Honeydew production and honeydew sugar composition of polyphagous black bean aphid, *Aphis fabae* (Hemiptera: Aphididae) on various host plants and implications for ant-attendance

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**Abstract.** The black bean aphid, *Aphis fabae*, is polyphagous and its life cycle involves seasonal migration between summer and winter host plants. The aphids are regularly tended by honeydew-collecting ants. This study investigates whether differences exist in honeydew production and honeydew sugar composition for *A. fabae* subspecies feeding on various host plants and whether such differences reflect differences in the intensity of ant-attendance (*Lasius niger*).

*A. f. fabae* feeding on the perennial summer host, *Tanacetum vulgare*, produced twice the amount of honeydew (ca. 110 µg per aphid h−1) than when feeding on the annual host plants *Vicia faba* or *Chenopodium album*. Honeydew production of *A. f. cirsiicanthoides* feeding on the creeping thistle *Cirsium arvense* was the highest measured in this study (ca. 150 µg per aphid h−1). Total sugar concentration of the honeydew of *A. f. fabae* did not differ when feeding on various summer hosts, whereas the honeydew of *A. f. cirsiicanthoides* on *C. arvense* contained a significantly higher amount of total sugars. The trisaccharide melezitose was the dominant sugar in all the honeydew samples, except for the honeydew of *A. f. fabae* and *A. f. evonymi* feeding on the woody winter host plant *Evonymus europaeus*. The highest proportion of melezitose (80% of total sugars) was found in the honeydew of *A. f. cirsiicanthoides* feeding on *C. arvense*. In this subspecies, the intensity of ant-attendance was also highest. The results confirm our hypothesis, that the sugar richness of the honeydew (rate of honeydew secretion × total sugar concentration) along with the presence of the attractant sugar melezitose are the critical factors in determining the extent of ant-attendance.

In *A. f. fabae* feeding on the spindle tree *E. europaeus*, the total sugar concentration of the honeydew as well as the sugar composition differed significantly between generations.

**INTRODUCTION**

Aphids (Aphidoidea) have commonly developed mutualistic relationships with ants. The ants cover their carbohydrate demand by collecting “honeydew”, which is excreted by the aphids as a sweet waste product of their sugar-rich but amino acid-poor diet of phloem sap (Way, 1963). Ants in return defend the aphids from natural enemies. In general, the intensity of ant-aphid mutualism is positively correlated with honeydew quantity and quality as well as the sugar demand of the ants (Bristow, 1984; Bonser et al., 1998; Fischer et al., 2001). Honeydew is an aqueous solution consisting mainly of various sugars and amino acids (Auclair, 1963; Maurizio, 1985). Honeydew production and honeydew composition vary considerably between aphid species (e.g. Hendrix et al., 1992; Völk et al., 1999) but may also vary within a particular aphid species in an age-specific pattern (Hertel & Kunkel, 1977; Fischer et al., 2002) or when feeding on different host plants. Douglas (1993) demonstrated differences in the amino acid pattern of the polyphagous aphid species *Aphis fabae* on different host plants. Fischer & Singleton (2001) showed that the honeydew of *Chaitophrus populii* and *C. populeti* contained higher proportions of the trisaccharide melezitose when feeding on *Populus tremula* than on *P. alba*. Finally, honeydew production and honeydew composition may vary within a particular species on a definite host plant in response to changes in the aphid’s interaction with ants (Del-Claro & Oliveira, 1993; Fischer et al., 2001).

The black bean aphid, *A. fabae*, is regularly tended by honeydew-collecting ants and benefits from this mutualism. The life cycle of the host-alternating *A. fabae* involves seasonal migration between unrelated summer (herbaceous or secondary) and winter (woody or primary) host plants. In autumn, winged females are produced (gynoparae) which locate the primary host, the spindle tree *Evonymus europaeus* and deposit sexual females that develop, mate and lay eggs before leaf fall. During the summer, *A. fabae* is polyphagous. For example, summer forms (virginoparae) of the sub-species *A. f. fabae* are able to colonize approximately 100 plant species (Powell & Hardie, 2001). Under field conditions *A. f. fabae* is not found on the winter host plant during the summer, but they will feed and reproduce on spindle trees in laboratory tests (Powell & Hardie, 2000). Another subspecies of *A. fabae*, *A. f. cirsiicanthoides* colonizes the creeping thistle, *Cirsium arvense*, as a summer host, whereas the non-host alternating subspecies *A. f. evonymi* is found on spindle trees throughout the whole year.

This study investigates whether differences exist in the honeydew composition of *A. fabae* subspecies feeding on various hosts and whether such differences in honeydew composition may result in differences in ant-attendance.

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First, we examined the honeydew production and the honeydew sugar composition of *A. f. fabae* on various host plants, a primary host, *E. europaeus* and three herbaceous summer hosts, the goosefoot *Chenopodium album* (Chenopodiaceae), the tansy *Tanacetum vulgare* (Asteraceae), and the broad bean *Vicia faba* (Fabaceae). Second, we compared the sugar composition of the honeydew from three *A. fabae* subspecies, *A. f. fabae*, *A. f. cirsiiacanthoides*, and *A. f. evonymi* feeding on their respective summer hosts, and third we investigated the relationship between honeydew composition in *A. fabae* ssp. colonies on four herbaceous summer host plants and ant-attendance. Finally, we analyzed the sugar composition of the honeydew during the life cycle of *A. f. fabae* on its winter host *E. europaeus*.

**MATERIAL AND METHODS**

**Plant and insect material**

*Chenopodium album* plants were grown from seeds in plant pots (diameter 25 cm) in a defined garden soil (obtained from Bayreuth Botanical Garden) to standardize plant quality. The pots were kept in a climate chamber (20°C, 16L:8D, 60% relative humidity, 8000 lux from the ceiling) where they grew to a height of about 20 cm within two to three weeks. *Vicia faba* (cv minor) plants were grown from seeds as described earlier (Engel et al., 2001). All plants were about 20 cm tall when used in the experiments. The two perennial herbs *Cirsium arvense* plants (minimum height of 30 cm) and tansy (*Tanacetum vulgare*) were obtained from rhizomes dug out from ruderal areas at the University campus and potted in garden soil (diameter of pots 30 cm). We only used plants that were about to flower for the experiments. All plants were 40 to 50 cm tall (Fischer et al., 2002). Young spindle trees (*Evonymus europaeus*) (50 to 70 cm tall) were obtained from the field (Neunkirchen, 10 km east of Bayreuth) and potted in defined soil.

To obtain an *A. f. fabae* stock culture, one large colony (> 100 individuals) was collected in the field. Twenty-five to 30 individuals of this colony were transferred separately to potted tansy plants in a climate chamber (for conditions, see above). For our experiments, aphids from this stock culture were transferred to their various host plants and used when the respective colonies reached a size of 50 to 100 individuals. In the case of *A. f. cirsii-acanthoides*, aphids were collected from *C. arvense* in the field and directly transferred to the experimental plants. The potted plants were kept at a sunny site in earth-filled planting pots (1 m × 1 m × 0.5 m; 1 × b × h) each of which contained a nest of the common garden ant *Lasius niger* (approx. colony size: 1000–2000 workers). In each pot, the plants were placed close to the nest in waterfilled bowls to prevent uncontrolled ant-attendance. The experiments were carried out between end of June and mid-August.

For the determination of the honeydew composition of *A. f. fabae* on *E. europaeus* during their life cycle and for the studies of *A. f. evonymi* on *E. europaeus*, established spindle trees at the university campus (approx. 3 m high) were used. The life cycle experiments were carried out from mid-April to mid-May (fungatrix, fungatrigenia, winged females/alates) and in October (alate gonoparae, oviparae).

**Measuring the intensity of ant-attendance**

For these experiments 15 potted plants each of *C. alba*, *C. arvense*, and *T. vulgare*, and 7 pots (10 at the start of the experiment) with *V. faba* were used. Five plants of one species were placed in a planting pot containing an ant nest. Three *V. faba* plants were destroyed by phytophagous insects during the experiment. This trial was excluded from the evaluation of the relative ant-attendance but could be included into the data used for Table 1. Thus, we used in total 10 large sized planting pots with one ant colony in each pot. Workers of *L. niger* had access to each plant (and thus to the aphids) through an arrangement of wooden sticks. By this means, we could control for ant-attendance at each individual plant.

At the beginning of the experiment each aphid colony was standardized to 20 individuals. For each colony we counted the number of aphids as well as all ant workers which left the aphid colony (on the wooden sticks) during a 10-min interval every day (at the same daytime) over a period of three weeks. For evaluation, we separated this time interval: The first period (approx. one week) included the time before swarming of ant nests (i.e., ant nests contained sexuals and thus had a high demand for honeydew), the second period the time after swarming (i.e., with a reduced demand for honeydew). For comparison of host plants, we choose only the days 2–5 before swarming and 6–9 after swarming of each colony to correct for a potential bias during pre-swarming, swarming and post-swarming days.

Although we started with a standardized colony size, colonies grew differently during the nine days experimental period. To obtain a parameter for the intensity of ant-attendance that was independent of aphid colony size, we divided the number of ants by aphid colony sizes (ants per aphid = relative ant-attendance) as justified by the specific positive linear correlation of ant-attendance (Fischer et al., 2001) (see Results).

**Quantification of honeydew production**

We measured the honeydew collected by the ants by comparing the weight of ant workers with empty and filled gasters (Herzig, 1937). For experimental details see Fischer et al. (2001, 2002). Since all excreted honeydew droplets were collected by *L. niger* workers, we estimated the honeydew production of an individual aphid per hour using the following formula: [(average weight of ants with filled gasters leaving *A. fabae* – average weight of ants with empty gasters) × average number of ants leaving the colony of *A. fabae*)/number of aphids per *A. fabae* colony]. Ants with empty gasters were collected when they approached the aphid colony via a stick. For each colony, we collected prior to the swarming period 20 ants with empty gaster and 20 ants with filled gaster. During this time period, the amount of honeydew collected in the gather provides a good estimate of the maximum potential honeydew production (see Fischer et al., 2001).

**Honeydew collection**

Honeydew was collected twice a week directly from the anus using a micropipette (volume 0.5 or 1 μl). After collection, samples were capped with Parafilm and stored at −20°C until analysis. For a comparison between plants, 10–15 individual samples per plant species were analysed, collected from fourth instar larvae and adult aphids which provided the highest amount of honeydew. For a comparison between various morphs on *E. europaeus*, we obtained 5–12 samples per morph. Since the honeydew sugar composition does not differ significantly between aphid age classes (Fischer et al., 2001), a random selection of only two age classes should not bias the results.

**Analysis of honeydew sugar composition**

The honeydew was analyzed using high-pressure liquid chromatography (HPLC). A Carbo PakPA100, 4 × 250 mm column (Dionex, Sunnyvale, CA) was used with a 0.5 M NaOH gradient of 3–70% in Milli-Q water for 30 min at a flow rate of
For the statistical evaluation of the data we followed procedures outlined by Sokal & Rohlf (1981) by using SPSS/PC+ 3.3 and WinSTAT 3.1. In Mann-Whitney-U tests, the Bonferroni correction was considered where necessary.

RESULTS

Honeydew production of *A. f. fabae* on various host plants

The honeydew production of *A. f. fabae* differed considerably between host plants (Fig. 1). Aphids feeding on the two annual plants *V. faba* and *C. album* produced less than half the amount of honeydew per aphid per hour than conspecifics on the perennial *T. vulgare*. Honeydew production of *A. f. fabae* virginoparae on the winter host *E. europaeus* was also high (135.5 µg per aphid per hour).

Sugar concentration and sugar composition of the honeydew of *A. f. fabae* on various host plants

Up to ten sugars could be identified in the honeydew of *A. f. fabae* on the various host plants. Honeydew was generally dominated by the trisaccharide melezitose (Fig. 2), the disaccharide trehalose, and the monosaccharides glucose and fructose. Sucrose, maltose and erlose were present in low concentrations in all honeydew samples, whereas traces of xylose (*V. faba*), turanose (*C. album*) and raffinose (*E. europaeus*) were detected only in single samples. The level of melezitose by far exceeded that of other sugars except for *A. f. fabae* on *E. europaeus* where the honeydew contained approximately equal amounts of melezitose, trehalose, glucose and fructose. The total sugar concentration of the honeydew (Fig. 3) did not differ significantly between various hosts (27 to 42 µg sugar per µl honeydew).

Honeydew production and sugar composition of the honeydew for three subspecies of *A. fabae*

*A. f. cirsiiacanthoides* feeding on *C. arvense* produced an average of 151.5 ± 35.5 µg honeydew per aphid per hour (means ± SE, n = 10; not shown), whereas *A. f. fabae* feeding on *V. faba* produced ca. 55 µg . aphid⁻¹ . h⁻¹ only (see Fig. 1). The honeydew production of *A. f. evoynmi* on *E. europaeus* was not measured. The honeydew of *A. f. cirsiiacanthoides* contained a significantly higher amount of total sugars (ca. 70 µg . µl⁻¹) than the honeydew of *A. f. fabae* on its summer host (Fig. 4; see also Fig. 3). A similar concentration of total sugars as for *A. f. fabae* on its summer host was present in the honeydew of
A. f. evonymi on spindle trees. The honeydew of A. f. cirsiiacanthoides on the creeping thistle contained the highest proportion of melezitose (80% of total sugars) (Fig. 5), whereas in the honeydew of A. f. evonymi fructose was dominant (34%). Moreover, this honeydew contained considerable amounts of turanose and raffinose, which were not detected in either of the other subspecies feeding on herbaceous plants (but found in a sample of A. f. fabae on E. europaeus, see above).

**Ant-attendance of A. fabae ssp. on herbaceous summer hosts by L. niger**

In a first set of experiments we showed that in three of the four host plants a positive correlation existed between aphid colony size and the number of attending ants (Table 1). Within our range of colony sizes the relationships were linear. Since ants began to swarm during the observation period (end of July), the observation period had to be divided into two parts: Pre-swarming at day 2–5 and swarming at days 6–9 (Table 2). Before swarming, significant differences in ant-attendance were observed between aphids on the three host plants with highest intensity of visiting ants for the A. f. cirsiiacanthoides colonies on C. arvense and least intensity of ant-attendance for A. f. fabae on T. vulgare. During swarming aphids on C. arvense were still more attractive for ants than on the other two host plants.

### Table 1. Correlation of ant-attendance (L. niger) with the number of aphids in A. fabae ssp. colonies on four herbaceous summer host plants; n, number of observations.

| Plant species | r | n | p    |
|---------------|---|---|------|
| C. album      | 0.511 | 90 | <0.001 |
| T. vulgare    | 0.638 | 87 | <0.001 |
| C. arvense    | 0.636 | 90 | <0.001 |
| V. faba       | 0.339 | 21 | 0.132 |

**Sugar composition of the honeydew from various generations of A. f. fabae**

Total sugar concentration in the honeydew of A. f. fabae on E. europaeus changed significantly during the experimental period (Fig. 6). Sugar concentration was very low (< 10 µg µl⁻¹) in the fundatrix generation which had hatched from overwintering eggs in April, but increased to 25–35 µg µl⁻¹ in the following two generations (fundatrigenia, alates; April to mid-May). Individuals of the third winged generation usually migrate to the summer host. Alate gynoparae that had returned from the summer host at the end of the vegetation period (October) again showed a low amount of sugars in the honeydew (ca. 10 µg µl⁻¹), but another increase could be observed in the oviparae by the end of October. The percent distribution of the sugars in the honeydew did not significantly change over the generations (Table 3) except that the honeydew of the fundatrigenia contained about twice the amount of melezitose as in the other generations (Fig. 7).

### DISCUSSION

The mutualism between aphids and ants is influenced by a number of biotic factors. For the ant, the quantity and the quality of the supplied honeydew are important (Bristow, 1984), but also the predictability of the honeydew source (Noe & Hammerstein, 1994). The quantity of the honeydew produced by aphids as well as the honeydew composition varies among species. Within given species, the rate of honeydew production and the sugar composition of the honeydew may vary along their developmental (seasonal) cycle and between age classes (Hertel & Kunkel, 1977; Fischer et al., 2002), but also with variation in the host plant (Hendrix et al., 1992; Fischer & Shingleton, 2001). Furthermore, honeydew production and honeydew composition change in response to interactions with ants (Del-Claro & Oliveira, 1993; Fischer et al., 2001).
Ants respond most intensively to honeydew containing high amounts of the trisaccharide melezitose (Kiss, 1981; Völkl et al., 1999). Some aphids, such as *Metopeurum fuscoviride* on tansy, excrete honeydew that contains up to 70% melezitose, and such species are strongly attended by ants (Fischer et al., 2001, 2002). Other species, for example *Macroplio Nothaplantarctea* on tansy, contain virtually no melezitose in their honeydew, and are not usually tended by ants (Hendrix et al., 1992; Völkl et al., 1999). In a recent study on the hierarchy of ant-attendance in eight species of aphids feeding on tansy (Woodring et al., 2004) it was demonstrated that a combination of the sugar richness of the honeydew excreted by the aphid (rate of honeydew production × total sugar concentration in the honeydew) together with a high content of the attractant sugar melezitose determines the extent of ant-attendance. There was no evidence that any single amino acid or group of amino acids in the honeydew act as an attractant for ant-attendance in these aphids.

In *A. f. fabae* feeding on four different herbaceous host plants, honeydew production differed significantly. For example, the amount of honeydew produced by *A. f. fabae* feeding on tansy was more than twice (ca. 110 µg per aphid h⁻¹) as high as when feeding on goosefoot, *C. album* (ca. 40 µg per aphid h⁻¹). The honeydew of *A. f. fabae* on tansy also contained a higher amount of total sugars, and a slightly higher proportion of melezitose, than when feeding on *C. album*. Another subspecies of *A. fabae*, *A. f. cirsiiacanthoides*, which occurs on the creeping thistle, *C. arvense*, during the summer, produced as much as 150 µg honeydew per aphid per hour, and this honeydew contained by far the highest amount of total sugars (ca. 67 µg/µl) as well as the highest portion of melezitose (80%). The calculated value for richness of honeydew was 8.7, compared to 3.9 for *A. f. fabae* on tansy, and only 1.0 for *A. f. fabae* on goosefoot. The results confirm our earlier hypothesis that the sugar richness of the honeydew along with the presence of melezitose are the critical factors in determining the extent of ant-attendance. Indeed, in the present study the highest intensity of visiting ants was observed for *A. f. cirsiiacanthoides* colonies on *C. arvense*. However, richness of honeydew does not explain the differences between goosefoot and tansy. Although having a lower richness of honeydew, aphids on goosefoot were less attended than aphids on tansy. We hypothesize that differences in plant structure might contribute to this finding. Ants have much better access to the aphid colonies feeding on goosefoot leaves and inflorescences than on the stem of tansy. *A. f. evonymi*, which is found on the woody spindle tree all year round, produced a honeydew which was low in total sugars and contained only low amount of melezitose with fructose being the dominant sugar. The sugar quality did not change during the season (Fischer, 2001). Ants were seldom observed visiting *A. f. evonymi* (Börner, 1952, Völkl & Fischer, unpubl.).

There are also important ecological implications. First, a higher ant-attendance should provide a better protection against natural enemies such as predators and parasitoids (Fischer et al., 2001). Second, there might exist indirect effects between colonies of the same species on different host plant species. Considering that ants compete for mutualists (e.g., Del-Claro & Oliveira, 1993; Fischer et al., 2001) we may hypothesize that aphid colonies on creeping thistle might distract ants from neighbouring aphid colonies on tansy or goosefoot which often grow in the same habitat. Again, these colonies may suffer from increased predation and parasitism (Fischer et al., 2001) due to reduced ant guarding. Thus, they may have lower

### Table 3. Kruskal-Wallis-ANOVA (K-W $\chi^2$) for the proportion of sugars in the honeydew from various generations of *A. f. fabae* on their winter host, *E. europaeus*. Means sharing the same letter do not differ significantly at $p < 0.05$ (Mann-Whitney-U test).

| Sugars | $\chi^2$ | df | p   |
|--------|----------|----|-----|
| % Tre  | 5.21     | 4  | 0.266|
| % Glu  | 1.51     | 4  | 0.825|
| % Fru  | 3.05     | 4  | 0.550|
| % Suc  | 1.22     | 4  | 0.875|
| % Mel  | 9.09     | 4  | 0.050|
| % Mal  | 2.05     | 4  | 0.727|
| % Eri  | 8.40     | 4  | 0.078|

*Fig. 6. Total sugar concentration (in µg sugar per µl honeydew) (mean ± SE) in the honeydew of various generations of *A. f. fabae* on their winter host, *E. europaeus*. Means sharing the same letter do not differ significantly at $p < 0.05$ (Mann-Whitney-U test).*

*Fig. 7. Proportion (in % of total content) (mean ± SE) of the identified sugars in the honeydew of various generations of *A. f. fabae* on *E. europaeus*. Means sharing the same letter within the trisaccharide melezitose do not differ significantly at $p < 0.05$ (Mann-Whitney-U test).*
overall fitness since they will be expected to produce fewer sexuals in autumn.

In another experiment, we studied the honeydew quality in various generations of *A. f. fabae* on their winter host, *E. europaeus*. In contrast to *A. f. evonymi* feeding on the spindle tree, honeydew of *A. f. fabae* showed drastic changes in sugar content as well as in sugar composition during the season. The honeydew of the fundatrigenia and alates in late spring contained significantly higher total sugar concentrations than of the other generations and the honeydew of the fundatrigenia also was the richest in melezitose. These two generations fed on the developing leaves, and thus we might expect that plant metabolism affects even honeydew concentration. This assumption is supported by the differences that plant metabolism affects even honeydew concentration.

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