Olfactory eavesdropping between two competing stingless bee species

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Abstract Foragers can improve search efficiency, and ultimately fitness, by using social information: cues and signals produced by other animals that indicate food location or quality. Social information use has been well studied in predator–prey systems, but its functioning within a trophic level remains poorly understood. Eavesdropping, use of signals by unintended recipients, is of particular interest because eavesdroppers may exert selective pressure on signaling systems. We provide the most complete study to date of eavesdropping between two competing social insect species by determining the glandular source and composition of a recruitment pheromone, and by examining reciprocal heterospecific responses to this signal. We tested eavesdropping between Trigona hyalinata and Trigona spinipes, two stingless bee species that compete for floral resources, exhibit a clear dominance hierarchy and recruit nestmates to high-quality food sources via pheromone trails. Gas chromatography–mass spectrometry of T. hyalinata recruitment pheromone revealed six carboxylic esters, the most common of which is octyl octanoate, the major component of T. spinipes recruitment pheromone. We demonstrate heterospecific detection of recruitment pheromones, which can influence heterospecific and conspecific scout orientation. Unexpectedly, the dominant T. hyalinata avoided T. spinipes pheromone in preference tests, while the subordinate T. spinipes showed neither attraction to nor avoidance of T. hyalinata pheromone. We suggest that stingless bees may seek to avoid conflict through their eavesdropping behavior, incorporating expected costs associated with a choice into the decision-making process.

Keywords Social information · Interceptive eavesdropping · Decision making · Dominance · Foraging · Cephalic labial glands

Abbreviations
GC-MS Gas chromatography–mass spectrometry
LGE Labial gland extract
w/w Weight/weight

Introduction
Animals at multiple trophic levels actively search for patchily distributed food such as mobile prey, flowering
or fruiting trees, or carrion. Such consumers can improve search efficiency, and ultimately fitness, by using information provided by the food itself or other organisms in the vicinity (Giraldeau 1997; Dornhaus and Chittika 2004). Use of social information (sensu Danchin et al. 2004) by foragers appears to be widespread. Information can come from signals (features or behaviors that have evolved to alter the behavior of the receiver in a specific way) or cues, which did not evolve because of such effects (Maynard Smith and Harper 2003). When signals provide such information, unintended receivers that use it are exhibiting “interceptive eavesdropping.” Because signals evolve through selection for information flow, they are vulnerable to selective pressures exerted by eavesdroppers (Peake 2005). Evolutionary and ecological effects of eavesdropping may be particularly strong and diverse in the context of foraging because resultant increases in food discovery efficiency cascade through food webs (Kean et al. 2003). Many examples show predators and prey benefiting from social information to locate prey and avoid predators, respectively (e.g., sources in Stowe et al. 1995; Peake 2005; Seppänen et al. 2007; Valone 2007). However, social information can also improve search efficiency within a trophic level. In this latter context, heterospecific eavesdropping (on signals) and “spying” (using social information provided by cues; Wisenden and Stacey 2005) can affect community structure. Such strategies can (1) increase the frequency of interaction among competitors (Seppänen et al. 2007) or (2) drive the formation and maintenance of foraging groups (Goodale et al. 2010) that provide benefits (e.g., protection) that overcome costs of food sharing (Stevens and Gilby 2004).

Despite the ecological implications of eavesdropping, little is known about how dominant and subordinate species competing for food use social information. To date, only a handful of studies have investigated interceptive eavesdropping on food location or quality signals by heterospecifics. Exploitation of heterospecific food location cues has also received some attention, primarily with social insects. Experiments suggest that subordinate species can avoid competitors (e.g., Pimm et al. 1985; Fletcher 2008; Evans et al. 2009; Slaa and Hughes 2009) or depleted resources (e.g., Nakashima et al. 2002; Yokoi et al. 2007) by using heterospecific visitation signals and cues. This latter phenomenon may be quite sophisticated; bumble bees can learn to be attracted or repelled by social information depending on their past experience with the food source, use it to determine when flowers replenish their nectar, and increase rejection of visited flowers when the visitor was an aggressive species. Avoidance of depleted resources through detection of chemical or visual cues likely saves bees time (Goulson 2009), thereby increasing foraging efficiency. Dominant species, on the other hand, are sometimes attracted to food cues and signals of heterospecific competitors (reviewed in Slaa and Hughes 2009; Goodale et al. 2010). This diversity of responses suggests the rules governing eavesdropping and spying within a trophic level are more complex than in predator–prey situations and require more sophisticated decision making (Coolen et al. 2003). To understand these rules, and the effects of social information on communities and signaling systems, we must further investigate within-trophic-level eavesdropping and spying across a range of species.

Highly social insects are an excellent system for studying eavesdropping and spying by competitors. Species often compete for food with sympatric relatives (Hubbell and Johnson 1977; Hölldobler and Wilson 1990; Eltz et al. 2002) and exhibit clear dominant–subordinate relationships (e.g., Fellers 1987; Lichtenberg et al. 2010). Social insects combine excellent associative learning (Dukas 2008) with powerful olfactory detection (Greenfield 2002) for successful foraging. Using social information may thus provide fitness benefits through improved search efficiency. Both ants and several genera of eusocial stingless bees (Apidae, Meliponini) recruit nestmates by depositing attractive pheromones at a high-quality food source or as a trail between food and nest (Hölldobler and Wilson 1990; Nieh 2004). Because recruitment pheromones are signals present in the public domain, they are susceptible to eavesdropping. Stingless bee trails may be particularly at risk because the trails are much less heavily guarded than are ant foraging trails (EML, personal observation). Where ant eavesdropping has been observed, it is typically between parabiotic “garden ant” species that share nests in epiphytes and thus are highly likely to encounter each others’ trails (Slaa and Hughes 2009).

Research shows that both intra- (Boogert et al. 2006; Jarau 2009) and interspecific (Nieh et al. 2004a) eavesdropping occurs in stingless bees. Coupled with patterns of response to the visual presence of heterospecifics on flowers (Slaa et al. 2003) and anecdotal observations (Kerr et al. 1963; Johnson and Hubbell 1975; Johnson 1983), these studies suggest that stingless bees actively avoid food sources of decreased resource quality or to which they will have limited access. In particular, stingless bees appear to avoid resources occupied by dominant species, thereby steering clear of conflict (the dominance motivation hypothesis). Dominant species may also benefit by following subordinates’ pheromone trails, using this social information to discover high-quality food sources that they can take over. Because stingless bees serve as major pollinators of tropical plants (Endress 1994), eavesdropping interactions between sympatric colonies may significantly affect bees’ foraging patterns and, ultimately, plant gene flow.

Here, we tested olfactory eavesdropping between two trail-laying stingless bee species that have a clear domi-
nance relationship. *Trigona hyalinata* and *Trigona spinipes* overlap in distribution (Camargo and Pedro 2007), exhibit similar floral utilization (Lichtenberg et al. 2010; unpublished data), and likely compete for resources. Both species use odor trails to recruit large numbers of nestmates to rich resources such as mass-flowering trees and sucrose feeders (Nieh et al. 2003; Nieh et al. 2004b). *Trigona hyalinata* foragers easily displace *T. spinipes* from food sources (Lichtenberg et al. 2010), both by arriving at a food source en masse and by attacking individual *T. spinipes* foragers (see Supplemental movies). In *T. spinipes* and all other trail-laying stingless bee species studied to date, recruitment pheromones come from the cephalic labial glands (Jarau 2009). An eavesdropper must both detect the target pheromone and distinguish it from its own. To show that eavesdropping is possible between these two species, we determined the chemical composition and attractiveness to nestmates of *T. hyalinata* labial gland secretions. The pheromone of *T. spinipes* is already known to have one main component: octyl octanoate (Schorkopf et al. 2007). We then tested eavesdropping between these species with preference tests. Under the dominance motivation hypothesis, we predicted that the subordinate *T. spinipes* (Lichtenberg et al. 2010) would avoid *T. hyalinata* recruitment pheromone, while the dominant *T. hyalinata* would be attracted to *T. spinipes* pheromone.

**Materials and methods**

**Study site and colonies**

We conducted research at the Universidade de São Paulo, Ribeirão Preto, in southeastern Brazil. This site is in the cerrado ecoregion and provides suitable stingless bee habitat in an otherwise urban and agricultural landscape. Multiple colonies of both *T. hyalinata* and *T. spinipes* inhabit the campus, although only a few colonies were accessible for experimentation because both species tend to nest close to the crowns of trees. We tested eavesdropping between two heterospecific colony pairs that were 100 m (2008 field season) and 500 m (2009) apart, and thus within flight range of each other (Kerr 1959). Only these four colonies could be paired with heterospecific colonies within flight range and were sufficiently low to the ground (<10 m) to allow us to train foragers to feeders.

Bees were trained to visit a training feeder located 10–15 m from their nest (exact distance depended on topography) and providing 15% weight/weight (w/w) sucrose solution (0.46 M). One *T. hyalinata* colony was less motivated to visit the feeder and thus was fed 30% w/w (0.99 M) sucrose solution. Stingless bees are known to collect nectar ranging from 5% to 67% w/w sugar, with an average nectar quality of 41% (Roubik et al. 1995). We used the relatively weak 15% sucrose solution, which limits overly intense recruitment, because each colony was visiting feeders for at least a month and we did not want nests to become filled with stored honey. To train bees, we placed cotton saturated with sucrose solution at the nest entrance. When necessary, we used a pole and climbed a ladder or the tree to do this. Once bees began feeding on the sucrose solution, we gradually moved the cotton away from the nest entrance and to the final location, ensuring bees followed each move. At the training location, we transferred bees from cotton to feeders. Each feeder consisted of a small inverted jar on a grooved plastic plate (von Frisch 1967). This design provides a constant supply of sucrose solution. Feeders were supported on plastic horizontal surfaces attached to the top of tripods approximately 1 m high. Before each trial, we let bees recruit to their training feeder until there were sufficient foragers (typically 50–200) for meaningful data collection (see below).

**Recruitment pheromone chemical analysis**

We determined the chemical composition of *T. hyalinata* recruitment pheromone via gas chromatography–mass spectrometry (GC-MS). We dissected labial glands from five foragers from the nest used in 2009 under a stereo microscope, carefully separating the glands from all other tissues. All glands were combined in 300 μL pure hexane (Labsynth) and dissolved at room temperature for 24 h before being stored in a freezer. To prevent contamination from alarm pheromone or other substances on the bees’ cuticles, we rinsed bees five times in hexane before dissection.

We carried out the GC-MS with a Shimadzu QP-2010 GCMS system. The GC was equipped with two different columns: first a DB-5MS (30 m×0.25 mm, J&W Scientific, Folsom, CA, USA) and then a more polar DB-WAX column (30 m×0.25 mm, J&W Scientific) for better separation of similar-weight compounds. Helium was used as a carrier gas (constant linear flow rate 40.0 cm/s with the DB-5MS column, 39.7 cm/s with the DB-WAX column). With the DB-5MS column, temperature was held at 50°C for 5 min, then increased by 5°C/min to 300°C, where it was held for the final 5 min. With the DB-WAX column, temperature started at 50°C, then increased by 3°C/min to 240°C, where it was held for the final 5 min. We made preliminary compound identifications using the Wiley mass spectral library (Mclafferty 2000), then confirmed all identifications with synthetic standards. Analytical standards’ sources were: Sigma-Aldrich Corp. (St. Louis, MO, USA), hexyl octanoate and octyl octanoate; CTC Organics (Atlanta, GA, USA), octyl decanoate; Jocelyn Millar (Riverside, CA, USA), octyl hexanoate; Stefan Jarau (Ulm, Germany), decyl hexanoate and hexyl decanoate.
Preference tests

We assessed bees’ responses to con- and heterospecific recruitment pheromones via preference tests, where individuals chose between two feeders each bearing different odors. Before the start of a trial we covered the training feeder, which encouraged bees to search for alternate food sources. Thus, participating bees functioned as scouts, individuals using internal information (knowledge of food types, sensory input, etc.) to locate food sources previously unknown to them (Biesmeijer and de Vries 2001). Preference tests were conducted 20–25 m from the nest and 15–20 m from the training feeder (see Fig. 1). Because bees scout independently and the potential search area increases with distance from the training feeder, new feeders farther from the training feeder will be found more slowly. Thus, we chose distances at which bees would arrive singly, yet frequently enough to gather meaningful data (at least ten decisions) during a trial. We chose a 15-min trial duration, within the 20-min retention time found for T. spinipes recruitment pheromone (Nieh et al. 2004b). Training feeders were set out when bees became active in the morning (c. 0900 hours) and trials continued through mid-day. Because the effects of low humidity on recruitment pheromone and bee scouting behavior are unknown, we conducted trials only when the relative humidity was above 50%. Only one colony was tested each day, participating in one to four trials of the various types described below. Solvent control and the treatment trials were interspersed.

At the testing location, we set up two feeders with 40 cm between their centers. This distance was short enough that arriving bees could smell both feeders and allowed simultaneous observation by a single observer. Recruitment pheromones have an active space of approximately 20 m under calm conditions (D. Schorkopf, personal communication). Preliminary trials showed 40 cm was far enough that bees would distinguish between the two feeders and treat them as separate odor sources. Bees arriving at the test location made a choice by landing on one of the two testing feeders. To avoid any potentially confounding effect of food presence, these feeders were empty. We immediately removed bees once they made a choice to eliminate visual local enhancement: orientation of foragers to the visual presence of other bees (Slaa et al. 2003). These bees were marked with enamel paint, released, and not counted in subsequent trials to avoid pseudoreplication.

Pheromones

Recruitment pheromone sources were (1) fresh pheromone and (2) labial gland extract (LGE). To collect fresh pheromone, we trained one forager of the marking species to visit a feeder with 30% or 45% (1.5 M) sucrose solution (using the higher concentration when recent recruitment levels were low) approximately 10–20 m further from the nest than the training feeder. This feeder bore strips of paper upon which bees readily deposited pheromone by rubbing their mouthparts on the paper’s edge while running along it. An odor mark was defined as one such rubbing event. Once the trained bee was observed odor marking and a group of recruits arrived at the feeder (indicating strong recruitment, and corresponding to the “pulses” in Nieh et al. 2004b), we harvested pheromone. We cut off the three to five strips of paper upon which bees had spent the most time odor marking, yielding seven to ten recent marks. The papers were then quickly transported (approximately 2 min in 2008, 10 min in 2009) to the other species’ testing location. In 2009, when the two nests were farther apart, we stored the marked papers in a clean glass vial in a cooled container during transport to slow volatilization. These strips were placed in a slit on the platform supporting one of the testing feeders. Clean paper was placed in a slit on the other feeder.

Use of LGE facilitates the application of precise, reliable pheromone doses and yields the same orientation behaviors as naturally deposited pheromone in the three Trigona species previously tested (Jarau 2009). We prepared LGE in a manner similar to the sample prepared for chemical analysis, except that each bee’s glands were separately dissolved in 100 μL of hexane. For LGE trials, we attached a small strip of filter paper (20 × 5 mm) to each testing feeder. In solvent control trials testing the effect of hexane, one randomly selected strip remained dry while the other bore 10 μL of hexane. For the treatments described below, we applied 10 μL of LGE to one of the filter paper strips. Ten microliters equaled 0.1 bee equivalents, a concentration to which closely related species respond (Jarau et al. 2004). One bee equivalent is the full labial gland content from one bee. In intraspecific preference tests, we used LGE harvested from the colony being tested. Each replicate trial used LGE from a different individual. We verified that bees’ responses to LGE matched behavior exhibited

![Fig. 1 General layout of feeders during preference tests. Layout and exact locations used depended on topography, avoiding dense vegetation, steep slopes, roads, buildings, etc. Distances were: nest to training feeder 10–15 m, training feeder to testing feeders 15–20 m, nest to testing feeders 20–25 m, between the center of each testing feeder 40 cm](image-url)
towards fresh pheromone. All equipment was cleaned with lab detergent or ethanol between trials.

Recruitment pheromone glandular source—intraspecific preference tests

*T. spinipes* produces recruitment pheromone in the cephalic labial glands and will follow an artificial trail made from labial gland extract (Schorkopf et al. 2007). In intraspecific preference tests, we determined whether *T. hyalinata* cephalic labial gland secretions are attractive to *T. hyalinata*. We compared bees’ responses to LGE from nestmates to responses in solvent control trials, repeating treatment and control five times per colony. We also tested *T. spinipes* in order to have a common methodology for comparing bees’ responses to nestmate recruitment pheromone with the eavesdropping responses described below. The two colonies of each species used in the eavesdropping experiment were also used for this one.

Interspecific eavesdropping

We used preference tests to determine heterospecific eavesdropping responses between *T. hyalinata* and *T. spinipes* under two conditions: (1) In the one-pheromone treatment, bees chose between a dry strip of paper and paper with 10 μL of heterospecific LGE (five trials per colony) or between a dry strip of paper and freshly deposited heterospecific pheromone (two to three trials per colony). To test the generality of eavesdropping responses, we conducted two additional one-pheromone trials per colony using LGE from bees captured from a different location (approximately 500–1,000 m away). These latter bees were likely from a different colony, and may or may not have been encountered by test colony foragers before experimentation. (2) In the two-pheromone treatment, bees chose between nestmate LGE and heterospecific LGE (five trials per colony).

Data analysis

For each trial, we determined the proportion of bees landing on the feeder with hexane (solvent control trials), the feeder with nestmate LGE (intraspecific preference tests) or the feeder with heterospecific LGE (eavesdropping trials). All proportions were transformed using Anscombe’s arcsine transformation (Zar 1999) to meet parametric assumptions. We assessed responses to nestmate pheromone and eavesdropping with two-factor ANOVAs, including colony as a fixed effect. For each species, we conducted five separate analyses, corresponding to separate questions (see Table 1). (A1) Did the average proportion of bees preferring the feeder with nestmate LGE in intraspecific preference tests differ from the proportion that preferred the hexane feeder in solvent control trials? This tested whether the labial glands are a source of recruitment pheromone. (A2) In the one-pheromone treatment, did bees show the same eavesdropping response to heterospecific LGE and fresh heterospecific pheromone? Due to the small number of trials with fresh pheromone, we used the non-parametric Mann–Whitney *U* test and untransformed proportions for this analysis. (A3) Did bees respond the same to heterospecific LGE from a nearby and a more distant heterospecific colony? This analysis employed Kruskal–Wallis tests and untransformed proportions due to the small number of trials with the distant colonies. (A4) Was there a difference in the average proportion of bees preferring hexane in solvent control trials, heterospecific LGE in the one-pheromone eavesdropping treatment or heterospecific LGE in the two-pheromone treatment? This tested heterospecific eavesdropping responses and was followed by a Tukey HSD test to determine pairwise significant differences

| Question | Treatments compared |
|----------|---------------------|
| (A1) Recruitment pheromone glandular source | Solvent control | Nestmate LGE |
| (A2) Eavesdropping on heterospecific labial gland extract (LGE) vs. natural odor marks | One-pheromone treatment | Fresh odor marks |
| (A3) Response to labial gland extracts from different heterospecific colonies | One-pheromone treatment, nearby colony | One-pheromone treatment, distant colony |
| (A4) Eavesdropping on heterospecific recruitment pheromone | Solvent control | One-pheromone treatment | Two-pheromone treatment |
| (A5) Response to own recruitment pheromone in the presence or absence of heterospecific pheromone | Nestmate LGE | Two-pheromone treatment (proportion in opposite direction of A4 analysis) |

See text for treatment descriptions
Results

Recruitment pheromone chemical analysis

*T. hyalinata* labial gland extract contained six major components, all carboxylic esters (Fig. 2, Table 3). Octyl octanoate, the major component of *T. spinipes* recruitment pheromone (Schorkopf et al. 2007), was the most abundant component of *T. hyalinata* recruitment pheromone.

Recruitment pheromone glandular source—intraspecific preference tests (question A1)

*T. hyalinata* LGE was highly attractive to conspecifics, strongly suggesting it is the source gland of recruitment pheromone for this species. Compared to their responses to hexane in solvent control trials, foragers of each species chose a feeder with nestmate LGE significantly more often than a paired feeder with no odor (A1; Fig. 3; Table 4). Preferences did not differ between conspecific colonies (Table 4). The *T. spinipes* nest we used in 2008 fell from its host tree and died before we completed our experiment, so we were only able to conduct three solvent control trials with this colony.

Table 2 Sample sizes and bee participation for each of the treatments, separated by species

| Treatment                  | Number of trials conducted | Mean number of bees choosing a feeder (min, max) |
|----------------------------|----------------------------|--------------------------------------------------|
| *Trigona hyalinata*        |                            |                                                  |
| Solvent control            | 10                         | 20.6 (10, 38)                                    |
| Nestmate LGE               | 11                         | 21.2 (13, 28)                                    |
| One-pheromone treatment    |                            |                                                  |
| Nearby colony              | 9                          | 18.4 (11, 26)                                    |
| Distant colony             | 4                          | 15.3 (12, 18)                                    |
| Fresh odor marks           | 4                          | 14.8 (5, 24)                                     |
| Two-pheromone treatment    | 11                         | 18.4 (11, 32)                                    |
| *Trigona spinipes*         |                            |                                                  |
| Solvent control            | 8                          | 20.6 (10, 49)                                    |
| Nestmate LGE               | 10                         | 19.7 (11, 34)                                    |
| One-pheromone treatment    |                            |                                                  |
| Nearby colony              | 10                         | 17.2 (10, 29)                                    |
| Distant colony             | 5                          | 14.4 (10, 19)                                    |
| Fresh odor marks           | 6                          | 17.7 (12, 30)                                    |
| Two-pheromone treatment    | 10                         | 17.5 (12, 25)                                    |

The fresh odor marks trial with only five bees was included because death of the marking colony prevented us from repeating that trial.

Interspecific eavesdropping (questions A2–A4)

For both species, response to heterospecific recruitment pheromone did not vary with pheromone source (A2, fresh pheromone vs. LGE: *T. hyalinata*, 29% vs. 34%, *U* = 34, *p* = 0.39; *T. spinipes*, 52% vs. 51%, *U* = 39, *p* = 0.67), the identity of the colony donating the LGE (A3, nearby vs. far: *T. hyalinata*, 37% vs. 29%, *K* = 3.84, *p* = 0.028; *T. spinipes*, 51% vs. 53%, *K* = 0.62, *p* = 0.89), or the colony being tested (ANOVA results in Table 4). Compared to solvent control trials, *T. hyalinata* strongly avoided *T. spinipes* recruitment pheromone (A4; Fig. 4; Table 4) in both one- and two-pheromone treatments. When choosing between conspecific and heterospecific LGE, the same proportion of *T. hyalinata* foragers preferred their own pheromone as in intraspecific preference tests (A5; Fig. 5; Table 4). *Trigona spinipes* foragers’ response to *T. hyalinata* pheromone was the same as their response to hexane (A4; Fig. 4; Table 4). Although *T. spinipes* foragers appeared to choose the feeder with *T. hyalinata* LGE less frequently in the two-pheromone treatment than in the one-pheromone treatment, this difference was not statistically significant (*p* = 0.06). However, the presence of *T. hyalinata* pheromone significantly reduced *T. spinipes* preference for their own LGE (A5; Fig. 5; Table 4).

Discussion

Recent evidence suggests that social information use by foragers is widespread, but our understanding of how animals use such information remains limited. Most examples of interceptive eavesdropping and “spying” occur between trophic levels or emphasize copying behavior (reviewed in Dall et al. 2005; Peake 2005; Seppänen et al. 2007; Valone 2007; Goodale et al. 2010). Our results provide the most complete example to date of interceptive eavesdropping by competing social insects: we determined the composition and probable glandular source of the chemical signal, and examined reciprocal heterospecific responses to this signal in preference tests with multiple colonies. We show heterospecific avoidance eavesdropping by a stingless bee: *T. hyalinata* avoids the recruitment
pheromone of *T. spinipes*. Among pollinating social bees, this within-trophic-level social information can help foragers avoid unprofitable resources or conflict (e.g., Stout et al. 1998; Nieh et al. 2004a).

Chemical analysis of *T. hyalinata* and *T. spinipes* LGE demonstrates that the pheromones should be (1) detectable by both species because both contain octyl octanoate in relatively high concentrations and (2) differentiable because *T. spinipes* recruitment pheromone consists of one major component while *T. hyalinata* has six (Fig. 2). One of these, hexyl octanoate, is reported for the first time as a component of stingless bee recruitment pheromones. Unlike the pattern reported for other social insects (Slaa and Hughes 2009), eavesdropping responses did not depend on relative dominance of the eavesdropping and signaling species. In our study, the dominant species, *T. hyalinata*, avoided the recruitment pheromone of the subordinate species. *Trigona spinipes* showed no attraction to or avoidance of the dominant species’ pheromone. Under the dominance motivation hypothesis, if eavesdropping decisions were based solely on relative dominance, we would have seen attraction by *T. hyalinata* foragers and avoidance by *T. spinipes* foragers to heterospecific recruitment pheromone.

Recruitment pheromones

Four trail-laying stingless bee species (in three genera), in addition to our study species, are attracted to labial gland secretions (Jarau 2009; Stangler et al. 2009). These results, taken with the identical chemistry of *T. spinipes* LGE and odor marks (Schorkopf et al. 2007), indicate that stingless bees’ recruitment pheromones are secreted by the labial glands. Our finding that *T. hyalinata* foragers are strongly attracted to nestmate LGE, and chemical similarity with congener’s LGEs, strongly suggests that recruitment pheromone comes from the labial glands in this species as well.

*Trigona hyalinata* recruitment pheromone composition is consistent with recruitment pheromones of congeners (Jarau 2009) and other odor-marking stingless bees (Stangler et al. 2009), which contain carboxylic and terpene esters. Octyl hexanoate, octyl octanoate, hexyl decanoate, octyl decanoate, and decyl hexanoate are shared with other species (Jarau 2009; Jarau et al. 2010), while hexyl octanoate is reported for the first time as components of stingless bee recruitment pheromones. Behavior of other *Trigona* species suggests that foragers require the entire blend of chemicals to exhibit natural trail-following behavior (Jarau 2009). Interestingly, the esters found in *T. hyalinata* recruitment pheromone are also found in other glandular extracts thought to have an attractive function. These include the Dufour’s gland in *Andrena* (Fernandes et al. 1981; Hefetz 1987), *Dufourea* (Wheeler et al. 1985), and *Svastra* (Duffield et al. 1984) bee species, and mandibular and preputial glands that likely produce sex pheromones in several *Myrmecocystus* ant species (Lloyd et al. 1989) and the Brandt’s vole (Zhang et al. 2007), respectively.

| Peak | Compound         | Retention time (min) | Relative abundance (%) |
|------|------------------|----------------------|------------------------|
| A    | hexyl octanoate  | 29.374               | 13.06                  |
| B    | octyl hexanoate  | 29.425               | 1.92                   |
| C    | octyl octanoate  | 36.180               | 45.46                  |
| D    | hexyl decanoate  | 36.249               | 12.64                  |
| E    | decyl hexanoate  | 36.329               | 13.33                  |
| F    | octyl decanoate  | 42.460               | 13.59                  |
In intraspecific preference tests, *T. spinipes* showed weaker preference (65%) than did *T. hyalinata* (80%). *Trigona spinipes* preferences were also weaker than in other preference experiments (73%, Nieh et al. 2004b; 90%, Schorkopf et al. 2007), while *T. hyalinata* shows greater consistency across studies (81%, Nieh et al. 2003). Three major differences in life history traits between these species could be related to this, although the last two seem less likely. First, *T. spinipes* appear to be highly generalist in their floral visitation, visiting 51% of 562 plant species for which we have collated stingless bee resource use data. *Trigona hyalinata*, however, were found on only 5% of the plants and seem to specialize on dense floral patches: trees and shrubs (unpublished data). The relationship between floral preference patterns and reliance on social information is not clear and bears further investigation. Second, at the species level *T. hyalinata* are more dominant and aggressive than *T. spinipes* (Lichtenberg et al. 2010). Third, *T. hyalinata* colonies have almost three times the number of workers as *T. spinipes* colonies have (D.W. Roubik,

![Fig. 3](image1.png) Responses of *T. hyalinata* and *T. spinipes* foragers to labial gland extract (LGE) from nestmates in intraspecific preference tests, compared to responses to hexane in solvent control trials (question A1). Bars show the average proportion of foragers preferring the given odor (±SE). Asterisks indicate statistically significant differences (α=0.05)

![Fig. 4](image2.png) Interspecific eavesdropping analysis (question A4), comparing preferences for hexane in solvent control trials with heterospecific LGE in the absence (“One-pheromone treatment”) and presence (“Two-pheromone treatment”) of nestmate LGE. Bars show the average proportion of foragers preferring the given odor (±SE). Asterisks indicate statistically significant differences (α=0.05)

### Table 4 ANOVA results for recruitment pheromone glandular source and heterospecific eavesdropping questions asked in the “Data analysis” subsection of the “Materials and methods” section, separated by responding species

| Research question | ANOVA output | Effect | F | p     | df  |
|-------------------|--------------|--------|---|-------|-----|
|                   |              |        |   |       |     |
| **Trigona hyalinata** |              |        |   |       |     |
| (A1) Recruitment pheromone glandular source |              |        |   |       |     |
| Trial type        | 67.28        | <0.0001| 1, 18 |
| Colony            | 1.03         | 0.32   | 1, 18 |
| (A4) Eavesdropping on heterospecific recruitment pheromone |              |        |   |       |     |
| Trial type        | 20.01        | <0.0001| 2, 30 |
| Colony            | 1.66         | 0.21   | 1, 30 |
| (A5) Response to own recruitment pheromone in the presence or absence of heterospecific pheromone |              |        |   |       |     |
| Trial type        | 0.003        | 0.96   | 1, 19 |
| Colony            | 2.62         | 0.12   | 1, 19 |
| **Trigona spinipes** |              |        |   |       |     |
| (A1) Recruitment pheromone glandular source |              |        |   |       |     |
| Trial type        | 37.23        | <0.0001| 1, 15 |
| Colony            | 1.54         | 0.23   | 1, 15 |
| (A4) Eavesdropping on heterospecific recruitment pheromone |              |        |   |       |     |
| Trial type        | 3.19         | 0.06   | 2, 29 |
| Colony            | 0.08         | 0.78   | 1, 29 |
| (A5) Response to own recruitment pheromone in the presence or absence of heterospecific pheromone |              |        |   |       |     |
| Trial type        | 12.57        | 0.002  | 1, 17 |
| Colony            | 0.004        | 0.95   | 1, 17 |

All interactions were not significant, and were removed from analyses. All analyses in this table used LGE. Statistics for questions A2 and A3 are given in the text.
personal communication; Wille and Michener 1973). This is likely related to the dominant status of *T. hyalinata*; species with larger colonies tend to be more dominant (Lichtenberg et al. 2010). It is unlikely that species’ dominance affects degree of reliance on information provided by nestmates, since subordinate *Melipona* species show strong attraction to nestmates’ chemical “footprints” (e.g., 91%; Nieh 1998).

Most previous research on stingless bee recruitment pheromones utilized artificial trails rather than presenting odors at a food source (the feeder), as we did. This raises the possibility that pheromones deposited at the food source differ from those used in odor trails. However, no evidence indicates that stingless bees have two separate recruitment pheromones. Our protocol used the same pheromone-producing gland as artificial trail studies. In one case, chemically analyzed odor marks were collected from the feeder (Schorkopf et al. 2007). In addition, *T. spinipes* and *T. hyalinata* create polarized odor trails, depositing the majority of their pheromone within 1 m of the feeder (Nieh et al. 2004b). Because the trail is an extension of odor marks at the food source, it is reasonable to assume that pheromone deposited on the feeder is the same as trail pheromone, and that the bees obtain the same meaning from our experimental setup and odor trails.

The pheromone concentration that we used, 0.1 bee equivalents, elicited a natural response in a congener, *Trigona recurva* (Jarau et al. 2004), and in our experiments. Eavesdropping responses of *T. hyalinata* and *T. spinipes* were the same whether the treatment was fresh pheromone that had elicited strong natural recruitment or LGE (*T. hyalinata*, 29% vs. 34% of bees choosing the feeder with pheromone; *T. spinipes*, 52% vs. 51%). Each species also showed a highly significant (*p*<0.001) preference for nestmate recruitment pheromone at 0.1 bee equivalents (Fig. 3; *T. hyalinata*, 80% choosing the feeder with pheromone; *T. spinipes*, 65%). The strength of these preferences was similar to those shown by each species in preference tests that employed odor trails (Nieh et al. 2003; Nieh et al. 2004b). Greater or lesser amounts of LGE may elicit different eavesdropping responses than those reported here. Predatory ants eavesdropping on fig volatiles exhibit such a dose-dependent response, showing greater attraction to larger quantities of figs (Ranganathan and Borges 2009). However, our results show that 0.1 bee equivalents are sufficient to elicit both conspecific and heterospecific responses that are the same as responses to natural-deposited pheromone at approximately the same concentration.

### Interspecific Eavesdropping

The limited amount of natural habitat near the laboratory and *Trigona* preferences for nesting high in trees limited the number of colonies that we were able to work with. Despite this, we feel that our results reflect species-typical behaviors. Results were highly consistent across replicate colonies (Table 4), and each colony showed the same response to pheromone from two different heterospecific colonies (A3). This consistency across colonies and months also indicates that a species’ eavesdropping behavior does not vary much, if at all, with the current food needs of the colony.

*Trigona spinipes* foragers clearly could detect the presence of heterospecific pheromone. Preference for nestmate pheromone decreased significantly, albeit slightly (Fig. 5; going from 65% to 59% choosing the *T. spinipes* LGE; *p*=0.002), when bees chose between conspecific and heterospecific pheromones. Thus, *T. spinipes* does recognize *T. hyalinata* pheromone as different. Despite this detection ability, *T. spinipes* foragers exhibited a behavioral lack of choosiness in eavesdropping tests, showing no preference between feeders with no odor and *T. hyalinata* pheromone (Fig. 4). A similar failure to use social information has been found for three-spined sticklebacks, although the related nine-spined sticklebacks use similar social information (Cooil et al. 2003). *Trigona spinipes* are attracted to footprint cues of the subordinate *Melipona rufiventris* at certain locations (Nieh et al. 2004a), suggesting they have a species- and context-specific response to social information, and heterospecific signals and cues do not always alter their movements. Bumble bees also facultatively use social information, exhibiting visual local enhancement only when approaching unfamiliar flower types (Kawaguchi et al. 2007). Alternately, *T. spinipes* may ignore *T. hyalinata* pheromones when they cannot also see foragers on the marked food source. *Apis mellifera* ignore olfactory information when sufficient visual information is available (Gürfa et al. 1994).

Contriary to our expectation, *T. hyalinata* foragers showed strong avoidance of the subordinate species’ recruitment pheromone (Fig. 4). This result was surprising,
given previous patterns of social information use by social insects (Slaa and Hughes 2009) and the highly dominant behavior exhibited by *T. hyalinata* (Lichtenberg et al. 2010). It is unlikely that our results reflect avoidance of all non-nestmate recruitment pheromones by *T. hyalinata* foragers. Individual *T. hyalinata* foragers will fly to and attack other species at food sources (Lichtenberg et al. 2010). Given that dominant stingless bee species such as *T. hyalinata* appear to be relatively poor at discovering new food sources (the dominance-discovery trade-off, Fellers 1987; Nagamitsu and Inoue 1997), avoiding all non-nestmate odors would severely limit food intake by *T. hyalinata* colonies.

Our findings are consistent with a hypothesis of conflict avoidance through eavesdropping decisions. Attacking to gain control of an occupied resource can inflict mortality losses even for highly dominant species (Johnson and Hubbell 1974; Nieh et al. 2005). The recruitment pheromone we presented to eavesdropping *T. hyalinata* in trials with fresh odor marks and LGE corresponds to the presence of numerous subordinate foragers. In our fresh odor mark trials, we collected marks once a “pulse” (Nieh et al. 2004b) of at least 30 bees arrived at the feeder. Stingless bee trails must be actively maintained by bees interrupting their food collection, and begin to fade out after approximately 20 min without such maintenance. Thus, recruitment pheromones provide current information on both resource availability and abundance of bees already present at the resource. Under such conditions, attack may be costly for a *T. hyalinata* colony because it may require the participation of hundreds of bees, which could otherwise be recruited to non-contested food sources. *Trigona hyalinata* foragers’ decisions to not choose a resource at which social information predicts high numbers of subordinate heterospecifics may be similar to the failure of dominant *Trigona silvestriana* to drive away large numbers of subordinate bees (Johnson and Hubbell 1974). Indeed, while *T. hyalinata* can easily displace a group of foraging *T. spinipes*, they do not attempt to do so every time they encounter the subordinate species (personal observation). This behavior merits further study. One possible explanation is that social insect eavesdropping decisions include expected costs associated with each choice; research that we are currently conducting investigates this.

Different lines of evidence suggest that eavesdropping on signals and spying on cues affect the movements of social bees. First, we have observed *T. hyalinata* depositing odor marks on flowers. Second, feeders are discovered more quickly by other stingless bee species when they bear recruitment pheromone or a large quantity of footprints (Johnson 1983). Finally, interspecific interactions increase between-plant movement of honey bees (Greenleaf and Kremen 2006) and bumble bees (Kawaguchi et al. 2007), and may do the same for eavesdropping stingless bees. Our results indicate that social information use by competitors is governed by complex rules. Potentially large ecological and evolutionary impacts make this an important area for future investigation.

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**Conflict of interest** The authors declare that they have no conflict of interest.

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