Repellent foraging scent recognition across bee families

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Abstract – Honeybees and bumblebees avoid probing flowers that have been recently depleted by conspecifics, presumably repelled by odours deposited by the previous visitor (foraging scent marks). Here we show that females of the solitary wool-carder bee Anthidium manicatum (Megachilidae) discriminate against previously visited inflorescences (Stachys officinalis), and that discrimination is equally strong regardless of whether the previous visitor is conspecific or belongs to a different bee family (Bombus terrestris, Apidae). Conversely, workers of B. terrestris responded differentially to different previous visitors, with previous visits by A. manicatum eliciting the most pronounced repellent effect. This finding may have resulted from the bumblebees’ avoidance of impending aggression by territorial A. manicatum males. Our results emphasize that foraging scent mark recognition is not necessarily linked to sociality, but a trait of individuals foraging in an unpredictable flower visitor community.

scent marks / footprints / flower discrimination / Bombus / Anthidium

1. INTRODUCTION

Plant-bee interactions are frequently characterised by a high degree of generalization, meaning that sympatric species of bees broadly overlap in their choice of food plants (Waser et al., 1996). Although evidence is scant, this implies that the extent of competition for floral resources among members of different species rivals that among conspecifics, and that foraging decisions of individuals should maximize energy gains in response to the behaviour of a range of competing species. This hypothesis has been applied to guilds of bumblebees (Bombus sp.) faced with the decision of whether to land on and probe a particular flower or inflorescence (Stout et al., 1998). Bumblebees were shown to reject flowers that have been recently visited by conspecifics, thus discriminating against flowers with little or no nectar content (Stout et al., 1998; Stout and Goulson, 2001a; Goulson et al., 1998). Doing cross-species tests with four species of Bombus, Stout et al. (1998) found that the likelihood of flower rejection was largely independent of the species affiliation of the previous visitor. Although other possible mechanisms have not been completely ruled out, mounting evidence suggests that flower discrimination is based on detection of residues of volatile chemicals deposited during previous visits (Goulson et al., 2000). Experiments have demonstrated that bumblebees reject flowers that were treated with extracts of Bombus tarsi, containing mixtures of long-chain hydrocarbons. These mixtures were similar (but not identical) among species of Bombus, thus allowing for the recognition of heterospecific scent marks (Goulson et al., 2000). It is not clear whether the deposition of scent marks is an active process, e.g.
mediated by imminent tarsal gland secretion, or whether the volatile residues constitute unavoidable ‘footprints’, left wherever bumblebees walk (Stout et al., 1998).

Using information of scent deposits on flowers is not restricted to bumblebees. Honeybees (*Apis mellifera* L.) are known to respond to repellent substances secreted by their conspecifics’ mandibular glands (Giurfa, 1993; Giurfa and Nunez, 1992). Furthermore, female solitary *Anthophora plumipes* (Pallas) have been claimed to use non-visual cues for discriminating among flowers, with recognition of conspecific scents being the likely mechanism (Gilbert et al., 2001).

Because the repellent scent compounds found in bumblebees (long-chain alkanes and alkenes) are common to cuticles of most insects, it is possible that similar marks are deposited by a broad range of flower visitors (Goulson, 2003). It is an open question whether recognition of scent deposits occurs between different genera or families of bees, and whether different species of previous visitors elicit differential repellent effects (see, e.g., conflicting evidence for *Apis/Bombus* in Williams (1998) and Stout and Goulson (2001b)). In the present study we investigated whether acceptance of flowers by female solitary wool-carder bees, *Anthidium manicatum* (L.), is influenced by previous visits of worker bumblebees, *Bombus terrestris* (L.) and vice-versa, and whether bumblebees discriminate between different species of previous visitors.

### 2. MATERIALS AND METHODS

The observations were carried out within a 15 m² stand of *Stachys officinalis* (L.) in the Botanical Gardens of the Heinrich-Heine-Universität in Düsseldorf. Observations were made from 0800 h to 1600 h (local time) from July 14 to 28 in 2003. Throughout this period the weather was hot and dry. *Stachys officinalis* (Lamiaceae) has vertical inflorescences with one to 30 open heart-shaped flowers providing nectar at the base of a short corolla tube. During the time of the study *S. officinalis* was highly attractive to several species of bees, with workers of *Bombus terrestris* and *B. pascuorum*, and female *Anthidium manicatum* being the most abundant visitors (in the given order), which were therefore selected as focal species.

For measuring the rates of rejection of ‘unvisited’ (control) inflorescences by individual foragers, whole inflorescences were covered with mesh bags for at least 3 hours (mostly overnight) before behavioral tests were conducted, so that nectar could accumulate and possible scent-marks evaporate. For an individual acceptance/rejection test, an inflorescence was cut with a short stem and exposed in the foraging path of an individual bee using forceps. It was recorded as ‘rejected’ if the bee would approach the inflorescence but not land on it or land on it briefly but not probe any flower for nectar. Respectively, inflorescences were regarded as ‘accepted’ if the forager would land on it and probe at least one flower for nectar (see Stout et al., 1998). An analogous procedure was used to measure rates of rejection for inflorescences that had been visited shortly before the test situation. For this, we cut inflorescences immediately after they had been visited by a bee and presented them to foraging bees within the next 5 minutes. In order to contrast the effects of conspecific versus heterospecific visits on rejection probability, we conducted independent test series using different combinations of the previous visitor species and the test species. Complete reciprocal test series were conducted for females of *A. manicatum* and workers of *B. terrestris*. *Bombus pascuorum* workers were used as additional previous visitors in tests on *B. terrestris* (see Tab. I for an overview of test combinations).

In an attempt to quantify the degree of nectar depletion in inflorescences resulting from visits by a given bee species we assessed the percentage of flowers probed per inflorescence. For quantitative comparison between bee species we distinguished

<table>
<thead>
<tr>
<th>Previous flower visitor</th>
<th>Test flower visitor</th>
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<tbody>
<tr>
<td><em>B. terrestris</em></td>
<td><em>B. pascuorum</em></td>
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<tr>
<td><em>B. terrestris</em></td>
<td>X</td>
</tr>
<tr>
<td><em>B. pascuorum</em></td>
<td>X</td>
</tr>
<tr>
<td><em>A. manicatum</em></td>
<td>X</td>
</tr>
<tr>
<td>unvisited flowers</td>
<td>X</td>
</tr>
</tbody>
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Table I. Overview of test combinations.
Repellent foraging scents

3. RESULTS

3.1. Rejection of inflorescences previously visited by conspecifics

First, we compared the rejection rates of unvisited versus visited inflorescences within each of the three species separately. Inflorescences previously visited by conspecifics were rejected significantly more often than unvisited ones in *B. terrestris* (N = 86; df = 1; $\chi^2 = 5.51; P < 0.05$) and *A. manicatum* (N = 61; df = 1; $\chi^2 = 13.87; P < 0.001$). In *B. pascuorum* there was a tendency in the same direction, but it was not significant (N = 67; df = 1; $\chi^2 = 3.10; P < 0.1$). Of the three species, *A. manicatum* showed the highest absolute rate of rejection of visited inflorescences (70.9%; see Fig. 1). However, when compared statistically, there was no significant difference between species concerning the likelihood of rejecting inflorescences previously visited by conspecifics (N = 113; df = 2; $\chi^2 = 4.06$; n.s.).

3.2. Rejection of inflorescences previously visited by heterospecific foragers

We analyzed whether individuals of *B. terrestris* and *A. manicatum* also reject inflorescences in response to visits by foragers of a different species, and if so, whether the rate of rejection differs depending on the species of the previous visitor.

In comparison to unvisited controls foraging *B. terrestris* were significantly more likely to reject inflorescences previously visited by *A. manicatum* (N = 86; df = 1; $\chi^2 = 13.877; P < 0.001$), but there was no difference for those previously visited by *B. pascuorum* (N = 76; df = 1; $\chi^2 = 0.171$; n.s.). When we compared the rejection rates of *B. terrestris* after previous visits by conspecifics, previous visits by *B. pascuorum* and previous visits by *A. manicatum*, there was a significant difference (N = 125; df = 2; $\chi^2 = 10.14; P < 0.01$), with previous visits by *A. manicatum* eliciting the strongest repellent effect (see Fig. 2).

In comparison to unvisited controls foraging *A. manicatum* were significantly more likely to reject inflorescences previously visited by *B. terrestris* (N = 66; df = 1; $\chi^2 = 10.867; P < 0.001$). No data were available to compare the effect of previous visits by *B. pascuorum*. There was no significant difference when we
compared the rejection rate of *A. manicatum* after previous visits by conspecifics with that after previous visits by *B. terrestris*, (N = 67; df = 1; $\chi^2 = 0.54$; n.s.).

### 3.3. Foraging behaviour

The three species varied significantly in the percentage of flowers probed per visited inflorescence (N = 267; df = 2; $\chi^2 = 19.14$; $P < 0.0001$), with *B. terrestris* visiting the largest proportion of flowers, followed by *B. pascuorum* and *A. manicatum* (Fig. 3). The low percentage of flowers probed by foraging *A. manicatum* corresponded to our impression that they were prone to rapid switches between inflorescences and moved quickly through the patch.

### 4. DISCUSSION

Our data demonstrate that *A. manicatum* rejects flowers that have recently been visited by conspecifics. As in bumblebees, it is likely that this behaviour is mediated by volatile chemicals deposited by the previous visitor. This is the first report of such a repellent effect for a megachild bee and strengthens the view that use of scent marks is not confined to social Apidae. The only other solitary bee that reportedly uses scent-marks is *Anthophora plumipes* (Gilbert et al., 2001). Traditionally, scent marking has been regarded as a typical trait of social bees that enhances colony foraging efficiency, emphasizing attractive effects that guide nest mates to rewarding flowers and minimizes the time colonies have to invest in searching for food (Cameron, 1981; Schmitt et al., 1991). However, on an individual basis both attractive and repellent scent marks may also be beneficial, e.g. by reducing the risk of immediate revisits of depleted flowers by a given individual or by facilitating rapid revisits of rewarding flowers that cannot be completely depleted during one single visit. Our finding that the solitary *A. manicatum* uses scent-marks clearly emphasizes these individual benefits and suggests that repellent scent marks are not necessarily a way of social communication in the social Apinae. Generally, it is still not clear if the repellent scents are actively deposited or simply unavoidable footprints left by foragers, which is an important aspect when considering their function.

Solitary *A. manicatum* as well as social *B. terrestris* clearly rejected flowers that were previously visited by the other species respectively, suggesting that recognition of scent marks takes place across members of different bee families. In *B. terrestris* there was also a significant difference of the strength of the repellent effects elicited by different species of previous visitors, suggesting that *B. terrestris* can (and does) discriminate between scent marks of different bee species. But why should foraging decisions depend on the species affiliation of the previous visitor? Discrimination would clearly be beneficial if the extent of resource depletion differs between species, an assumption that may seem validated by the evident differences in the percentage of probed flowers per inflorescence. However, the thoroughness of foraging by the different species does not correspond to the observed repellent effects. Especially the strong repellent effect of previous visits by *A. manicatum* is unlikely related to lack of nectar in the visited flowers, since *A. manicatum* individuals are the least thorough in their foraging. This finding also emphasizes that the decision making process is not based on the direct perception of nectar levels present in the flowers.

We hypothesize that the strong repellent effect of previous visits by *A. manicatum* is related to the territorial behaviour of the species. Male *A. manicatum* establish and defend small resource-based mating territories around...
their females’ food plants. The males constantly patrol these territories and chase away other flower visitors, including bumblebees, by physically attacking them with abdominal spines. The harassed individuals often carry away severe wounds or even die after the attacks (Pechuman, 1967; Wirtz et al., 1988; Wirtz et al., 1992). The strongly decreased acceptance rate of flowers previously visited by *Anthidium* in *B. terrestris* could be a result of this aggressive behaviour. Bumblebees may associate perceived *Anthidium* scent with previously experienced attacks and avoid marked inflorescences.

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Résumé – Reconnaissance du marquage odorant répulsif suite au butinage parmi les familles d’abeilles. La recherche de nourriture occasionne pour les insectes butineurs des coûts sous forme de temps passé par visite de fleurs et d’énergie nécessaire pour exploiter les ressources. Afin de minimiser ces coûts, les butineurs doivent pouvoir différencier, avant même de les tester, les fleurs qui vont leur donner une récompense de celles qui ne vont rien fournir. De fait, il a été montré que les ouvrières de bourdons (*Bombus* sp.) discriminent les fleurs vides, i.e. celles déjà visitées par d’autres bourdons peu de temps auparavant. Des études antérieures ont suggéré que cette discrimination ne reposait pas sur une perception directe du nectar, mais que des marques odorantes laissées par le visiteur précédent étaient perçues et soumises à une évaluation indirecte de la teneur en nectar attendue. Cette capacité n’est pas limitée à l’espèce, car des marques odorantes peuvent être perçues réciproquement par d’autres espèces au sein du genre *Bombus* (Stout et al., 1998; Stout et Goulson, 2001a; Goulson et al., 1998). Dans cette étude nous montrons que l’enregistrement et l’utilisation des marques odorantes existent aussi chez les abeilles solitaires et que la perception réciproque a lieu aussi entre membres de différentes familles d’abeilles. Des femelles de l’abeille solitaire « tapisière » ou « cotonnière », *Anthidium manicatum* (L.) (Megachilidae), qui butinent principalement la Bétoine officinale (*Stachys officinalis*) discriminaient dans la même mesure les fleurs qui avaient été visitées peu auparavant par des individus de la même espèce et celles qui avaient été visitées par des bourdons terrestres (*Bombus terrestris*, Apidae). Chez les bourdons ter-

**Bombus / Anthidium / discrimination / butinage / marquage odorant / marquage par les tarse**

Zusammenfassung – Erkennung der abweisen-

den Duftmarkierung nach Sammelaktivität von

unterschiedlichen Bienenfamilien. Bei der

Nahrungssuche entstehen Blütenbesuchern Kosten

in Form der pro Blütenbesuch aufzuzwän-

den Zeit und der zur Ausbeutung der Ressourcen

notwendigen Energie. Um diese Kosten zu mini-

mieren, sollten Blütenbesucher deshalb bereits vor

der Beprobung in der Lage sein, zwischen beloh-

nenden und unbelohnenden Blüten zu unterschei-

den. Tatsächlich diskriminieren Hummelarbeiteri-

nen (*Bombus*) nachweislich gegen vor kurzer Zeit

von anderen Hummelindividuen besuchte, d.h. 
gelerte Blüten bzw. Blütenstände. Durch frühere

Studien wurde nahegelegt, dass diese Diskriminie-

rungh nicht auf einer direkten Perzeption von Nectar

beruht, sondern dass Duftmarken der vorherigen

Besucher wahrgenommen und zur indirekten

Bewertung des zu erwartenden Nektarangebots 

herangezogen werden. Diese Fähigkeit ist nicht auf

die Artgrenzen beschränkt, denn die Duftmarken 

können innerhalb der Gattung *Bombus* von Ange-

hörigen unterschiedlicher Spezies wechselseitig 

wahrgenommen werden (Stout et al., 1998; Stout 

und Goulson, 2001a; Goulson et al., 1998). In der 

vorliegenden Untersuchung zeigen wir, dass die 

Registrierung und Verwertung der Duftmarken 

auch bei solitären Bienenarten vorkommt und 

dass wechselseitige Wahrnehmung auch zwischen 

Angehörigen verschiedener Bienenfamilien statt-

findet. Weibchen der solitären Wollbiene (*Anhi-

dium manicatum* (L.), Megachilidae), die am 

Gewöhnlichen Ziest (*Stachys officinalis*) foragier-

ten, diskriminierten in gleichem Maße gegen Blü-

tenstände, die vor kurzem von Artgenossen besucht 

worden waren, als gegen solche, die vorher von 

Erdhummeln (*Bombus terrestris*, Apidae) besucht 

worden waren. Bei Erdhummeln dagegen war die 

Ablehnungswahrscheinlichkeit von der Artzuge-

hörigkeit des Vorbesuchers abhängig. Am stärksten

wurden Blütenstände abgelehnt, die vorher von
Wollbienen besucht worden waren. Wahrscheinlich gründete dieser Effekt nicht auf einem besonders niedrigen Nektargehalt von Wollbienen besuchten Blütenständen, sondern reflektiert die Tendenz von Hummeln, sich der Aggression territorialer Wollbienennägchen zu entziehen (durch Vermeidung von Wollbienenengrruch).


**REFERENCES**


