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Inter-caste communication in social insects Christoph Grüter¹ and Laurent Keller



Social insect colonies function as highly integrated units despite consisting of many individuals. This requires the different functional parts of the colony (e.g. different castes) to exchange information that aid in colony functioning and ontogeny. Here we discuss inter-caste communication in three contexts, firstly, the communication between males and females during courtship, secondly, the communication between queens and workers that regulate reproduction and thirdly, the communication between worker castes that allows colonies to balance the number of different worker types. Some signals show surprising complexity in both their chemistry and function, whereas others are simple compounds that were probably already used as pheromones in the solitary ancestors of several social insect lineages.

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Introduction

Insect colonies often consist of thousands — and sometimes millions — of individuals and the success of individuals depends crucially on the success of the colony [1,2]. Colonies show two kinds of division of labor. First, there is a reproductive division of labor between queens (and kings in termites) and the largely sterile workers. Second, there is division of labor among the workers for tasks like brood rearing, colony defense or foraging [1,2]. Communication between and within the different castes (queens, males and different worker groups) is fundamental for the efficient functioning of a colony. In order for colonies to respond to the often changing needs, workers — like the cells of multicellular organisms need to respond to signals in ways that are beneficial to the whole complex system. Most communication is based on chemical signals (or pheromones) that are produced by exocrine glands [1–4]. Hundreds of chemicals produced in more than 60 different glands have been identified in social insects [3,5], which has led researchers to refer to social insects as chemical factories [1]. Traditionally, pheromones have been divided into two classes, primer and releaser pheromones [1]. A releaser pheromone initiates an immediate behavioral response, whereas a primer pheromone alters more long-term endocrine and reproductive systems in the recipient [6]. However, it has become clear that there are pheromones that have both releaser and primer effects [6–8]. The pheromone signals are perceived via olfactory sensillae on the antennae $[3,9,10^{\circ},11^{\circ}]$ before being further processed by the olfactory system [12].

In this review we focus on recent advances in our understanding of inter-caste communication in three important contexts: firstly, communication between male and female reproductives that results in mating and, subsequently, colony foundation, secondly, communication between queens and workers to regulate reproduction and thirdly, communication between different functional groups of workers (sometimes called sub-castes) that allows colonies to balance the number of workers performing different tasks (for communication within castes, for example, among foragers during resource collection or during house hunting see [13–16]). Recent research has highlighted the importance of chemical and behavioral complexity, context, and dose for communication [6]. Furthermore, the recent identification of several queen signals that inhibit reproduction in workers [7,17^{••},18] or other queens [19] has improved our understanding of the evolution of reproductive division of labor. New tools like calcium imaging or genomics have started to reveal how pheromone signals are processed in the nervous system [12,20] and how external cues and signals induce important behavioral modifications that are associated with large scale changes in the pattern of gene expression in the brain (e.g. [20–22]).

Communication between males and females

Before starting a new colony reproductive individuals must find a mating partner. Chemical communication plays a fundamental role in this process and males in particular show numerous adaptations that help them find females [3]. These include large compound eyes, strong wing muscles or antennae with large numbers of odor receptors [23,24]. Most mating patterns fall into two broad categories, the 'female calling syndrome' and the 'male aggregation syndrome' [1,3]. In species with the 'female calling syndrome', females are often wingless and do not travel far from their natal nest (sometimes they call from within the nest). They release sex pheromones to 'call' the winged males. These species typically form small colonies and mating flights do not seem to be synchronized across colonies. In some cases it is the workers that 'call' males. In *Megaponera analis*, for example, males follow recruitment trails laid by workers to guide them to the nest [1]. In the stingless bee *Scaptotrigona postica*, 2alcohols attract males from long range, but additional substances are then required to elicit copulations by males [25]. Once males have settled near a nest containing a virgin queen, the males themselves become attractive to other males, leading to large aggregations (Figure 1a) [26,27]. Thus, male aggregations in many stingless bee species are probably the outcome of both female and male calling [26–28].

Species with larger colonies frequently exhibit a 'male aggregation syndrome', whereby males from many colonies gather at specific sites where they are later joined by

Figure 1



(a) A male aggregation in the Neotropical stingless bee *Tetragonisca angustula*. Aggregations of several hundred males are usually found close to a nest that contains a virgin queen or will soon contain a virgin queen (Photo by C. Grüter). (b) A fire ant (*Solenopsis invicta*) queen and her retinue workers. The workers feed and groom the queen and remove her eggs (Photo by W.R. Tschinkel).

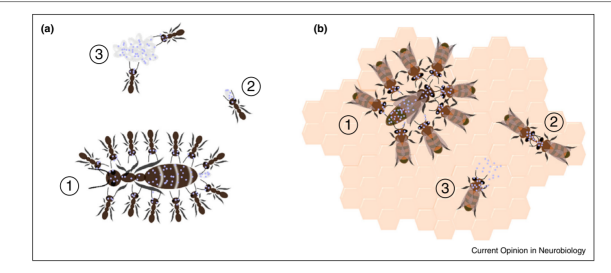
females [3]. In these species mating flights are usually tightly synchronized between many colonies thereby decreasing the probability of inbreeding and hybridization between closely related species. In species such as some *Camponotus*, Atta and Acromyrmex, the departure of both sexes from the nest is coordinated by the release of mandibular pheromones by males as they leave the nest. This, in turn, triggers the mass exodus of females [3]. Honey bee (Apis mellifera) drones gather at congregation areas (ranging from 30 to 200 m in diameter) where they wait for virgin queens [23,29]. A modification to this pattern of male aggregations is found in bumble bees where males patrol flight paths which they mark with labial gland secretions [30]. In Bombus terrestris, and probably other Bombus species, females are preferentially attracted to flight paths marked by many males, hence selecting for males to aggregate to attract females [3]. Another alteration to the 'male aggregation syndrome' is found in many polistine wasps, where males mark objects in their territories with scent by dragging their legs or abdomens over the substrate or release pheromones into the air [31,32].

How females locate male aggregations is not well understood but males' mandibular glands have been implicated as the source of the sex attractant in numerous ant genera and conclusively demonstrated in several Pogonomyrmex species [1,3]. The main compound found in mandibular glands in Pogonomyrex (4-methyl-3-heptanone) seems likely to play an important role in female attraction [3], although controlled experiments are needed to confirm this. Once females have found the aggregation femalederived pheromones are thought to attract males at close range. In Formica lugubris, for example, the source of the attractant is the females Dufour's gland, which contains undecane (90%), (Z)-4-tridecene (4%), and tridecane (4%). Synthetic undecane causes a strong male response similar to the response observed in nature [33]. One of the first identified sex pheromones is the honey bee 'queen substance', which originates from the queens' mandibular glands [34]. Mandibular gland extracts of honey bee queens attract drones from a distance of $\sim 50 \text{ m}$ [3,29]. The most active compound (9-keto-(E)-2-decenoic acid or 9-ODA) is almost as active as the whole blend [34]. Interestingly, Asian honeybee species also use 9-ODA as the main male attractant [35]. However, differences in the timing and location of mating flights and visual cues of sexual partners seem to limit heterospecific sexual encounters [3]. After mating, queens of some species quickly become unattractive to males. This can be caused by changes in the queens' cuticular hydrocarbon (CHC) profile (e.g. Leptothorax gredleri [36]) or by tactile signals produced by the queen (e.g. *Pogonomyrmex* spp. [1]).

Communication between queen and workers

An important prerequisite for the functioning of social insect colonies is the ability of queens to signal their presence and good health. To this end queens produce a





Distribution of queen pheromones (indicated as blue dots). (a) In ants and termites, workers acquire queen pheromone by physically interacting with a queen, for example, during grooming or feeding (1), when carrying queen-laid eggs (2) or when contacting egg piles (3). (b) Honey bee workers acquire queen pheromone when contacting the queen (1), when interacting with workers that had recently been in contact with the queen, for example, via trophallaxis with 'messenger' workers (2) or via the wax comb (3).

chemical signal that often has several effects, among which are the inhibition of the rearing of new reproductives [6,37[•]], attraction of workers to the queen (Figure 1b) [6,37[•]], the suppression of worker reproduction [6,7,17^{••},38–40] and chemical marking of eggs, which allows workers to recognize whether eggs are queen-laid or worker-laid [39,41]. The absence of the queen signal, which generally is associated with the death of the queen, usually elicits important changes in the colony, including the rearing of new queens [42] or the activation of worker ovaries to produce male brood. There has been considerable controversy about whether queen signals are manipulative tools allowing queens to prevent workers from reproducing or whether they are cooperative signals that allow workers to increase their fitness [5,43–45]. The idea behind the second hypothesis is that because worker reproduction may impose costs for colony productivity, workers may increase their inclusive fitness by not reproduction in the presence of the queen [39,46]. Several lines of evidence support the hypothesis that queen signals are indeed honest signals of fertility, rather than an attempt by the queen to manipulate the workers [47[•]]. First, the 'honest signal' hypothesis predicts a positive correlation between signal production and fertility and there is increasing support for this prediction in ants [48,49]. A possible explanation for this positive correlation is that queen signals are derived from chemical by-products of ovarian development [38,41,47[•]]. Second, comparative studies show that workers often ignore queen signals and try to reproduce in the presence of a fertile queen if it is in their genetic interest to do so [47°,50]. Third, the 'queen manipulation' hypothesis predicts an evolutionary arms race between queens and workers that leads to the

this prediction, recent research suggests that structurally related non-volatile hydrocarbons function as conserved queen signals in several lineages of social insects, including ants, wasps, bumble bees and stingless bees [17^{••},18,38,47[•],51], but see [52[•]], suggesting that queen pheromones are derived from ancestral communication systems that were already present in the common solitary ancestor of ants, wasps and bees (with the exception of the honey bee, see below) [17^{••},47[•]]. However, definitive evidence for queen pheromones being honest signals will require a better mechanistic understanding of how the compounds affect workers and queens, for example whether queen-produced pheromones exclusively bind to antennal receptors (which would support the view that they are honest signals) or whether they also enter the worker circulatory system and mediate hormonal changes directly affecting reproduction (which would be consistent with pheromonal queen control) [43].

rapid evolution of chemical signals [43,45]. Contrary to

One challenge for large colonies is an efficient distribution of the queen signal. Research in ants and termites shows that the signal is located on both the queen and on her eggs [7,19,37°,39]. This means that the queen presence can be felt in parts of the nest where the queen is not present (Figure 2a). In honey bees, eggs are not carried around, but there are specialist 'messengers' whose role is to actively distribute the queen pheromone in the colony [42]. Small amounts of pheromone are also transmitted via the wax comb (Figure 2b) [53]. Again, 9-ODA plays an important role but, unlike in the attraction of drones, worker attraction requires the combined presence of four additional mandibular compounds (some strains may require even more compounds) [6,54]. The combination of 9-ODA and these four compounds constitute the queen mandibular pheromone which has been shown to alter the pattern of expression of several hundred genes in the brain of adult workers [20,55], including genes implicated in the dopamine pathways [56] and genes associated with the behavioral maturation of workers from nurses to foragers [20].

Communication between worker sub-castes

Division of labor among workers is an important reason for the ecological success of social insects [1,2]. A key challenge for a colony is to allocate an appropriate number of workers to the different tasks. The number of soldiers in a colony, for example, should match the level of threat a colony faces [57,58]. Research has shown that pheromones that function as negative and positive feedbacks play a crucial role in balancing the number of workers performing different tasks. In honey bees, for example, the number of workers that progress from nurse duties to foraging duties as they age depends on the number of foragers already present in the colony [59,60]. Foragers produce ethyl oleate, which acts as a chemical inhibitory factor delaying the onset of foraging [61]. The removal of foragers leads to increased levels of juvenile hormone (JH) titers, which is correlated with the onset of foraging behavior [62]. Ethyl oleate is synthesized de novo and is present in highest concentrations in the bee's crop. Thus, behavioral maturation and the propensity to become a forager is modulated via trophallaxis [61], a form of food exchange that plays a prominent role as an information channel in various contexts [63]. A pheromone produced by larvae has a similar effect: colonies treated with a synthetic blend of this pheromone show delayed foraging activities compared to workers in untreated control colonies [8].

Colony defense is another important task and many species rely on a specialist soldier caste for their safety [1,2,64,65]. Colonies in some species are able to adjust soldier production according to the dangers in the environment [57,58]. An overproduction of new soldiers is prevented because the presence of existing soldiers inhibits the production of new soldiers. Such soldier inhibition has been in demonstrated in aphids [66], termites [67] and ants [68] but the pheromones causing this effect have not yet been identified.

Caste specific response to pheromones

Recent research has started to elucidate caste-specific differences in the olfactory system that underlie caste-specific responses to pheromones. For instance, males, queens and workers differ in the number of sensory sensillae on the antennae [69,70] and in the expression of a range of odorant receptors (ORs), some of which are known to respond to components of queen pheromone [9,71]. The number of sensory sensillae, in turn, has been

shown to correlate with the number of glomeruli, the functional units of the antennal lobes (AL, the primary olfactory center) [69]. In honeybees, queens and workers show morphological differences in both the primary (AL) and secondary (mushroom bodies) olfactory center: queens have smaller glomeruli and fewer microglomeruli [72]. In several ants and the honeybee, males and females differ in the number and size of glomeruli in the antennal lobes. Females have more glomeruli, but males have some large macroglomeruli that are less common or absent in the female castes [12,73-75]. Additionally, glomeruli in males may contain different types of innervations than in queens and workers [73]. Calcium imaging technology has revealed that odors, including pheromones, trigger specific activity patterns in the glomeruli [12,76–78]. This was shown in honeybees where harnessed individuals were simultaneously exposed to floral or pheromonal odors while measuring the spatio-temporal excitation patterns evoked in the glomeruli [76–78]. These studies demonstrate that odor identity in the AL is coded in odor-specific activity patterns that involve the combined activity of a number of glomeruli and that different classes of odors tend to activate different glomeruli.

Concluding remarks

Communication systems are a prerequisite for the functioning of complex biological systems in general and animal societies in particular. Yet, despite decades of research we still have a superficial understanding of the identity of the chemicals and the corresponding receptors that are involved and their location of action. The development of new molecular tools has started to shed light on these mechanistic questions and future work will allow us to gain a better understanding of how systems of chemical communication evolved to regulate the behavior and physiology of individuals and societies.

Conflict of interest statement

Nothing declared.

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References and recommended reading

Papers of particular interest, published within the period of review, have been highlighted as:

- · of special interest
- •• of outstanding interest
- 1. Hölldobler B, Wilson EO: *The Ants*. Cambridge, MA: The Belknap Press of Harward University; 1990.
- Hölldobler B, Wilson EO: The Superorganism: The Beauty, Elegance, and Strangeness of Insect Societies. New York: W. W. Norton & Company; 2009.

- Ayasse M, Paxton R, Tengö J: Mating behavior and chemical 3. communication in the order Hymenoptera. Ann Rev Entomol 2001. 46:31-78
- 4. Richard F-J, Hunt J: Intracolony chemical communication in social insects. Insectes Sociaux 2013, 60:275-291
- Heinze J, d'Ettorre P: Honest and dishonest communication in 5. social Hymenoptera. J Exp Biol 2009, 212:1775-1779.
- 6. Slessor KN, Winston ML, Le Conte Y: Pheromone communication in the honeybee (Apis mellifera L.). J Chem Ecol 2005, 31:2731-2745.
- 7. Holman L, Jørgensen CG, Nielsen J, d'Ettorre P: Identification of an ant queen pheromone regulating worker sterility. Proc R Soc Lond B: Biol Sci 2010, 277:3793-3800.
- Le Conte Y, Mohammedi A, Robinson GE: Primer effects of a 8. brood pheromone on honeybee behavioural development. Proc R Soc Lond B: Biol Sci 2001, 268:163-168.
- Wanner KW, Nichols AS, Walden KK, Brockmann A, Luetje CW, 9 Robertson HM: A honey bee odorant receptor for the queen substance 9-oxo-2-decenoic acid. Proc Natl Acad Sci U S A 2007, 104:14383-14388.
- 10. Villar G, Baker TC, Patch HM, Grozinger CM: Neurophysiological mechanisms underlying sex-and maturation-related variation in pheromone responses in honey bees (Apis mellifera). J Compar Physiol A 2015, 201:731-739.

This study shows a link between the expression of the antennal receptor AmOR11 and behavioral sensitivity to a major compound of the honey bee queen pheromone, 9-ODA

11. Sharma KR, Enzmann BL, Schmidt Y, Moore D, Jones GR,

Parker J, Berger SL, Reinberg D, Zwiebel LJ, Breit B: Cuticular hydrocarbon pheromones for social behavior and their coding in the ant antenna. *Cell Rep* 2015, **12**:1261-1271.

The authors identify and study the olfactory sensillae on antennae that allow ants to identify other individuals based on cuticular hydrocarbons.

- Sandoz J-C: Odour-evoked responses to gueen pheromone 12. components and to plant odours using optical imaging in the antennal lobe of the honey bee drone Apis mellifera L. J Exp Biol 2006, 209:3587-3598.
- 13. Morgan DE: Trail pheromones of ants. Physiol Entomol 2009, 34:1-17.
- Seeley TD: Honeybee Democracy. Princeton: Princeton University 14. Press; 2010.
- Franklin EL: The journey of tandem running: the twists, turns 15. and what we have learned. Insectes Sociaux 2014, 61:1-8.
- 16. Czaczkes TJ, Grüter C, Ratnieks FLW: Trail pheromones: an integrative view of their role in colony organization. Ann Rev Entomol 2015, 60:581-599.
- 17. Van Oystaeyen A, Oliveira RC, Holman L, van Zweden JS,
 Romero C, Oi CA, D'Ettorre P, Khalesi M, Billen J, Wäckers F et al.: Conserved class of queen pheromones stops social insect workers from reproducing. Science 2014, 343:287-290.

The authors newly identify 3 queen pheromones (in an ant, a wasp and a bumblebee) and provide a phylogenetic analysis that reveals the conserved nature of queen pheromones in many social insect species.

- Smith AA, Millar JG, Suarez AV: A social insect fertility signal is 18. dependent on chemical context. Biol Lett 2015, 11:20140947.
- 19. Yamamoto Y, Matsuura K: Queen pheromone regulates egg production in a termite. Biol Lett 2011, 7:727-729.
- 20. Grozinger CM, Sharabash NM, Whitfield CW, Robinson GE: Pheromone-mediated gene expression in the honey bee brain. Proc Natl Acad Sci U S A 2003, **100**:14519-14525.
- 21. Dolezal AG, Toth AL: Honey bee sociogenomics: a genomescale perspective on bee social behavior and health. Apidologie 2014, 45:375-395.
- 22. Zayed A, Robinson GE: Understanding the relationship between brain gene expression and social behavior: lessons from the honey bee. Ann Rev Genet 2012, 46:591-615.

- 23. Winston ML: The Biology of the Honey Bee. Cambridge, Massachusetts: Harvard University Press; 1987.
- 24. Brockmann A, Brückner D: Structural differences in the drone olfactory system of two phylogenetically distant Apis species, A. florea and A. mellifera. Naturwissenschaften 2001, 88:78-81.
- 25. Engels W, Engels E, Francke W: Ontogeny of cephalic volatile patterns in queens and mating biology of the neotropical stingless bee, Scaptotrigona postica. Invertebr Reprod Dev 1997, **31**:251-256.
- 26. Galindo López JC, Kraus FB: Cherchez la femme? Site choice of drone congregations in the stingless bee Scaptotrigona mexicana. Anim Behav 2009. 77:1247-1252
- 27. Fierro MM, Cruz-López LDS, Villanueva-Gutiérrez R, Vandame R: Queen volatiles as a modulator of Tetragonisca angustula drone behavior. J Chem Ecol 2011, 37:1255-1262.
- 28. Verdugo-Dardon M, Cruz-Lopez L, Malo E, Rojas J, Guzman-Diaz M: Olfactory attraction of Scaptotrigona mexicana drones to their virgin queen volatiles. Apidologie 2011, 42:543-550.
- 29. Brockmann A, Dietz D, Spaethe J, Tautz J: Beyond 9-ODA: sex pheromone communication in the European honey bee Apis mellifera L. J Chem Ecol 2006, **32**:657-667.
- 30. Bergman P, Bergström G: Scent marking, scent origin, and species specificity in male premating behavior of two Scandinavian bumblebees. J Chem Ecol 1997, 23:1235-1251.
- 31. Kasuya E: Male mating territory in a Japanese paper wasp, Polistes jadwigae Dalla Torre (Hymenoptera, Vespidae). Kontyu 1981, 49:607-614.
- 32. Beani L, Calloni C: Leg tegumental glands and male rubbing behavior at leks in Polistes dominulus (Hymenoptera: Vespidae). J Insect Behav 1991, 4:449-462
- Walter F, Fletcher DJC, Chautems D, Cherix D, Keller L, Francke W, Fortelius W, Rosengren R, Vargo EL: Identification of 33. the sex pheromone of an ant, Formica lugubris (Hymenoptera, Formicidae). Naturwissenschaften 1993, 80:30-34
- 34. Gary NE: Chemical mating attractants in the gueen honey bee. Science 1962, 136:773-774.
- 35. Koeniger G, Koeniger N, Phiancharoen M: Comparative reproductive biology of honeybees. Honeybees of Asia. Springer; 2011:: 159-206.
- 36. Oppelt A, Heinze J: Mating is associated with immediate changes of the hydrocarbon profile of Leptothorax gredleri ant queens. J Insect Physiol 2009, 55:624-628.
- 37. Matsuura K, Himuro C, Yokoi T, Yamamoto Y, Vargo EL, Keller L:

Identification of a pheromone regulating caste differentiation in termites. *Proc Natl Acad Sci U S A* 2010, **107**:12963-12968. The authors report the identification of the first pheromone that inhibits reproduction in termites.

- 38. Nunes TM, Mateus S, Favaris AP, Amaral MF, von Zuben LG, Clososki GC, Bento JM, Oldroyd BP, Silva R, Zucchi R: Queen signals in a stingless bee: suppression of worker ovary activation and spatial distribution of active compounds. Sci Rep 2014, 4:7449.
- 39. Endler A, Liebig J, Schmitt T, Parker JE, Jones GR, Schreier P, Hölldobler B: Surface hydrocarbons of queen eggs regulate worker reproduction in a social insect. Proc Natl Acad Sci USA 2004, 101:2945-2950.
- 40. Tan K, Liu X, Dong S, Wang C, Oldroyd BP: Pheromones affecting ovary activation and ovariole loss in the Asian honey bee Apis cerana. J Insect Physiol 2015, 74:25-29.
- 41. Oi CA, Van Oystaeyen A, Oliveira RC, Millar JG, Verstrepen KJ, van Zweden JS, Wenseleers T: Dual effect of wasp queen pheromone in regulating insect sociality. Curr Biol 2015, **25**:1638-1640.
- Seeley TD: Queen substance dispersal by messenger workers 42. in honeybee colonies. Behav Ecol Sociobiol 1979, 5:391-415.
- 43. Keller L: Adaptation and the genetics of social behaviour. Philos Trans R Soc B: Biol Sci 2009, 364:3209-3216.

- 44. van Zweden JS: The evolution of honest queen pheromones in insect societies. *Commun Integr Biol* 2010, **3**:50-52.
- 45. Keller L, Nonacs P: The role of queen pheromones in social insects: queen control or queen signal? *Anim Behav* 1993, 45:787-794.
- 46. Ratnieks FLW, Foster KR, Wenseleers T: Conflict resolution in insect societies. Ann Rev Entomol 2006, 51:581-608.
- 47. Oi CA, Van Zweden JS, Oliveira RC, Van Oystaeyen A,
- Nascimento DL, Wenseleers R: The origin and evolution of social insect queen pheromones: novel hypotheses and outstanding problems. *BioEssays* 2015, 37:808-821.

The authors review the debate about queen pheromones and their function as either honest signals vs. tools of worker manipulation.

- Endler A, Liebig J, Hölldobler B: Queen fertility, egg marking and colony size in the ant Camponotus floridanus. Behav Ecol Sociobiol 2006, 59:490-499.
- Holman L, Dreier S, D'Ettorre P: Selfish strategies and honest signalling: reproductive conflicts in ant queen associations. Proc R Soc Lond Ser B-Biol Sci 2010, 277:2007-2015.
- Wenseleers T, Ratnieks FLW: Enforced altruism in insect societies. Nature 2006, 444:50.
- van Zweden JS, Bonckaert W, Wenseleers T, d'Ettorre P: Queen signaling in social wasps. Evolution 2014, 68:976-986.
- 52. Amsalem E, Orlova M, Grozinger CM: A conserved class of queen pheromones? Re-evaluating the evidence in bumblebees (Bombus impatiens). Proc R Soc B: Biol Sci 2015, 282:20151800

The findings presented in this study challenge the finding that queen pheromones in bumblebees are conserved across species and groups of species.

- Naumann K, Winston ML, Slessor KN, Prestwich GD, Webster FX: Production and transmission of honey bee queen (Apis mellifera L.) mandibular gland pheromone. Behav Ecol Sociobiol 1991, 29:321-332.
- Keeling CI, Slessor KN, Higo HA, Winston ML: New components of the honey bee (*Apis mellifera* L.) queen retinue pheromone. Proc Natl Acad Sci U S A 2003, 100:4486-4491.
- Chandrasekaran S, Ament SA, Eddy JA, Rodriguez-Zas SL, Schatz BR, Price ND, Robinson GE: Behavior-specific changes in transcriptional modules lead to distinct and predictable neurogenomic states. Proc Natl Acad Sci U S A 2011, 108:18020-18025.
- Beggs KT, Glendining KA, Marechal NM, Vergoz V, Nakamura I, Slessor KN, Mercer AR: Queen pheromone modulates brain dopamine function in worker honey bees. Proc Natl Acad Sci U S A 2007, 104:2460-2464.
- Passera L, Roncin E, Kaufmann B, Keller L: Increased soldier production in ant colonies exposed to intraspecific competition. *Nature* 1996, 379:630-631.
- Segers FHID, Von Zuben LG, Grüter C: Local differences in parasitism and competition shape defensive investment in a polymorphic eusocial bee. *Ecology* 2015. (in press).
- Robinson GE: Regulation of division of labor in insect societies. Ann Rev Entomol 1992, 37:637-665.
- Huang Z, Robinson GE: Regulation of honey bee division of labor by colony age demography. Behav Ecol Sociobiol 1996, 39:147-158.
- Leoncini I, Le Conte Y, Costagliola G, Plettner E, Toth AL, Wang M, Huang Z, Bécard JM, Crauser D, Slessor KN et al.: Regulation of behavioral maturation by a primer pheromone produced by adult worker honey bees. Proc Natl Acad Sci U S A 2004, 101:17559-17564.

- Huang Z, Robinson GE: Honeybee colony integration: workerworker interactions mediate hormonally regulated plasticity in division of labor. Proc Natl Acad Sci U S A 1992, 89:11726-11729.
- Farina WM, Grüter C: Trophallaxis a mechanism of information transfer. In Food Exploitation by Social Insects: Ecological, Behavioral, and Theoretical Approaches. Edited by Jarau S, Hrncir M. CRC Press; 2009:173-187.
- 64. Tian L, Zhou X: The soldiers in societies: defense, regulation, and evolution. Int J Biol Sci 2014, 10:296-308.
- Grüter C, Menezes C, Imperatriz-Fonseca VL, Ratnieks FLW: A morphologically specialized soldier caste improves colony defence in a Neotropical eusocial bee. Proc Natl Acad Sci U S A 2012, 109:1182-1186.
- Shibao H, Kutsukake M, Fukatsu T: Density-dependent induction and suppression of soldier differentiation in an aphid social system. J Insect Physiol 2004, 50:995-1000.
- Haverty MI, Howard RW: Production of soldiers and maintenance of soldier proportions by laboratory experimental groups of *Reticulitermes flavipes* (Kollar) and *Reticulitermes virginicus* (Banks) (Isoptera: Rhinotermitidae). Insectes Sociaux 1981, 28:32-39.
- Passera L: Production des soldats dans les sociétés sortant d'hibernation chez la fourmi Pheidole pallidula (Nyl.) (Formicidae, Myrmicinae). Insectes Sociaux 1977, 24:131-146.
- Mysore K, Shyamala BV, Rodrigues V: Morphological and developmental analysis of peripheral antennal chemosensory sensilla and central olfactory glomeruli in worker castes of *Camponotus compressus* (Fabricius, 1787). Arthropod Struct Dev 2010, 39:310-321.
- Esslen J, Kaissling K-E: Zahl und Verteilung antennaler Sensillen bei der Honigbiene (Apis mellifera L.). Zoomorphologie 1976, 83:227-251.
- Zhou X, Rokas A, Berger SL, Liebig J, Ray A, Zwiebel LJ: Chemoreceptor evolution in Hymenoptera and its implications for the evolution of eusociality. *Genome Biol Evol* 2015, 7: 2407-2416.
- Groh C, Rössler W: Caste-specific postembryonic development of primary and secondary olfactory centers in the female honeybee brain. Arthropod Struct Dev 2008, 37: 459-468.
- Zube C, Rössler W: Caste-and sex-specific adaptations within the olfactory pathway in the brain of the ant Camponotus floridanus. Arthropod Struct Dev 2008, 37:469-479.
- 74. Mysore K, Subramanian K, Sarasij R, Suresh A, Shyamala BV, VijayRaghavan K, Rodrigues V: Caste and sex specific olfactory glomerular organization and brain architecture in two sympatric ant species Camponotus sericeus and Camponotus compressus (Fabricius, 1798). Arthropod Struct Dev 2009, 38:485-497.
- Kuebler LS, Kelber C, Kleineidam CJ: Distinct antennal lobe phenotypes in the leaf-cutting ant (Atta vollenweideri). J Compar Neurol 2010, 518:352-365.
- Joerges J, Küttner A, Galizia CG, Menzel R: Representations of odours and odour mixtures visualized in the honeybee brain. *Nature* 1997, 387:285-288.
- Galizia CG, Menzel R: Odour perception in honeybees: coding information in glomerular patterns. *Curr Opin Neurobiol* 2000, 10:504-510.
- Galizia CG, Menzel R: The role of glomeruli in the neural representation of odours: results from optical recording studies. J Insect Physiol 2001, 47:115-130.