A method to feed individual bees (Hymenoptera: Apiformes) known amounts of pesticides

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Abstract – We devised a simple method (“flower”) to feed bees individually, and compared it with two other methods commonly used (“film canister” and “glass vial”). We tested the three methods on two solitary species, Osmia lignaria and Megachile rotundata, and one social species, Apis mellifera, under four different light regimes (natural, artificial, plant growth and darkness). The flower method was the most effective for all three bee species: 90–95% of the bees fed under natural light, 80–95% under artificial light, 75–100% under plant growth light, and 45–70% in darkness. Percent success was 0–50% with the film canister method, and 0–60% with the glass vial method. The flower method may allow more comprehensive future evaluation of pesticide effects on bees.

oral toxicity test / individual feeding / Osmia lignaria / Apis mellifera / Megachile rotundata

1. INTRODUCTION

Bee populations are often exposed to pesticide treatments in both natural and agricultural ecosystems (Kevan, 1975; Crane and Walker, 1983; Johansen and Mayer, 1990; Peach et al., 1993), and insecticide poisoning is considered one of the main causes of bee population declines worldwide (Dias and Raw, 1999; Tepedino and Ginsberg, 2000). Given the ecological and economic importance of bees as pollinators of wild flowers and cultivated plants (Kevan, 1991; Southwick and Southwick, 1992), it is surprising that our knowledge on bee toxicity is so fragmentary and mostly restricted to one species, the honey bee, Apis mellifera L. (Apidae) (Johansen and Mayer, 1990). Information on pesticide toxicity to non-Apis bees is dismally scarce, and limited to a handful of species managed for crop pollination (review in Taséi, 2002). A common first step in bee toxicity studies is the establishment of LD50 (median lethal dose) values, i.e. the dose, expressed in µg of active ingredient per insect, inducing 50% mortality following application (contact or oral) of measured amounts of active ingredients or commercial pesticide formulations. The methods used for determining LD50 values on A. mellifera are defined by official guidelines, both in Europe and in the USA (US EPA, 1996; OEPP/EPPO, 2001; Cluzeau, 2002). Methods used to study contact toxicity in A. mellifera can be easily applied to other species. However, in studies of oral toxicity in A. mellifera a common feeder is provided to a group of workers assuming that, through trophallaxis, all individuals will receive similar doses of test solution (OEPP/EPPO, 2001). Group feeding is not applicable to most other bee species, which do not perform trophallaxis and thus require individual feeding. In fact,
oral tests on non-\textit{Apis} bees are rarely conducted because individual feeding is considered time consuming (Felton et al., 1986).

In 2002, we initiated a study on the effect of pesticide sprays on the solitary bee \textit{Osmia lignaria} Say (Megachilidae). We tried to feed individual \textit{O. lignaria} known amounts of sugar solution using some of the methods available from the literature (Taséi, 2002), but our success rate (percent individuals that would feed) was very low. Thus, we devised a new method, which we tested in comparison to the two most commonly used methods available (Johansen et al., 1984; van der Steen et al., 1996; Bortolotti et al., 2002; Patetta et al., 2002). All three methods were tested on two solitary bee species \textit{O. lignaria} and \textit{Megachile rotundata} (Fabricius) (Megachilidae), and the social bee \textit{A. mellifera}. All three species are used as crop pollinators (Richards, 1984; Free, 1993; Bosch and Kemp, 2001), and thus are frequently exposed to pesticides.

2. MATERIALS AND METHODS

\textit{Apis mellifera} workers were captured in the morning at the entrance of a hive and brought to the laboratory, where they were chilled for a maximum of 30 min at 4 °C prior to being assigned to the different feeding methods. \textit{Osmia lignaria} and \textit{M. rotundata} wintering females within their cocoons were incubated until emergence at 25 °C and 29 ºC, respectively. Upon emergence, females were transferred to a holding cage (40 × 30 × 30 cm) to allow them to deposit the meconium. Females were starved overnight and then assigned to the different feeding methods. No chilling was necessary for either \textit{O. lignaria} or \textit{M. rotundata}.

We used the following three individual feeding methods: (1) Film canister method (after van der Steen et al., 1996; Bortolotti et al., 2002; Patetta et al., 2002). Bees were individually transferred to black film canisters (3 cm diameter, 5 cm height) with a small hole drilled on the side, near the base. The test solution was pipetted onto a microscope slide next to the hole. (2) Glass vial method (after Johansen et al., 1984). Bees were individually transferred to 12-mL glass vials with plastic snap caps. The test solution was injected into a segment of plastic tubing (length 15 mm, inside diameter 2.9 mm), fitted snugly into the plastic snap caps. (3) Flower method (Fig. 1). A tiny plastic ampoule (inside diameter 2 mm, outside diameter 3 mm, height 5 mm) was inserted into the calyx of a flower, whose reproductive column had been previously removed with a pair of forceps. The plastic ampoule was built by approaching one tip of a 5-mm section of polyethylene tubing to a flame or some other heat source. The heat caused the polyethylene tip to melt, sealing the tubing orifice. Because the polyethylene never actually touched the heat source, no charring occurred, and ampoules had no obvious odor. The test solution was pipetted into the ampoule. Flowers and bees were individually housed in ice cream cups made of waxed cardboard (8 cm diameter, 5 cm height) covered with a plastic Petri dish lid. Flowers were positioned on an inverted glass vial stopper. A wire mesh screen insert (mesh size 2 × 1 mm) in the Petri dish lid provided adequate aeration. To facilitate flower manipulation, we used large, actinomorphic flowers with open corollas. In preliminary trials, we tested two or
more flowers per bee species, including cherry (Prunus avium L.), apple (Malus domestica Borkh), morning glory (Convolvulus arvensis L.) and periwinkle (Vinca minor L.). All flowers in these preliminary trials yielded similar results. In the actual experiments we used cherry flowers for the early-flying O. lignaria and morning glory flowers for the later-flying M. rotundata and A. mellifera. A trained technician could prepare 100 test flowers in 40 minutes.

In all three methods, a 10-µL-drop of test solution (25% volume of sucrose to 75% volume of water) was offered to the bee for one hour. We tested each method under four different light regimes: (1) Natural light. Feeding units placed outdoors (on sunny to partially-cloudy days); (2) Artificial light 1. Feeding units placed in an incubator with two 15W Cool White Sylvania® fluorescent tubes (wavelength peaks at 405, 430, 545 and 575 nm) placed 15 cm above the feeding units; (3) Artificial light 2. Feeding units placed in an incubator with two 20W Gro-Lux/Aquarium Standard Sylvania® fluorescent tubes (usually used to help plant growth; wavelength peaks at 460, 545, 630 and 660 nm) placed 15 cm above the feeding units; (4) Darkness. Feeding units placed in an incubator with no light. We tested darkness because current guidelines for group feeding toxicology assays are usually conducted in darkness (Arzone and Vidano, 1980; OEPP/EPPO, 2001). Temperatures during all tests (including those outdoors) were 22 ± 2 °C for O. lignaria and 25 ± 2 °C for M. rotundata and A. mellifera. We used a lower temperature for O. lignaria because this spring-flying bee is active at lower temperatures than M. rotundata or A. mellifera (Burril and Dietz, 1981; Lerer et al., 1982; Bosch and Kemp, 2001).

Sample sizes were 20 individuals of each species (workers in A. mellifera, females in O. lignaria and M. rotundata) per feeding method/light regime. We analyzed the proportion of feeding bees as a function of feeding method and light regime. Because different bee species were tested under different temperatures and with different flowers, we analyzed the results of each species separately. Due to the number of cells with frequencies of < 5 for one of the species (O. lignaria, Tab. I), we attempted to fit an exact Logit model using LogXact (Version 5.0). However, the computational size was too large for this exact method to work (even on a sufficiently equipped computer). In addition, the complete separation of the data prevented the computation of Likelihood Ratio and Wald test statistics. Therefore, because the structure of the O. lignaria data prevented the use of an exact test, we used an asymptotic Logit model (GENMOD procedure, SAS Institute Inc., 1989; Allison, 1999). Because of the pronounced pattern of the results (O. lignaria was the species for which differences among feeding methods were most extreme), it is unlikely that the use of this statistical analysis would compromise our interpretations.

3. RESULTS

The flower method was, by far, the most effective for all three bee species (Tab. I): 90–95% of the bees fed under natural light, 80–95% under Cool White fluorescent tubes, 75–100% under Gro-Lux fluorescent tubes, and 45–70% in darkness. All bees that fed with the flower method consumed all 10 µL of the test solution. Percent success was 0–60% with
Table II. Results of Logit analysis of feeding success in Osmia lignaria females, Megachile rotundata females, and Apis mellifera workers offered 10 µL of sugar solution using three feeding methods under four light regimes.

<table>
<thead>
<tr>
<th>Species</th>
<th>Source</th>
<th>df</th>
<th>X²</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>O. lignaria</td>
<td>method</td>
<td>2</td>
<td>166.2</td>
<td>&lt; 0.0001</td>
</tr>
<tr>
<td></td>
<td>light</td>
<td>3</td>
<td>6.87</td>
<td>0.076</td>
</tr>
<tr>
<td></td>
<td>method × light</td>
<td>6</td>
<td>7.10</td>
<td>0.312</td>
</tr>
<tr>
<td>M. rotundata</td>
<td>method</td>
<td>2</td>
<td>107.66</td>
<td>&lt; 0.0001</td>
</tr>
<tr>
<td></td>
<td>light</td>
<td>3</td>
<td>5.59</td>
<td>0.133</td>
</tr>
<tr>
<td></td>
<td>method × light</td>
<td>6</td>
<td>14.36</td>
<td>0.026</td>
</tr>
<tr>
<td>A. mellifera</td>
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<td>39.97</td>
<td>&lt; 0.0001</td>
</tr>
<tr>
<td></td>
<td>light</td>
<td>3</td>
<td>44.16</td>
<td>&lt; 0.0001</td>
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<tr>
<td></td>
<td>method × light</td>
<td>6</td>
<td>11.29</td>
<td>0.080</td>
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</table>

It might be argued that extending exposure time of bees to the feeding units could improve success rates. Johansen et al. (1984) left bees in the feeding glass vials for three hours, but found that most bees fed within 1 hour. We observed that, with all three feeding methods, bees would usually feed within 30 minutes or not at all. Furthermore, current guidelines discourage individual confinement of A. mellifera for more than one hour (Felton et al., 1986; OEPP/EPPO, 2001). To standardize procedures, it would be better to use similar conditions with all bee species. Because artificial and natural light tests yielded similar results, we recommend the use of artificial light.

A common problem in bee toxicology studies is the reluctance of bees to feed on solutions containing chemicals. A recent review points out that repellency is less of a problem in field trials than in laboratory tests, probably because the attractiveness of the flowers overrides the repellency of the chemical application (Thompson, 2003; see also Naumann et al., 1994). Using the flower method, chemicals such as Captan and Neem known to be repellent when dissolved in sucrose (Solomon and Hooker, 1989; Naumann et al., 1994) were readily ingested by A. mellifera and O. lignaria, even at near-saturation concentrations (E.L., unpublished).

With the flower method, a source of flowers not treated with chemicals needs to be secured throughout the study. Two of the flowers that we tried, V. minor and C. arvensis were available in large numbers and for long periods (more than two months) in gardens in spring and summer, respectively. In preliminary, off-season, trials we used flowers grown in greenhouses. The flower method is more labor-intensive than the other two methods tested. However, because it is much more efficient, it actually saves time since fewer feeding units need to be prepared and fewer bees need to be collected and/or reared. This is especially important when studying species that cannot be obtained in large numbers. As mentioned,
as many as 100 flowers can be prepared in 40 minutes. To further save time, we tried to deposit the droplet of sugar solution directly on the flower corolla (without the polyethylene ampoule). However, at least when systemic pesticides were added to the sugar solution, the droplet was partially absorbed by the flower tissues. In a subsequent study, the flower method (with the ampoule and under artificial light) was used to assess the toxicity of several pesticides to O. lignaria and A. mellifera. Success (feeding) rates were 97.7% for the first species, and 88.2% for the second (n = 871 and 755, respectively) (E.L., unpublished data). In conclusion, the flower method is a simple, highly effective procedure that could help simplify and standardize oral toxicity tests on a diversity of bee species in the laboratory. To further standardize procedures, future studies should investigate the possibility of substituting natural flowers by artificial flowers properly colored and scented.

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Résumé – Méthode pour administrer individuellement aux abeilles (Hymenoptera, Apiformes) une quantité définie de pesticides. Les études en laboratoire sur l’action des pesticides sur les abeilles écartent souvent les tests de toxicité orale à cause de la difficulté d’administrer des quantités définies de solutions à tester. Les études sur l’Abeille domestique (Apis mellifera L.) reposent sur des méthodes de nourrissement de groupe et considèrent que des quantités sensiblement égales de solution sont ingérées par chaque ouvrière à travers les échanges de nourriture (trophallaxie). Néanmoins la plupart des abeilles (s.l.) ne pratiquent pas la trophallaxie. Nous avons mis au point une méthode simple (la méthode « de la fleur ») pour nourrir des abeilles individuellement et l’avons comparée à deux autres méthodes couramment employées (les méthodes de « la boîte pour pellicule photo » et celle du « flacon en verre »). Nous avons testé les 3 méthodes sur 2 abeilles solitaires, Osmia lignaria Say et Megachile rotundata (Fabricius) (Megachilidae), et sur une espèce sociale, A. mellifera, sous 4 régimes de lumière différents (lumière naturelle, tubes fluorescents blanc froid, tubes fluorescents Gro-Lux et obscurité). Dans la méthode de la fleur, les abeilles étaient confinées individuellement dans une coupe à glace. On leur donnait une fleur dont le pistil avait été remplacé par une fine ampoule dans laquelle la solution à tester était pipetée (Fig. 1). La méthode de la fleur a été la plus efficace chez les 3 espèces d’abeilles (Tab. I), quel que soit le type d’éclairage : 90–95 % en lumière naturelle, 80–95 % avec tubes fluorescents blanc froid, 75–100 % avec tubes fluorescents Gro-Lux et 45–70 % en obscurité. Le pourcentage de succès a été compris entre 0 et 50 % avec la méthode de la boîte pour pellicule photo et entre 0 et 60 % avec la méthode du flacon de verre. La préparation des unités de nourrissement demandait plus de travail dans la méthode de la fleur mais avec de la pratique, on peut préparer 100 fleurs en 40 min. En raison de sa grande efficacité, la méthode peut en fait économiser du temps (en termes de récolte d’abeilles et de préparation) et du matériel animal.

Osmia lignaria / Apis mellifera / Megachile rotundata / test de laboratoire / toxicité orale / nourrissement individuel

oraler Toxizitätstest / individuelle Fütterung / Osmia lignaria / Apis mellifera / Megachile rotundata

REFERENCES


