for the first time. As in many other hymenopterans, the second valvifer of *B. rostrata* possesses two fields of styloconic sensilla (hair plates) of proprioceptive function. The presence of two paired fields of campaniform sensilla on the second valvula and second valvifer is first shown in an apoid wasp. The first and the second valvulae bear scattered sensilla-like structures on the external surface, more numerous apically. The first valvula has two subapical barbs externally and a pair of valvilli on its inner surface, whereas the outer surface of the second valvula is smooth. The third valvula is sclerotized externally, consisting of proximal and distal parts, and bearing four sensilla morphotypes of mechanoreceptive and probably chemoreceptive functions. The inner surface of the valvulae and the membranous cuticle that is touching the sting have microstructures of different shapes directed distally. Functional aspects of characters studied are discussed.

Keywords

Morphology, sting apparatus, ovipositor

Introduction

The Hymenoptera are the sole group of endopterygote insects with a well-developed ovipositor, a plesiomorphic retention that has been considered one of the key factors in their diversification (Gauld and Bolton 1988). Besides oviposition, it can perform a number of other functions including sawing a substrate; location, discrimination

and marking of the host; envenomation; stinging etc. (Quicke et al. 1999). In most Aculeata, the ovipositor has lost the primary function of an egg transporting device and is used as a sting to paralyse and rarely to transport the prey or as defense weapon (Rasnitsyn 1980, Radović 1985, Radović and Sušić 1997).

There are numerous classic works on morphology and functions of the hymenopterous ovipositor (e.g., Snodgrass 1956, Oeser 1961, Scudder 1961, Smith 1970, Quicke et al. 1999, Vilhelmsen 2000, Vilhelmsen et al. 2001, Packer 2003). Several publications dealt especially with microsculptural characters which have been used for phylogenetic reconstructions in some groups, mainly parasitic wasps (e.g., Quicke et al. 1992, 1999, Le Ralec et al. 1996, Nénon et al. 1995, Rahman et al. 1998). Both the comparative and functional aspects of the sting apparatus have been investigated extensively throughout some groups of the Apoidea (for review see Radović 1985, Packer 2003, Cardinal and Packer 2007), however only a few species of apoid wasp were explored in this respect (see review in Gadallah 2001). As a sister group of bees, the Crabronidae have gained insufficient attention during the last 10 years (Gadallah 2001, Packer 2003, Gadallah and Assery 2004). Generalizations concerning the sting structure in apoid wasps are still needed, though some attempts to compare sting morphology across several groups of apoid wasps have been made (Gadallah 2001, Gadallah and Assery 2004). Surprisingly, microstructure characters including sensory structures have not been discussed in the literature, except for a few general comments on the presence and distribution of some sensilla (Radović and Sušić 1997, Gadallah and Assery 2004). To the best of my knowledge, more detailed information on the sting microsculpture in apoid wasps is not available. The purpose of this study is to describe sting microsculpture in the digger wasp *Bembix rostrata* (Fabricius, 1781) (Hymenoptera, Crabronidae) using scanning electron microscopy in order to consider its possible functional use.

Materials and methods

Six females of *B. rostrata* were collected in Central Ukraine (Kyiv Province, Vyshgorodsky District, surroundings of village Khotyanivka, July 2008; 50°34'55"N 30°33'51"E). The gaster was removed from the wasp body using forceps, cut open slightly and macerated in 10% KOH. The sting apparatus was subsequently excised from the genital chamber, washed in water and examined in glycerine under a stereo microscope. For SEM study, the cuticular parts were washed in distilled water, dehydrated in a graded ethanol series and acetone, critical point dried (OM CPD 7501), coated with gold-palladium (OM-SC7640) and examined with a Zeiss EVO-50 SEM (Museum of Zoology, Natural History Senckenberg Collections Dresden, Germany).

The terms used are preferably from Vilhelmsen et al. 2001, but also from Quicke et al. 1999, Gnatzy and Volknandt 2000 and Packer 2003. The following abbreviations were used: ap, apical process of inner membranous wall of 3rd valvula; au, aulax;

blb, bulb of the 2nd valvula; dv, ductus venatus (= venom duct); rh, rhachis; rp, rostral process of 2nd valvifer; r1, ramus of the 1st valvula; r2, ramus of the 2nd valvula; T9, tergum 9; vlv, valvilli; 1vf, 1st valvifer; 1vv, 1st valvula; 2vf, 2nd valvifer; 2vv, 2nd valvula; 3vv, 3rd valvula.

Results

General organization of the sting apparatus (Figs 1, 2A, B). The sting apparatus of *B. rostrata* lies within the genital chamber formed by the partial infolding of the 8th and 9th abdominal segments (7th and 8th metasomal segments) into the 7th abdomi-

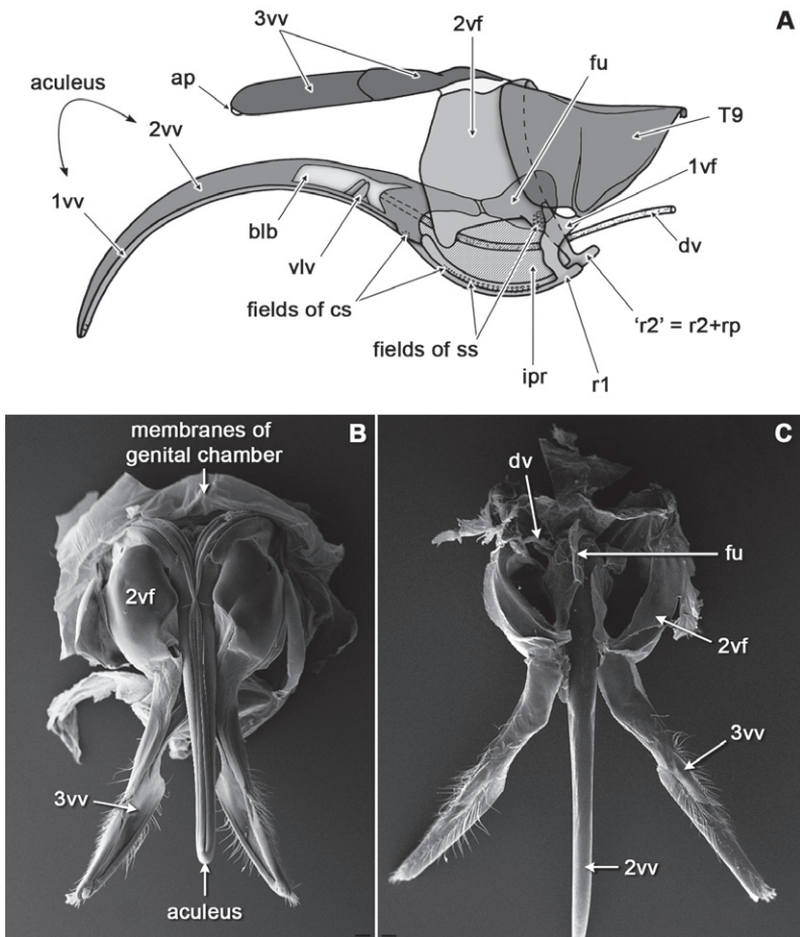


Figure 1. General organization of the sting apparatus in the digger wasp *Bembix rostrata*: **A** diagram showing relative position of main parts in a lateral view (aculeus is extended ventrally, furcula is turned anteriorly; bulb is shown in longitudinal section to show valvilli, the membranous incisura postarticularis is hatched); B-C, SEM micrograph of the sting in ventral **B** and dorsal **C** views. Scale bars: 100 µm.

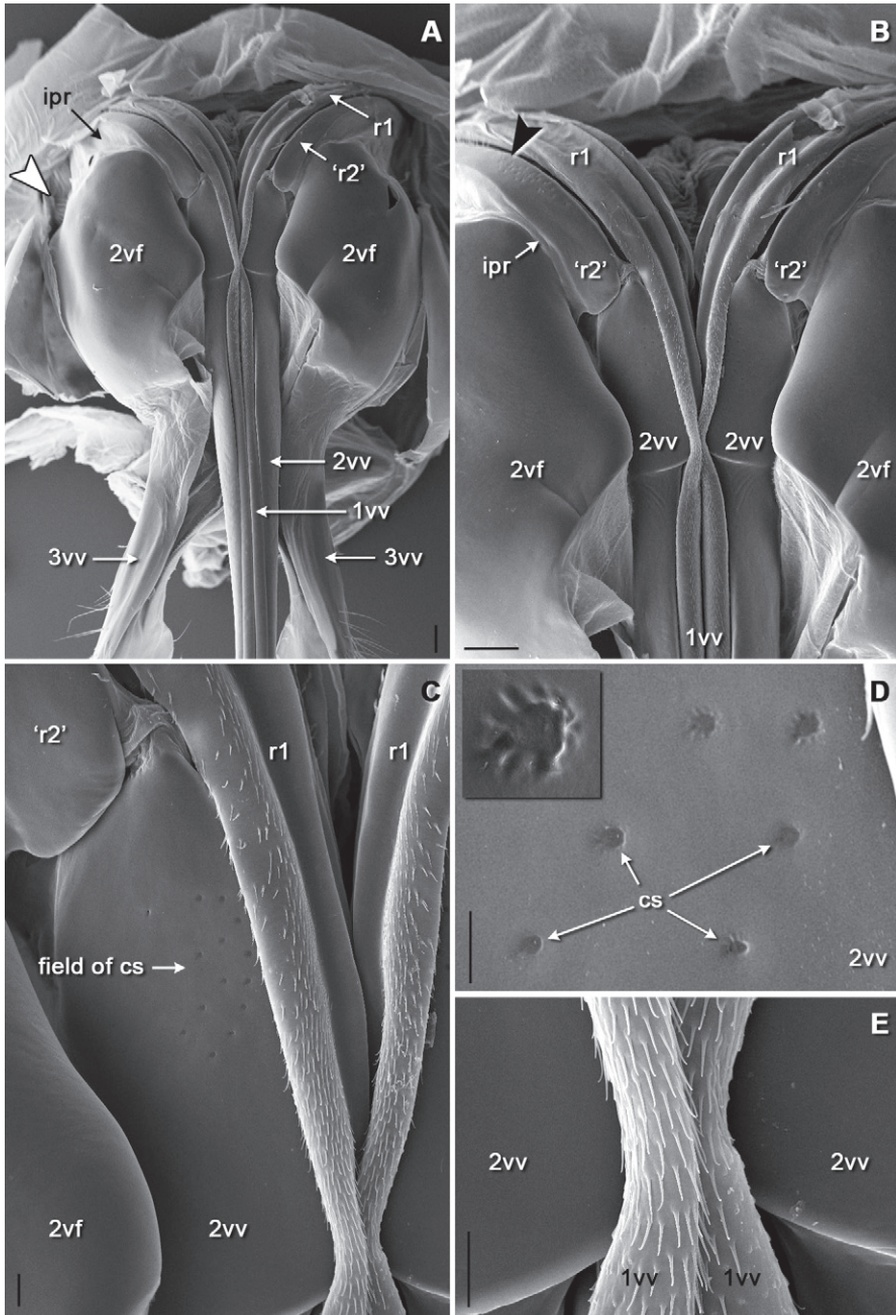


Figure 2. SEM micrographs of the sting basis, ventral view: **A** general aspects **B** basal region of the rami of both valves and of the second valvifera **C** field of campaniform sensilla on basis of the second valve **D** detail of the field of campaniform sensilla on basis of the second valve (one campaniform sensillum enlarged in inset) **E** distally directed microspination on membranous regions of the first valves. Arrowhead in A shows position of the field of styloconic sensilla on the second valvifer nearby its articulation with the first valvifer (enlarged in Fig. 3B). Scale bars: 100 μ m in A and B; 20 μ m in C and E; 10 μ m in D.

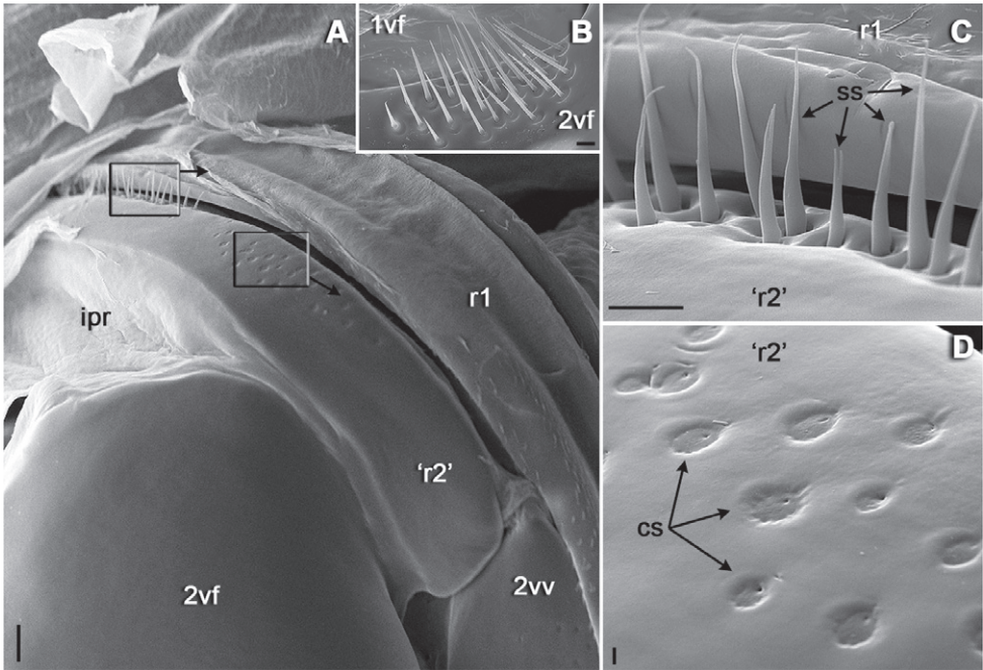


Figure 3. SEM micrographs of the sensilla on the second valvifer: **A** position of two fields of sensilla on the articular process of second valvifer, ventral view **B** field of styloconic sensilla nearby the articulation with first valvifer, lateral view **C** detail of the row of styloconic sensilla on the articular process of the second valvifer **D** detail of the field of campaniform sensilla on the articular process of the second valvifer distally to the row of styloconic sensilla. Insets in A are enlarged in C and D. Scale bars: 30 μm in A; 10 μm in B and C; 2 μm in D.

nal segment, thus only the apical part of the sting is visible outside at rest. The sting shaft itself (= aculeus) is massive, curved ventrally and comprises paired 1st valvulae (= gonapophyses of 8th segment, ventral valves and lancets) and unpaired 2nd valvulae (= medially fused gonapophyses of 9th segment, dorsal valve, and stylet). Basal parts of the 1st and 2nd valvulae are continuous with long curved dorsally processes (rami) which extend to the valvifers: the 1st ramus is connected to the 1st valvifer (= gonangulum; triangular plate), whereas the 2nd ramus is fused laterally with the anterior region of the 2nd valvifer, the region called the rostral process. The 1st and 2nd valvulae (and their rami) form a sliding interlocking mechanism called the olistheter which comprises a groove-like ventral component (= aulax) and a tongue-like dorsal one (= rhachis). The unpaired furcula attaches to the base of the 2nd valvula highly flexibly. The 1st valvifer possesses two articulations: the anterior one is with the modified 9th tergite (= quadrate plate), the posterior one with the 2nd valvifer (= proximal part of 9th gonocoxite). The very elastic membraneous incision (= incisura postarticularis, postincision) separates the anterior rostral process of the 2nd valvifer from its main body. The sting itself is ensheathed by paired 3rd valvulae (= distal part of 9th gonocoxite or sting sheaths) when

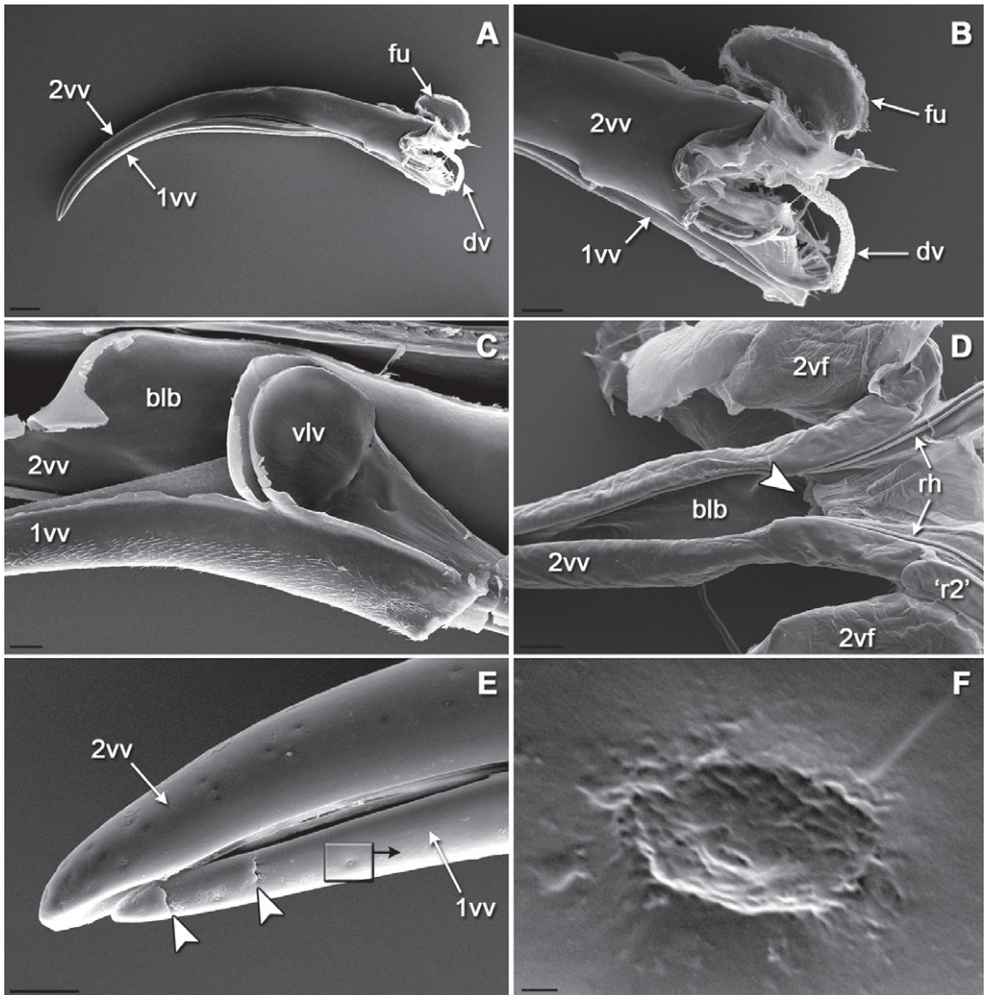


Figure 4. SEM morphology of the sting: **A** lateral view of sting with the furcula and venom duct **B** basal part of the sting **C** valvilli on the first valve positioned in the bulb of the second valve, medial view **D** ventral view of isolated second valve showing the bulb **E** lateral view of sting apex showing scattered sensilla-like structures on both valves and two subapical ridges on the first valve **F** single wrinkled sensilla-like structure that superficially resembles the secretory pores described by Nenon et al., 1995 (note characteristic contamination around the pore). Arrowheads indicate the position of the opening of the venom duct (in D) and the subapical barbs (in E). Scale bars: 200 μm in A; 100 μm in B and D; 30 μm in C and E; 300 nm in F.

not in use. The modified 8th tergite (= spiracle plate) is the outermost part of the sting apparatus and is connected neither with the sting valves nor with the valvifers.

Scanning electron microscopy. The entire external surface of the sting proper is mostly smooth, with only two subapical barbs on the 1st valvula (Figs 4A, E); however, there are numerous sensilla-like structures scattered over the entire surface, these are more numerous apically (Figs 4E, F). The dorsal surface of the 1st valvula forms an

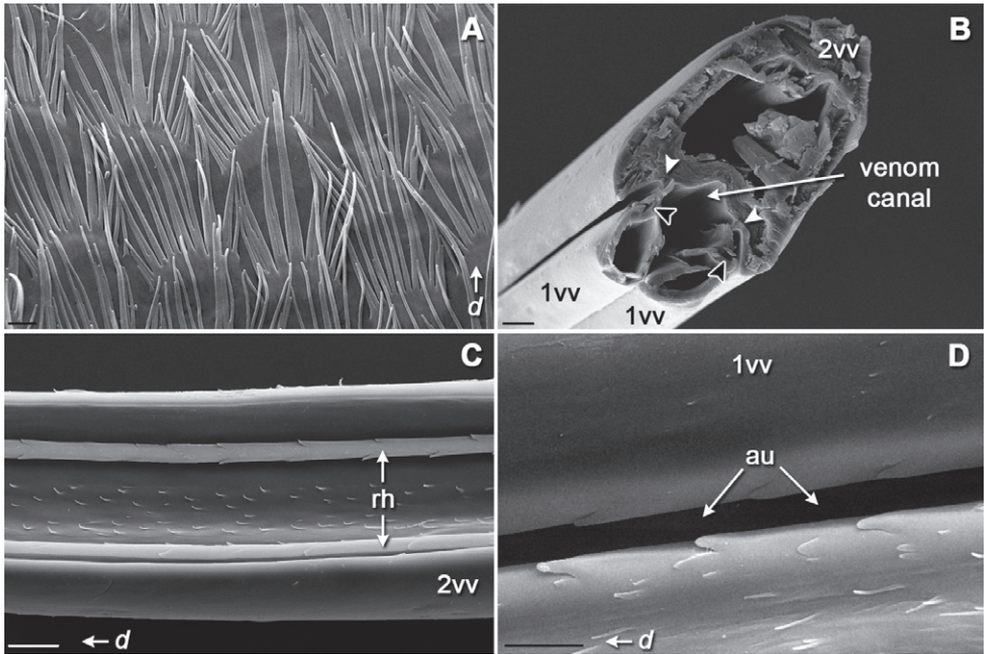


Figure 5. SEM micrographs of internal surfaces of the sting apparatus: **A** setose membrane of the genital chamber that contact the sting basis **B** transverse section of the aculeus showing position of the parts of the olistheter **C** ventral view of isolated second valve showing internal microsculpture between two rachises **D** dorsal view of aulax of first valve. *d*, distal direction. Arrowheads in **B** indicate position of rachises (white) and aulaxes (black) of the olistheter mechanism. Scale bars: 3 μm in **A** and **D**; 20 μm in **B** and **C**.

anterior swelling where two flaps called valvilli arise (Fig. 4C). These valvilli are housed within a broadened part of the 2nd valvula called the bulb. The venom duct enters the sting base at the anterior margin of the bulb (Figs 4A, B, D). Components of the olistheter are finely serrated, and the inner surface of the 2nd valvula bears distally directed microtrichia (Figs 5B–D). The setose membrane of the genital chamber covers the sting base (Fig. 5A). Three fields of sensilla are found on the 2nd valvifer. A row of styloconic sensilla is situated on the fused 2nd ramus and rostral process (Figs 3A, C). The seta of each sensillum in a row may touch the 1st ramus, and the more the 1st valvula protracts relative to 2nd valvula the more sensilla contact the 1st ramus. A field of 25–28 uniformly directed campaniform sensilla is situated more distally (Figs 3A, D). Another group of at least 35 styloconic sensilla that forms a setal plate is situated on the main sclerite of the 2nd valvifer where it articulates with the 1st valvifer (Figs 2A, 3B). The seta of each sensillum in this field may touch the 1st valvifer. A paired field of ca. 13 uniformly directed campaniform sensilla is also found on the anterolateral surface of the 2nd valvula where it articulates with the rostral process (Figs 2C, D). The well sclerotized 3rd valvula consists of two segments, the proximal and the distal (Figs 6A, C). Its inner wall is mainly membranous with a densely microsetose surface and bears

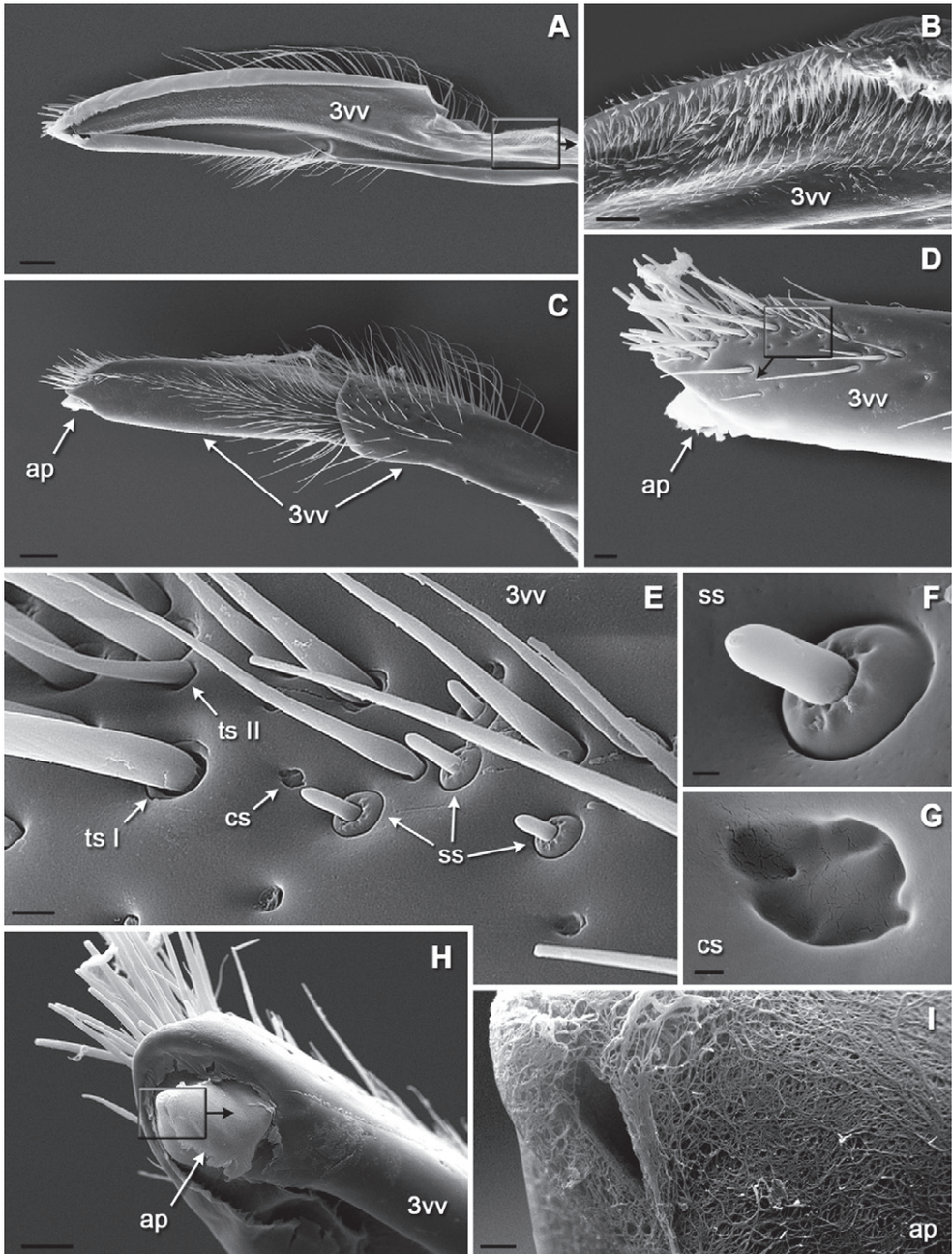


Figure 6. SEM morphology of the third valve: **A** medial view (inset enlarged in **B**) **B** microspination of membranous inner wall **C** lateral view **D** apex in lateral view (inset enlarged in **E**) **E** variety of sensilla on dorsoapical surface **F** socketed styloconic sensillum with apical pore **G** campaniform sensillum **H** apex in mediadorsal view showing membranous process (inset enlarged in **I**) **I** spongiform cuticle of an apical process. Scale bars: 100 μm in **A** and **C**; 20 μm in **B** and **D**; 5 μm in **E**; 1 μm in **F**; 0.5 μm in **G**; 30 μm in **H**; 2 μm in **I**.

a longitudinal ridge (Figs 6A, B). Apically, the inner membrane of the 3rd valvula forms a cone-shaped process composed of a peculiar spongy cuticle (Figs 6C, H, I). At rest, this apical process abuts closely to the dorsolateral surface of the 2nd valvula. There are at least four morphotypes of sensilla on the 3rd valvula (Figs 6D–G). Trichoid sensillum type I is socketed and has a rounded setal tip; these sensilla are aggregated apically. Trichoid sensillum type II is unsocketed and sharply pointed; these sensilla are the most abundant type, they are located all over the external surface but mostly at the apex and at the margin between proximal and distal parts of the 3rd valvula. Campaniform sensilla are rare and scattered among trichoid ones. Socketed styloconic sensilla bear a distinct pore on their rounded setal tip; these sensilla are found only on the dorsal surface of the 3rd valvulae between trichoid sensilla type I.

Discussion

Bembix rostrata is one of the most widely distributed and remarkable digger wasps species in Europe, and it often forms large colonies of dozen to hundred individuals. It prefers sandy and sunny habitats, where a female digs several burrows. As in most bembicine wasps, *B. rostrata* paralyzes the prey, which are predominantly large tabanid and syrphid flies, by inserting the sting through the venter of the thorax (Nielsen 1945, Evans 1957, Gadallah 2001). At this time, the sting of *B. rostrata* is functioning neither as an egg laying nor prey transporting device.

Radović (1985) has shown that the shape of the sting in apoid wasps is correlated with the degree of sclerotization and mobility of the prey: the wasps which prey on swift flying insects have a more strongly curved sting. This is the case in *B. rostrata* which has a markedly curved sting apparently correlated with the high speed of flight of their dipteran prey. The smooth external surface is a peculiar feature of the sting in most bembicine wasps studied so far (Gadallah and Assery 2004). Rare exceptions are three species of *Bembix* with barbed first valvulae: *B. arenaria* Handlirsch, 1893, *B. oculata* Panzer, 1801 (both after Gadallah and Assery 2004), and *B. rostrata* (this study). Apart from bembicine wasps (Crabronidae: Bembicinae), a barbed sting has been found in *Sericophorus relucens* F. Smith (Crabronidae: Crabroninae) which has barbs on first and second valvulae that may fasten the prey during its transportation on the sting (Radović and Sušić 1997). Interestingly, *S. relucens* also preys on flies. Similarly, species of the genus *Oxybelus* (Crabronidae: Crabroninae) possess spines on the first valvulae and transport their prey (muscid flies) impaled on the sting (Radović 1985). On the other hand, many unrelated species of apoid wasps have a barbed sting that is correlated with their preying on less sclerotized insects like caterpillars, aphids, cockroach nymphs, mantids, etc. (Radović 1985).

Several surfaces of the sting apparatus of *B. rostrata* are covered with unidirectional microstructures. The wall of the venom canal in *B. rostrata* is furnished with small microtrichia that are orientated distad. They are randomly scattered and relatively sparse, and seem to be somewhat reduced if compared with other non-aculeate hymenopter-

ans where they form a comb-like or ctenidial pattern (Quicke et al. 1999). This egg canal microsculpture is not restricted to Hymenoptera, but can be found across several insect orders that possess an ovipositor (Austin and Browning 1981). Here it functions as a “linear ratchet”, providing one-way movement of eggs along the ovipositor during longitudinal sliding of the valvulae. However, it is uncertain whether the microsculpture of the venom canal is involved in functioning of the sting in *B. rostrata*, which is not used for egg transport. The distally directed serration on the olistheter elements has been found in the sting of *B. rostrata* and also in the honeybee, *Apis mellifera* L. (Shing and Erickson 1982). The membranous cuticle of the genital chamber in *B. rostrata* that contacts with the sting base, as well as the inner walls of the third valvulae, are also covered with dense, distally directed microsetae of obscure functions. In parasitoid wasps, similar microsculpture on the inner walls of the third valvulae are supposed to be involved in cleaning of the ovipositor sensilla between oviposition episodes (Le Ralec et al. 1996, Quicke et al. 1999). On the other hand, a similar setose membrane in *A. mellifera* is known to produce and accumulate pheromones (Lensky et al. 1995, Martin et al. 2005). Noteworthy is that males of *B. rostrata* are able to differentiate between virgin or freshly copulated females and older females, by means of chemical cues which may be associated with the trunk (Schöne and Tengö 1981). It is not inconceivable that the female sting apparatus and associated structures can be a source of the aforesaid chemical cues.

The sensory equipment of the sting apparatus of *B. rostrata* is diverse. The sensilla can be divided into those that perceive intrinsic stimuli from the insect body (proprioceptors) and those detecting environmental factors (exteroceptors) (Quicke et al. 1999). Sensilla of the first type are represented in *B. rostrata* by two morphotypes, styloconic sensilla and campaniform sensilla, both often aggregated in fields. As in many other hymenopterans, *B. rostrata* possesses two fields of styloconic sensilla (hair plates) on the first valvifer that measure positional relationship of the first and second valvulae (Le Ralec et al. 1996, Quicke et al. 1999, Vilhelmsen et al. 2001, etc.). Two fields of campaniform sensilla have been found on the basal region of the second valvula and on its ramus for the first time in an apoid wasp. With regard to uniform transverse orientation of every sensillum in a field, they most likely function as detectors of lateral stresses and strains arising in the sting during prey penetration. Although campaniform sensilla on the third valvula are scattered, their orientation is also well-ordered. Exteroreceptors are located on the third valvula and are comprised of two morphotypes of trichoid sensilla and one morphotype of styloconic sensilla. Gadallah and Assery (2004) mentioned that trichoid sensilla on the third valvula in apoid wasps have a mechanosensory function. Possibly the differences in morphology of trichoid sensilla reflect some functional differences. The last group of sensilla that are of special interest are socketed styloconic sensilla with an apical pore, detected for the first time on the sting apparatus in an apoid wasp. It was found on the third valvula of *B. rostrata*. The presence of an apical pore implies that these sensilla are not exclusively mechanoreceptive and probably can perceive some chemical stimuli. No chemoreceptores have been recorded on the sting apparatus in apoid wasps so far.

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References

- Austin AD, Browning TO (1981) A mechanism for movement of eggs along insect ovipositors. *International Journal of Insect Morphology and Embryology* 10(2): 93–108.
- Cardinal S, Packer L (2007) Phylogenetic analysis of the corbiculate Apinae based on morphology of the sting apparatus (Hymenoptera: Apidae). *Cladistics* 23: 99–118.
- Evans HE (1957) Studies on the comparative ethology of digger wasps of the genus *Bembix*. Comstock Publishing Associates, Ithaca, New York, 248 pp.
- Gadallah NS (2001) A comparative morphological study of the skeletal parts of the sting apparatus in some *Stizus* species from Egypt (Sphecidae: Bembicinae). *Egyptian Journal of Zoology* 37: 255–265.
- Gadallah NS, Assery BM (2004) Comparative study of the skeletal parts of the sting apparatus in some sphecid species from Saudi Arabia (Hymenoptera: Sphecidae). *Linzer biologische Beiträge* 36(2): 1393–1412.
- Gauld ID, Bolton B (1988) *The Hymenoptera*. Oxford University Press, Oxford, 332 pp.
- Gnatzy W, Volkhardt W (2000) Venom gland of the digger wasp *Liris niger*: morphology, ultrastructure, age-related changes and biochemical aspects. *Cell and Tissue Research* 302(2): 271–284.
- Le Ralec A, Rabasse JM, Wajnberg E (1996) Comparative morphology of the ovipositor of some parasitic Hymenoptera in relation to characteristics of their hosts. *The Canadian Entomologist* 128: 413–433.
- Lensky Y, Cassier P, Tel-Zur D (1995) The setaceous membrane of honey bee (*Apis mellifera*) workers sting apparatus: structure and alarm pheromone distribution. *Journal of Insect Physiology* 41: 589–595.
- Martin SJ, Dils V, Billen J (2005) Morphology of the Dufour gland within the honey bee sting gland complex. *Apidologie* 36: 543–546.
- Nénon J-P, La Lannig J, Kacem N, Barbier R, Allo M-R (1995) Micromorphologie de l'ovipositeur des Hyménoptères et évolution des symphytes phytophages aux apocrites parasitoïdes. *Comptes rendus de l'Académie des sciences Paris, Sciences de la vie/Life Sciences* 318: 1045–1051.
- Nielsen ET (1945) *Moeurs des Bembex*. Monographie biologique avec quelques considérations sur la variabilité des habitudes. *Spolia Zoologica Musei Hauniensis* 7: 1–174.

- Oeser R (1961) Vergleichend-morphologische Untersuchungen über den Ovipositor der Hymenopteren. *Mitteilungen aus dem Zoologischen Museum in Berlin* 37: 3–119.
- Packer L (2003) Comparative morphology of the skeletal parts of the sting apparatus of bees (Hymenoptera: Apoidea). *Zoological Journal of the Linnean Society* 138: 1–38.
- Quicke DLJ, Fitton MG, Ingram SN (1992) Phylogenetic implications of the structure and distribution of ovipositor valvelli in the Hymenoptera (Insecta). *Journal of Natural History* 26(3): 587–608.
- Quicke DLJ, LeRalec A, Vilhelmsen L (1999) Ovipositor structure and function in the parasitic Hymenoptera with an exploration of new hypotheses. *Atti dell'Accademia Nazionale Italiana di Entomologia, Rendiconti* 47: 197–239.
- Radović IT (1985) Morphology and adaptive value of the sting apparatus of digger wasps (Hymenoptera : Sphecidae). *Acta entomologica Jugoslavica* 21 (1–2): 61–73.
- Radović IT, Sušić S (1997) Morphological characteristics of the sting and prey carriage mechanism in *Sericophorus relucens* F. Smith (Hymenoptera: Sphecidae: Larrinae). *Proceedings of the Entomological Society of Washington* 99(3): 537–540.
- Rahman MH, Fitton MG, Quicke DLJ (1998) Ovipositor internal microsculpture and other features in doryctine wasps (Insecta, Hymenoptera, Braconidae). *Zoologica Scripta* 21(4): 333–343.
- Rasnitsyn AP (1980) Origin and evolution of Hymenoptera. *Trudy paleontologicheskogo instituta akademii nauk SSR [Transactions of the Paleontological Institute, Academy of Science, USSR]* 174. Nauka Press, Moscow, 192 pp. (in Russian).
- Schöne H, Tengö J (1981) Competition of males, courtship behaviour and chemical communication in the digger wasp *Bembix rostrata* (Hymenoptera, Sphecidae). *Behaviour* 77(1–2): 44–65.
- Scudder GGE (1961) The comparative morphology of the insect ovipositor. *Transactions of the Royal Entomological Society of London* 113: 25–40.
- Shing H., Erickson EH (1982) Some ultrastructure of the honeybee (*Apis mellifera* L.) sting. *Apidologie* 13: 203–213.
- Smith EL (1970) Evolutionary morphology of the external insect genitalia. 2. Hymenoptera. *Annals of the Entomological Society of America* 63: 1–27.
- Snodgrass RE (1956) *Anatomy of the honey bee*. Cornell University Press, Ithaca, 334 pp.
- Vilhelmsen L (2000) The ovipositor apparatus of basal Hymenoptera (Insecta): phylogenetic implications and functional morphology. *Zoologica Scripta* 29 (4): 319–345.
- Vilhelmsen L, Isidoro N, Romani R, Basibuyuk HH, Quicke DLJ (2001) Host location and oviposition in a basal group of parasitic wasps: the subgenual organ, ovipositor apparatus, and associated structures in the Orussidae (Hymenoptera, Insecta). *Zoomorphology* 121: 63–84.