Foragers of sympatric Asian honey bee species intercept competitor signals by avoiding benzyl acetate from *Apis cerana* alarm pheromone

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While foraging, animals can form inter- and intraspecific social signalling networks to avoid similar predators. We report here that foragers of different native Asian honey bee species can detect and use a specialized alarm pheromone component, benzyl acetate (BA), to avoid danger. We analysed the volatile alarm pheromone produced by attacked workers of the most abundant native Asian honey bee, *Apis cerana* and tested the responses of other bee species to these alarm signals. As compared to nest guards, *A. cerana* foragers produced 3.38 fold higher levels of BA. In foragers, BA and (E)-dec-2-en-1-yl acetate (DA) generated the strongest antennal electrophysiological responses. BA was also the only compound that alerted flying foragers and inhibited *A. cerana* foraging. BA thereby decreased *A. cerana* foraging for risky sites. Interestingly, although BA occurs only in trace amounts and is nearly absent in sympatric honeybee species (respectively only 0.07% and 0.44% as much in *A. dorsata* and *A. florea*), these floral generalists detected and avoided BA as strongly as they did to their own alarm pheromone on natural inflorescences. These results demonstrate that competing pollinators can take advantage of alarm signal information provided by other species.

Alarm signalling, defined as one organism using signals to alert another about danger, is widespread and occurs in plants1, insects2, and vertebrates3, 4. Species at the same trophic level can transfer interspecific information about foraging and risk avoidance5. By using common information that reliably indicates predation, prey may benefit from shared information6, 7. Such information sharing has been demonstrated in tadpoles6, 8, fishes9, and social insects2. This transfer may be beneficial even when the species are competitors. For example, Asian honey bee foragers from different colonies and species can be rivals for limited nectar and pollen resources10. Can they use interspecific alarm pheromones for their individual benefit? Although alarm signalling may be individually costly, it can evolve via kin selection11 and reciprocal altruism12, as exemplified by alarm signals in eusocial organisms. Once such signals have evolved, different colonies of the same species and even different species could benefit by intercepting information about dangerous food locations. Theoretically, they could intercept alarm signal information13. As predicted, *A. cerana* can use olfactory eavesdropping to detect and avoid an alarm pheromone component in the sympatric *Apis dorsata* that *A. cerana* does not possess14.

Such interception within a species is a by-product of colonies having the same alarm pheromone. However, between species with different alarm pheromone compounds or ratios of these compounds, evolution could favour heterospecific sensitivity to alarm pheromones. Species that often encounter each other, like members of a pollinator guild, could benefit by learning to recognize alarm pheromones produced by heterospecifcics. This is similar to the phenomenon of bees avoiding heterospecific “footprint” cuticular hydrocarbon odours that indicate a specific flower has already been visited and therefore is less likely to be rewarding16. We hypothesize that different honey bees can learn to associate different honey bee alarm pheromones with danger and thereby reduce their

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risk of predation during foraging. Such avoidance has implications for pollination, because predators can impose significant non-consumptive effects by causing pollinators to avoid dangerous locations.\(^1\)

In honey bees, alarm pheromones can increase colony fitness by reducing colony recruitment to dangerous locations. For example, \(L.\) mellifera foragers that detected sting alarm pheromone at a food source significantly reduced their recruitment (less waggle dancing) and increased their production of inhibitory stop signals.\(^18\) Thus, alarm pheromones can also inhibit recruitment communication, providing an olfactory negative feedback signal against the positive feedback signal of the waggle dance.

Honey bee sting alarm pheromones are multi-component blends. Isopentyl acetate (IPA) is the major component of sting alarm pheromone in all honey bee species.\(^19\) The main other previously-reported sting alarm pheromone components (>10% by mass) of each species are benzyl acetate (BA, in \(L.\) mellifera),\(^20\) octyl acetate (OA, in \(L.\) mellifera, \(L.\) cerana, \(L.\) florea and \(L.\) dorsata),\(^21\) (E)-oct-2-en-1-yl acetate (OEHA, in \(L.\) mellifera),\(^22\) (E)-dec-2-en-1-yl acetate (DA, in \(L.\) cerana, \(L.\) dorsata, \(L.\) laboriosa and \(L.\) florea),\(^15, 20, 21, 23\) (Z)-eicos-11-en-1-ol (EH, in \(L.\) mellifera and \(L.\) cerana),\(^14, 23\) and gamma-octanoic lactone (GOL, in \(L.\) dorsata and \(L.\) laboriosa).\(^26, 27\)

The functions of these different compounds are best understood in the Western honey bee, \(L.\) mellifera. IPA, OA and BA play a major role in \(L.\) mellifera nest defence. IPA is most important for initiating an alarm response, but is so volatile that it is less effective at marking the intruder for further attacks. OA is less volatile, and therefore more persistent: it is important for orienting bees towards a moving target.\(^25\) In \(L.\) mellifera, BA is more effective at increasing the number of fanning workers in the hive, which may be part of a defensive response.\(^28, 29\) In \(L.\) mellifera workers, BA levels also depend upon task specialization.\(^30\) However, the function of BA is otherwise unclear.

Less is known about the effects of different alarm pheromone components in other honey bee species. IPA is the major alarm pheromone in \(L.\) mellifera, \(L.\) cerana, \(L.\) dorsata, and \(L.\) florea, but natural sting pheromone elicits a longer-lasting reaction than IPA alone.\(^21\) The more persistent DA may provide an orientation cue in \(L.\) dorsata and \(L.\) florea, as OA does in \(L.\) mellifera.\(^11, 23\) In \(L.\) cerana, EH is also more persistent than IPA and may provide orientation information.\(^24\)

In addition, components may exert different effects depending upon context. In the context of foraging, bees are not defending their colony but rather fleeing from danger and marking a location as dangerous.\(^26\) For example, GOL and DA are most effective at repelling \(L.\) dorsata and \(L.\) cerana foragers, even though GOL is not found in \(L.\) cerana. This example of \(L.\) cerana intercepting an alarm pheromone component of another bee species illustrates the complexity of forager responses to alarm pheromones.\(^15\)

\(L.\) cerana, \(L.\) dorsata, and \(L.\) florea are sympatric tropical Asian honey bee species,\(^11, 15\) face formidable predators at the nest and in the field,\(^15, 19, 31–33\), and are major native pollinators of agricultural crops and native plants in Asia.\(^34–37\) The different species vary in population density, with \(L.\) cerana as the most common (in order of abundance: \(L.\) cerana \(>\) \(L.\) dorsata \(>\) \(L.\) florea).\(^38\) In fact, \(L.\) cerana is more than three times more abundant than \(L.\) florea. The abundance of \(L.\) dorsata changes seasonally due to their annual migrations. In seasons when \(L.\) dorsata is sympatric with \(L.\) cerana, \(L.\) cerana is initially more abundant than \(L.\) dorsata, but \(L.\) dorsata eventually becomes as or more abundant than \(L.\) cerana.\(^4\) Thus, it should be advantageous for \(L.\) dorsata and \(L.\) florea to detect and intercept the alarm pheromone of \(L.\) cerana, the most abundant bee species.

Some of these honey bee species have alarm pheromone compounds, like GOL, that are not found in other honey bee species.\(^15\) However, the primary interspecific differences lie in the relative abundances of these different compounds. Because the relative abundances may be a source of information, it is possible that \(L.\) florea and \(L.\) dorsata do not respond or respond differently to \(L.\) cerana alarm pheromone. Our goal was therefore to better understand the function of different honey bee alarm pheromone components in \(L.\) cerana, to determine if BA varies according to the sympatric species, \(L.\) cerana and \(L.\) florea, can intercept and use this information.

### Materials and Methods

**Honey bee colonies and sample collection.** We used six \(L.\) cerana \(L.\) cerana colonies (three four-comb colonies and three two-comb observation colonies) at Yunnan Agriculture University and Kunming Botanic Garden in Kunming for the pheromone sampling and feeder experiments. At the Xishuangbanna Tropical Botanical Garden (XTBG), China, approximately 20 \(L.\) cerana colonies were sited close to where we conducted our inflorescence bioassay experiment. At XTBG, we used naturally foraging bees but conducted trials over several months (October 2015 to April 2016) over a broad area (56 plants distributed over 4 km²) and therefore likely used bees from multiple colonies. Sample sizes for each experiment are shown in Table S1.

Wild bee species were collected at XTBG and Nabanhe (Yunnan). \(L.\) dorsata and \(L.\) florea foragers were from both XTBG (two colonies per species) and Nabanhe (one colony per species). In field bioassays, we likely used more than three wild colonies of \(L.\) dorsata because we collected foragers at three different sites, each separated by at least 6.4 km.

**Exp. 1: Alarm pheromone analyses.** The headspace odours produced by alarmed \(L.\) dorsata have been previously determined.\(^15, 26\) To analyse and determine the source of \(L.\) cerana and \(L.\) florea alarm pheromone, we collected headspace odours emitted by an alarmed forager using the same procedure used for \(L.\) dorsata alarm pheromone analyses.\(^15, 26\) Our primary focus was \(L.\) cerana, but for our information interception experiments, we wished to determine (for \(L.\) florea) and confirm (for \(L.\) dorsata), which alarm pheromone components these species possess, using the same procedures with the same Gas Chromatography (GC) and Gas Chromatography-Mass Spectrometry (GC-MS) equipment.

We first carefully captured bees foraging on inflorescences without alarming them. To do so, we gently placed the wider opening of a clean glass funnel (2.0 cm and 0.8 cm diameter openings) over a foraging bee. The captured bee was then induced to walk through the funnel and into a clean 2 ml glass vial, attracted by 365 nm UV LED...
light (CREE, TW). This vial was sealed with a PTFE cap through which we inserted a needle (0.2 mm in diameter, 5.0 cm long) to disturb the bee into producing alarm pheromone. This needle made contact with, but did not pierce, the bee cuticle. We then removed this needle and collected all volatile alarm pheromones by inserting a 65 μm PDMS/DVB fibre (Supelco, CA) into the vial for Headspace Solid Phase Micro-Extraction (HS-SPME) at 30 °C for 30 min. As a control, in preliminary trials, we captured bees inside glass vials and did not disturb them with a needle. However, even in this situation, we were able to detect trace amounts of known alarm pheromone compounds, such as IPA because the bees were trapped for 30 min. We therefore collected control data on odours pumped from the headspace around undisturbed foragers in a hive and passed through a clean glass tube in which the SPME fibre was placed for 1 hour per trial. We conducted nine trials with three different colonies and found no compounds that matched those produced by needle-alarmed bees.

We also analysed the volatile compounds emitted by sting venom alone. We dissected a sting (stinger and sting gland) from a cold-anesthetized forager or guard and deposited it into a PTFE capped vial (1 sting per vial). After the dissected sting apparatus revolved at room temperature, it began to pulse and emit drops of venom. We then extracted the headspace odours using the HS-SPME procedure previously described. For each species, we compared 18 extracts (9 alarmed bees and 9 sting glands from bees from three different A. cerana colonies). For A. dorsata and A. florea, we were not able to collect directly from colonies and therefore collected bees foraging on floral resources at spaced at least 4 km apart.

Our primary focus was on A. cerana. To determine if A. cerana workers produced different alarm pheromones depending upon their task specialization, we used GC-FID quantification to compare guard bees and forager bees from colonies in Kunming. To collect guard bees, we struck the nest and captured exiting guards with a clean soft cotton sieve. To collect foragers, we captured bees returning to the nest with pollen in their corbiculae with a clean cotton sieve. Such bees are pollen foragers and therefore clearly identifiable as foragers.

For our coupled Gas Chromatography-Electroantennographic Detection (GC-EAD) analyses, we used A. cerana, A. dorsata, and A. florea foragers collected on inflorescences at Kunming and XTBG. EAD couples the measurement of antennal responses with a GC analysis of the compounds from a mixture, natural alarm pheromone. To test antennal responses to known amounts of pure compounds, we used Electroantennography (EAG) with A. cerana foragers collected on Kunming inflorescences. Finally, we sampled foragers from colonies for our GC-MS analysis. For our GC and GC-MS chemical analyses and EAG and GC-EAD procedures, we followed standard methods (see Supplemental Methods, S1).

Commercially available isopentyl acetate (IPA), 2-methyl butyl acetate (2MBA), phenyl methanol (PM), octan-1-ol (OH), (E)-oct-2-en-1-ol (OEH), isopentanol (IP), benzyl acetate (BA), octyl acetate (OA), gamma-octanoic lactone (GOL) and phenethyl acetate (PEA) were obtained from TCI Co. Ltd (Tokyo, Japan). (E)-3-Dec-2-en-1-ol (DH) was synthesized by NaBH₄ reduction of (E)-3-Dec-2-en-1-yl acetate (DA) and (E)-oct-2-en-1-yl acetate (OA) were synthetized by acetylation of (E)-3-Dec-2-en-1-ol and (E)-oct-2-en-1-ol, respectively, using acetyl chloride in hexane with triethylamine. Finally, and then purified with silica chromatography.

Exp. 2: A. cerana alarm flight experiment. We tested the behaviour of returning foragers responding to alarm pheromone (natural, synthetic mixtures, and synthetic individual components) released at the nest entrance, providing the context of encountering alarm odour in flight. Alarm flight bioassays were conducted on windless sunny days (above 20 °C) in Kunming. To determine which alarm pheromone components would trigger alarm flight, we applied compounds to a clean filter paper strip (15 mm by 4 mm) that was attached to the end of a clean wood stick (2 mm in diameter, 30 cm long) using a clean no. 0 insect pin. After evaporation of the dichloromethane for 10 s (see above), we positioned each filter paper 30 cm away from the hive entrance, at a 45° angle, to avoid blocking the entrance. Bees normally fly in a straight line directly into the colony. Bees responded to the natural release of alarm pheromone by turning away (fly to a track that was in an acute angle (< 90 °) with the straight normal returning trail) before entering the nest. We therefore used this turning behaviour as a bioassay of an alarmed bee.

To describe this turning behaviour qualitatively, flight motions were recorded with an HDR-CX450 video camera (Sony, Japan) at 50 frames per second (fps), providing a 2.7 × 1.5 m field of view. We randomly selected a subset of returning bees (15 bees from one colony) and digitized their flight tracks with Tracker v4.92 software (Douglas Brown, USA). We digitized the bee’s position each 20 s, beginning when she entered the field of view and ending when she entered nest. These flight tracks only capture a flattened, 2-D perspective, but illustrate the context of encountering alarm odour in flight. Alarm flight bioassays were conducted on floral resources at spaced at least 4 km apart.
Given that odours will dissipate over time, we conducted separate trials in which we analysed the number of bees that exhibited alarm behaviour over time, counting the number of alarmed bee each min over 5 min.

Exp. 3: A. cerana feeder foraging alarm experiment. Separately, we tested the behaviour of foragers that encounter alarm pheromone while foraging at a rich sucrose feeder. The most common natural analogue for the experiment would be foragers avoiding alarm pheromone released by a foraging conspecific that had been attacked by a predator. Individually marked foragers were trained to a feeder (an inverted 70 ml glass vial placed on a grooved base) that was 100 m away from the nest, and contained odourless, 35% (w/w) sucrose solution (analytical grade sucrose, AR, Xilongchem, CN) prepared in distilled water. Once the test began, we replaced the training feeder with a clean experimental feeder and, simultaneously, set out an identical control feeder, both displaced by 80 cm from the original training location and spaced 80 cm apart. Identical pieces of filter paper were placed under each feeder. We pipetted out four equidistant dots (2.5 µl each) of IPA, BA, OA, OEA and DA solution onto the filter paper of the experimental feeder and four equidistant dots (2.5 µl each) of DCM solvent on the control feeder. After complete evaporation of the solvent for 10 s, we recorded the choice of each forager. We tested the compounds in ascending quantity order (0.1 µg, 1 µg, 10 µg and 100 µg). Foragers were captured and removed after making a choice, and thus each forager was tested only once. For each sample (four concentrations of five compounds), we recorded the choices of 15 bees from each of three colonies. Each colony was tested on a separate day and under similar sunny weather (15 to 23 °C, moist content 55%).
Exp. 4A: Floral foraging information interception experiments with A. dorsata and A. cerana. Both A. dorsata and A. cerana forage at Calliandra haematocephala inflorescences (each inflorescence approximately 3–7 cm in diameter and separated by approximately 10 cm) at XTBG15. We therefore used these inflorescences to test if A. dorsata could detect and exhibit alarm to the primary components of A. cerana alarm pheromone (IPA, GOL, DA, OA, and BA). A. dorsata alarm pheromone shares IPA, DA, and OA with A. cerana alarm pheromone15, 26. However, A. dorsata alarm pheromone contains GOL, which A. cerana does not15. A. dorsata forager alarm pheromone does contain trace amounts of BA at 0.07% the level found in an A. cerana forager (value based upon GC-FID standard curve shown in Fig. 1E). To deliver odours, we built a stimulus controller (Fig. S4) consisting of a 550-CE air pump (4 ml/s, Nidec, Japan), an active charcoal filter (inner diameter of 1.5 mm), a HXL170 electromagnetic switching gas valve (Zile, China), a PTFE tube, and an odour pipette to deliver the test compounds to an inflorescence visited by honey bees. We only tested one bee at a time. When a bee landed on the inflorescence, we manually triggered a 3 s stimulus of clean, filtered air only (control phase), followed by 1.5 s of the odour stimulus (stimulus phase) added to the airflow. This method ensured that we could control for bee responses to the airflow alone. We recorded if the bee remained or departed during the stimulus phase. Only bees that did not depart during the control phase were used. Overall, only 15.3% of foragers departed in response to the clean airflow only. It was not possible to capture bees since they departed so rapidly in response to alarm pheromone and the same bee could therefore have returned to the same inflorescence, though this was unlikely given the abundance of nearby inflorescences in the C. haematocephala tree. We tested a concentration series of alarm pheromone components deposited on the filter paper strip in the odour pipette. DCM blank controls were tested first, and then the concentrations were tested in ascending order. To avoid odour cross-contamination, the odour paper, odour pipette, and the connecting glass joint (Fig. S4) were replaced with clean ones for each different compound.

Exp. 4B: Floral foraging information interception experiments with A. florea and A. cerana. We observed A. florea and A. cerana foraging together on the large, clustered inflorescences of the date palm, Phoenix dactylifera. We conducted nine replicates, three replicates per tree with three different trees. We conducted trials on three separate days between Dec 2015 and Jan 2016. On each day, we conducted three trials, each with a different tree. A. florea and A. cerana alarm pheromones share IPA, DA, OA, and BA, though A. florea alarm pheromone only contains trace amounts of BA (0.44% the quantity found in A. cerana, Fig. 1E). In this plant species, the flowers are tightly clustered in a large sheaf (30 cm in diameter, 50 cm in length with multiple small flowers <3 cm apart) and thus targeting the entire inflorescence with the odour pump was not possible. We therefore tested the group response of all bees foraging on a single large inflorescence to odour presented in a paper strip containing DCM as blank or 10 μg (2 to 10 A. cerana eq) of IPA, OA, BA, and DA or 5 eq of dissected A. cerana or A. florea sting glands impregnated on a filter paper strip, respectively, to the inflorescence. DCM was the solvent, and the same volume was used for control and experimental treatments. We waited 30 s to allow the odour to diffuse, and then counted the total number of A. cerana and A. florea foragers over the next 5 min. The odour was therefore presented for 5.5 min. We waited for approximately 45 min between tests to allow foraging to recover.

Statistics. For exp. 1A (GC-FID), we analysed component quantities with one-way Analysis of Variance (ANOVA) and used Tukey's Honestly Significant Difference (HSD) test for post-hoc comparisons. To compare the quantities of BA among the species that we studied, we log-transformed the quantities measured (ng) and used one-way ANOVA, and Tukey's Honestly Significant Difference (HSD) test for post-hoc comparisons with Sequential-Bonferroni corrected significance levels. We used SPSS 22 (IBM, US) for this analysis.

For exp. 1B (EAG), we rectified the response data (mV) by subtracting, per bee, the response to each compound from that bee's response to the blank control and then log-transforming the resulting data. Because bees exhibited no EAG responses to the lowest quantity (0.1 pg, Fig. 2C), we used Dunnett's method (which corrects for potential Type 1 error) to make all pairwise comparisons between 0.1 pg and each higher quantity. We included sample sites as a random effect. We used SPSS 22.0 (IBM, US) for this analysis.

For the exp. 2A compound comparison, we used a multiway frequency analysis, based on an ANOVA performed on a generalized linear model (GLM, Poisson model, log link) as described by Vokey22 to investigate the effects of different compounds on the number of alarmed bees at the nest entrance. We used a multiway frequency analysis with colony, chemical, and “bee state” (i.e., alarmed or quiet) as categorical factors and number of bees as cell values. For example, one row of data (corresponding to one cell on the 3-dimensional matrix) would have an associated colony number, a chemical, and a binary outcome category variable (e.g., “alarmed”), and the corresponding cell value would then be the number of alarmed bees that were from a given colony when a given chemical was used. We chose this analysis method because it is ideal for repeated-measures discrete data and does not suffer from some of the complexities associated with GLMM. In multiway frequency analysis, all effects, including colony, are treated as fixed. However, similar to specifying factors as random effects in GLM, this
technique allows the model effects to be assessed after the variance due to all other effects is removed (see 40 for full explanation). For this analysis, we report $G^2_{df}$, the likelihood ratio Chi-square statistic 40.

For exp. 2B (nest entrance BA quantity-response assay), we analysed the effect of different quantities of BA upon the number of alarmed bees at the nest entrance, also using a multiway frequency analysis with colony, concentration, and bee state as factors and number of bees as cell values.

For exp. 3, we tested if A. cerana foragers would avoid feeders with different treatment odors by using Chi-square tests. We used a null hypothesis expectation of equal visitation at both feeders if there was no effect of compound. We used Excel 2007 (Microsoft, US) for this analysis.

For exp. 4A (A. dorsata floral repellence assay), we analysed the A. dorsata data (number of alerted bees) using multiway frequency analysis (R v. 3.3.2). In exp. 4B (A. florea and A. cerana floral repellence assay), we also used a multiway frequency analysis.

Unless otherwise specified, we used JMP 12 (SAS, US) for ANOVA analysis. We present the mean ± 1 standard error. For our ANOVA, we used residuals analysis to confirm that the data met model assumptions. In the repeated-measures model, we chose a linear model based upon its fit with the data (repeated-measure discrete data). In the experiments that we analysed with ANOVA, each bee was analysed or tested only once to conform to expectations of data independence. For the multiway frequency analysis, the data do not need to conform to a particular distribution or be independent (see 40 for full discussion).

Results

A. cerana and A. florea alarm odours are produced by the sting glands. In A. cerana and A. florea, headspace SPME analysis of volatiles emitted by an alarmed forager had the same chemical composition as sting gland volatiles (match for all major peaks, Fig. 1). Thus, all A. cerana and A. florea alarm odours can be found in the sting gland (Fig. 1A). We therefore proceeded to use only dissected A. cerana and A. florea sting glands in our subsequent assays testing A. cerana and A. florea alarm pheromone.

Exp. 1: BA was more abundant and EAD responsive than other active alarm compounds in A. cerana foragers. In A. cerana, the sting gland is a major source of alarm pheromone produced by foragers (Fig. 1A). To identify compounds, we used authentic standards and compared the MS spectrum and retention times of these standards, run on the same equipment, with the analyses of natural alarm pheromones. GC-MS analyses showed that sting gland alarm pheromone in guards and foragers consists of the following main components: 3-methyl butanol (MB), isopentyl acetate (IPA), phenyl methanol (PM), (E)-oct-2-en-1-ol (OH), benzyl acetate (BA), octyl acetate+(E)-Oct-2-en-1-yl acetate (OA+OEA, not distinguishable on the HP-5 column, but are distinct on DB-WAX column, Fig. S1), phenethyl acetate (PA), (E)-dec-2-en-1-ol (DH), and (E)-dec-2-enyl acetate (DA).

There were significant overall differences in the relative abundance of compounds produced by forager and guard bees (ANOVA: $F_{7,112} = 22.21$, $P < 0.001$, Fig. 1B). Foragers produced significantly more BA than guard bees.

Figure 2. Antennae of foragers from all three bee species (A. cerana, A. dorsata, and A. florea) are highly sensitive to BA. (A) GC-EAD responses to the HS-SPME extracts of dissected A. cerana stings (1 eq). Relative abundances of each compound and representative EAD signals for each bee species are shown. (B) Responses of antennae (EAG) to different quantities of BA EAG (Ctrl = solvent only). The bar (∗) indicates quantities that elicited significantly higher responses as compared to the control (Dunnett’s test, $P < 0.01$). Means and standard errors are shown. The inset photo shows the antennal preparation.
BA, which was the only compound that significantly repelled foragers at repulsive quantities of each compound were: 100 µg of GOL, an abundant (Tukey’s HSD test, \( P < 0.05 \), Fig. 1B).

A. dorsata and A. florea share the same major volatile compounds in their alarm pheromones, including BA (Fig. 1C,D). However, BA is only found trace levels in A. dorsata and A. florea (Fig. 1E), and probably was not previously identified because it is present at trace levels and may have been confounded with the DA peak.

A. cerana forager antennae responded to peaks of IPA, PM, OH, BA, OA and DA (GC-EAD analysis, Fig. 2A). A. dorsata and A. florea antennae responded to IPA, BA, OA and DA.

Because we were primarily interested in A. cerana, we focused on this species. A. cerana antennae exhibited differential sensitivity (\( F_{6, 172} = 73.14, P < 0.001 \)). The strongest responses were to BA and DA (Tukey’s HSD test: \( P < 0.05 \), Fig. 2B). As expected, responses increased with higher quantities (EAG quantity within subject effect: \( F = 432.54, P < 0.0001 \)). There was no significant between subject effect of site (\( F_{5, 171} = 0.246, P = 0.937 > 0.05 \)) or within subject effect of the interaction site*quantity (\( F_{5, 171} = 1.365, P = 0.081 > 0.05 \)).

The difference threshold, the lowest quantity that elicited a significantly greater antennal response than the control, was at 0.1 ng (Dunnett’s method: \( P = 0.002 \)) for BA (Fig. 2C). For comparison, the EAG difference thresholds for IPA, OA and DA were 100 ng, 100 ng and 10 ng, respectively (data from [16] using the same method). Thus, A. cerana foragers were more sensitive to BA than to any other major alarm pheromone component.

Exp. 2: BA alarmed flying returning foragers. After determining that BA was an abundant alarm pheromone component that elicited strong antennal responses, we tested its efficacy in two different contexts: alarm at the nest entrance (exp. 2) and alarm at food sources (exp. 3&4). We presented the test compounds on paper strips at the nest entrance (Fig. 3A) to simulate sting alarm pheromone release following predator attack or detection. We used a multiway frequency analysis with colony, chemical, and “bee state” (i.e., alarmed or quiet) as categorical factors and number of bees as cell values. So, for example, one row of data (corresponding to one cell on the 3-dimensional matrix) would have an associated colony number, a chemical, and a binary outcome category variable (e.g., “alarmed”), and the corresponding cell value would then be the number of alarmed bees that were present from colony x when chemical x was used. We found a significant effect of compound (\( G^2 = 239.14, P < 0.0001 \)), but no effect of colony (\( G^2 = 3.44, P = 0.18 \)), and no interaction between colony and compound (\( G^2 = 13.18, P = 0.52 \)). Post hoc analyses (Chi-square Z-tests) revealed that BA and mixtures with BA increased alarm responses (\( P < 0.017 \), Fig. 3C). In addition, BA seemed to act on a different time scale. BA has a lower vapour pressure than IPA and was slower to take effect than IPA (Fig. 3B).

There was a significant effect of compound concentration (\( G^2 = 92.74, P < 0.0001 \) (see Fig. 3D), but no significant effect of colony (\( G^2 = 0.187, P = 0.91 \)) and no colony * concentration interaction (\( G^2 = 4.26, P = 0.83 \)). The ratios of alerted to observed bees at BA quantities of 0.1, 1 and 10 µg were significantly higher than the ratio of bees responding to the blank control (Chi-square tests: \( P < 0.017 < P_{\text{Sequential Bonferroni}} = 0.01 \)). Thus, an alarm response could be elicited by only 0.1 µg of BA, corresponding to 0.03 eq.

Exp. 3: BA repelled foragers from landing on feeders. We next tested the effects of A. cerana sting alarm pheromone compounds on bees foraging at a feeder, simulating the situation of a forager encountering alarm pheromone released by an attacked conspecific. Although the feeder offered highly rewarding, concentrated sucrose solution ad libitum, IPA, BA, and DA repelled A. cerana foragers (Fig. 4A). The minimum repulsive quantities of each compound were: 100 µg IPA (\( > 1 \) eq, \( \chi^2 = 15.52, n = 45 \) bees, \( P = 0.017 < 0.050 \)), 1.0 µg of BA (\( < 1 \) eq, \( \chi^2 = 8.49, n = 45 \), \( P = 0.037 < 0.050 \), Fig. 4A), and 10 µg of DA (\( > 1 \) eq, \( \chi^2 = 15.52, n = 45 \), \( P = 0.017 < 0.050 \)). These threshold differences are summarized in Fig. 4B. Thus, foragers were most sensitive to BA, which was the only compound that significantly repelled foragers at \( < 1 \) eq.

Exp. 4: Multiple bee species avoid A. cerana alarm pheromones on floral resources. Previously, we found that A. cerana could intercept and cedesdrop upon some of the alarm pheromone components of A. dorsata, but it was unknown if A. dorsata or A. florea could detect and respond appropriately to A. cerana alarm pheromone components.

In the floral bioassay of A. dorsata (exp. 4A), there were significant effects of compound (\( G^2 = 37.35, P < 0.0001 \)), concentration (\( G^2 = 237.0, P < 0.0001 \)) and the compound*quantity interaction (\( G^2 = 31.8, P = 0.011 \)) (Fig. 5). There was no significant effect of field site (\( G^2 = 1.3, P = 0.52 \)). None of the other interactions was significant. In pairwise post-hoc comparisons (all tests passed the Sequential Bonferroni correction), the minimum quantities that elicited significant avoidance were: BA at 1 µg (\( P = 0.0000 < 0.001 \)), GOL at 1 µg (\( P = 0.0000 < 0.001 \)), DA at 10 µg (\( P = 0.0001 < 0.001 \)), and IPA at 1 µg (\( P = 0.0065 < 0.01 \)). BA at 10 µg repelled more A. dorsata foragers than 1 µg of GOL (\( P = 0.010 < P_{\text{Sequential Bonferroni}} = 0.025 \)) or 10 µg of OA (\( P = 0.020 < P_{\text{Sequential Bonferroni}} = 0.025 \)). However, BA at 10 µg repelled as many foragers as 10 µg of GOL (\( P = 0.10 > 0.05 \)). Thus, A. dorsata was repelled by 0.3 eq of A. cerana BA (Fig. 1E). A. dorsata was also repelled, as expected, by 0.15 eq GOL, an A. dorsata-specific alarm component (Fig. 5A).

In the floral bioassay of A. florea and A. cerana (exp. 4B), there were no significant effects of tree (\( G^2 = 0.59, P = 0.75 \)) or bee species (\( G^2 = 0.02, P = 0.88 \)), and none of the interactions between any factors was significant (\( P < 0.05 \)). There was a significant effect of compound (\( G^2 = 133.67, P < 0.0001 \)) such that natural sting alarm pheromones and certain synthetic sting pheromone compounds significantly reduced forager visits. The most inhibitory treatments were A. cerana forager natural sting pheromone (5.0 eq) and 10 µg BA (2.8 eq Fig. 5, Tukey’s HSD, \( P < 0.05 \)).

Discussion
Signal interception, which can be a form of eavesdropping, has generally been thought of as providing a benefit to the interceptor at the disadvantage of the signaler, although there are exceptions[4]. There are multiple examples
of such information usage in stingless bees, ants, termites, social wasps. Here, we considered the case of a pollinator guild in which interspecific sensitivity to a major alarm pheromone component, benzyl acetate (BA), of one of the most common pollinators, *A. cerana*, should also benefit other bee species without, in theory, being detrimental to the signal sender. Such mutualistic or cooperative information sharing occurs in a wide variety of animals, vertebrate and invertebrate. In our case, this mutualism likely arose because many social insects have evolved alarm pheromones that help colonies deal with danger, and there is no evident disadvantage if colonies of the same or different species also use this information to avoid dangerous food or nest predators. In addition, closely related species like the different species of *Apis* share similar alarm signals. One might argue that such information could be used deceptively by one species to deter another from visiting a rich food source. However, the most important selective pressure for the evolution of social insect alarm signals is likely the benefit that alarm signals provide for the colony, the unit of reproduction. It makes little sense for bees to evolve alarm pheromones that deceive competitors but leave their own colonies vulnerable by decreasing the honest information content of alarm signals. Crying wolf is not effective if it reduces one's defenses against wolves. We therefore predict that such alarm information will be honestly produced and mutualistically used by sympatric social bee species. Our data support this hypothesis by showing that *A. dorsata*, *A. florea*, and *A. cerana* avoided BA on floral food and that *A. cerana* also avoided BA at its own nest.

**Figure 3.** Alarm responses of returning *A. cerana* bees to alarm pheromone components and mixtures in front of the nest. (A) Comparison of the flight track of foragers in response to 10 μg (5–10 eq) of the major components separately or in combination. The inset shows a scaled schematic of the hive, sample placement, and video field of view. Bees returning to the nest normally flew directly in (Control). Alarmed bees exhibited sharp turns and looping flights. (B) Examples of the effect of time on forager flights in response to BA and IPA. (C) Percentage of alarmed bees out of all approaching bees within 6 min in response to different components and mixtures. We presented 10 μg of each component (5–10 eq) or the dissected sting glands from five bees. (D) Percentage of alarmed bees to different quantities of BA. Different letters indicate significant differences (Chi-squared tests, *P* < 0.01, Sequential Bonferroni corrected). Means and standard errors are shown.
Our results also support the volatility hypothesis, that some alarm pheromone compounds have been selected based upon their ability to endure and provide lasting information. A potential advantage to intercepting olfactory signals about danger, as compared to typical visual or acoustic danger signals, is that an olfactory danger signal can persist after the signaller departs. However, it may also be important to activate initial defences rapidly. The most volatile compounds, like IPA, can trigger an initial alarm response, but more persistent compounds, such as BA and DA, have greater utility as longer-term markers of danger. The multi-component blends that we find in sting alarm pheromones may therefore have been shaped by their inherent toxicity, detectability, and temporal dynamics (persistence).

For example, we found that *A. cerana* forager sting alarm pheromone is particularly rich in BA, which is significantly less volatile than the most abundant component, IPA. BA has strong effects. At the nest entrance, returning bees, most likely foragers, showed aversive alarm responses to natural sting pheromone, a synthetic combination of four major components, BA, DA, and BA + DA. However, BA stood out as the single compound that most strongly elicited aversion when presented at the nest entrance (Fig. 3C). This behaviour matched the high sensitivity of forager antennae to BA (Fig. 2B).

Although BA has been previously identified in the sting alarm pheromone of *A. mellifera*22, we provide the first identification of BA in *A. cerana, A. dorsata* and *A. florea*, likely because the technique previously used, a
Carbowax column, coated with solid phase polyethylene glycol (PG) cannot separate DA and BA. Since A. dorsata and A. florea also produce BA in their respective alarm pheromones, albeit in trace amounts, we cannot state that these two species are eavesdropping upon the BA in A. cerana alarm pheromone. Eavesdropping can only be conclusively demonstrated when the eavesdropper does not produce the compound in question. However, we predict that other pollinating bees, particularly non-Apis species that are less likely to produce BA, may eavesdrop upon BA to avoid danger.

The higher levels of BA in A. cerana foragers as compared to guard sting alarm pheromone matches what is known in A. mellifera, in which prior research demonstrated higher levels of BA, BH, and 2-nonenol in foragers than in guards, fanning bees, or comb bees respectively. Elevated levels of BA in foragers may be reasonable, if BA is an important alarm pheromone compound that foragers use to mark dangerous food sources or when encountering danger upon returning to the nest entrance. What is the function of the other alarm pheromone components? In the context of foraging, prior work showed that (E)-dec-2-en-1-yl acetate (DA) was effective at repelling A. cerana foragers. OA was identified in all honey bees, but the function of OA in nest defence and repulsing foragers appears to be weak. OEA was first identified in A. mellifera, and our analysis also revealed it in A. cerana. OEA cannot be separated from OA by using general HP-5 columns; thus OEA may also occur in other Apis species and the functions of these two compounds are uncertain. EAD analyses showed that A. cerana antennae responded to a peak consisting of OA and OEA (Fig. 2A). It therefore unclear which of these two compounds A. cerana can detect. Possibly, the lack of avoidance to OA shown in our assays is due to an inability to detect OA. However, given the relatively high abundance of OA (Fig. S1) and the strong general ability of A. cerana to detect multiple volatile compounds within this size range (Fig. 2A), we suggest that A. cerana can detect OA.

We hypothesize that the likelihood of encountering BA shapes how useful it is for other bee species. BA is produced in large quantities by A. cerana, which also appears to be more common on floral resources than either A. dorsata or A. florea, but not known. Moreover, A. dorsata is a migratory species whose sympatric presence with A. cerana can seasonally fluctuate. Thus, testing this hypothesis about species abundance and information utility will require a detailed understanding of species populations in time and space.

Data accessibility. All data are accessible in supplemental datasheets.

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Author Contributions

Conceived and designed the experiments: P.W., K.T. and J.C.N. Performed the experiments: P.W., Y.C., S.D., Y.Q., H.Z. and J.L. Analysed the data: P.W., B.H. and J.C.N. Contributed reagents, materials, or analysis tools: P.W., H.Z., K.T. and J.C.N. Wrote the paper: P.W., B.H., K.T. and J.C.N.

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