Foraging behavior and prey interactions by a guild of predators on various lifestages of Bemisia tabaci

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Foraging behavior and prey interactions by a guild of predators on various lifestages of *Bemisia tabaci*

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**Abstract**

The sweetpotato whitefly, *Bemisia tabaci* (Gennadius) is fed on by a wide variety of generalist predators, but there is little information on these predator-prey interactions. A laboratory investigation was conducted to quantify the foraging behavior of the adults of five common whitefly predators presented with a surfeit of whitefly eggs, nymphs, and adults. The beetles, *Hippodamia convergens* Guérin-Méneville and *Collops vittatus* (Say) fed mostly on whitefly eggs, but readily and rapidly preyed on all of the whitefly lifestages. The true bugs, *Geocoris punctipes* (Say) and *Orius tristicolor* (Say) preyed almost exclusively on adult whiteflies, while *Lygus hesperus* Knight preyed almost exclusively on nymphs. The true bugs had much longer prey handling times than the beetles and spent much more of their time feeding (35-42%) than the beetles (6-7%). These results indicate that generalist predators vary significantly in their interaction with this host, and that foraging behavior should be considered during development of a predator-based biological control program for *B. tabaci*.

**Keywords:** Predator-prey interactions, prey choice, sequence analysis, biological control, insect behavior

**Abbreviation:**

ELISA enzyme-linked immunosorbent assay

**Introduction**

The sweetpotato whitefly, *Bemisia tabaci* (Gennadius), strain B (= silverleaf whitefly, *B. argentifolii* Bellows and Perring) and other whitefly species are worldwide pests of many ornamental, greenhouse, and agricultural crops (Perring 1996; Naranjo and Ellsworth 2001). An enormous effort has been dedicated to studying the natural enemies of these pests (Gerling and Mayer 1996; Naranjo and Ellsworth 2001). An enormous effort has been dedicated to studying the natural enemies of these pests (Gerling and Mayer 1996; Gerling et al. 2001). To date, most of the research on whitefly natural enemies has focused on evaluating the behavior and efficacy of parasitoids (Gerling 1986, 1992; Gerling et al. 2001). In contrast, while it is known that many predator species feed on whiteflies (Gerling 1990; Cocker 1994; Breene et al. 1994; Nordlund and Legaspi 1996; Gerling et al. 2001), there is little published information on their prey preferences or specific foraging patterns. For example, with the exception of a few lacewing species (Or and Gerling 1985; Gerling 1986, 1990; Butler and Henneberry 1988; Kapadia and Puri 1989; Breene et al. 1992, 1994; Legaspi et al. 1994), very little is known about any predator’s prey preference, consumption rate, or handling time for the various whitefly lifestages (Nordlund and Legaspi 1996; Hagler 2002).

Progress toward determining these basic foraging characteristics has been slow, due in part to the difficulty of observing whitefly predation in the field. Whiteflies and many of their predators are small, elusive, and cryptic. Additionally, adult whiteflies and their predators are highly mobile. As a consequence, direct field observations of whitefly predation are tedious, time consuming, and can be disruptive to the normal foraging process (personal observation). Furthermore, continuous observation of an individual predator over an extended period of time is not practical in the field for those species that disperse readily and forage on the underside of the leaf. Because of these technical challenges, indirect techniques are used to examine the influence of whitefly predators. For example, we developed a whitefly-specific enzyme-linked immunosorbent assay (ELISA) for identifying whitefly remains in the guts of field-collected predators. However, this assay had three limitations: (1) the whitefly “egg-specific” monoclonal antibody used in the assay did not differentiate between an egg and a gravid female meal, (2) it did not detect predation on whitefly nymphs and adult males, and (3) the assay was not quantifiable (Hagler et al. 1993). As a consequence, the gut content immunoassay could not detect how many whiteflies were consumed or which lifestage were consumed (i.e., an egg, adult female, or a mixture of both).

This study was conducted to learn more about the
interactions of whitefly predators with their host. We directly observed the behaviors of 130 individuals, representing five predator species, in feeding arenas containing an abundance of whitefly eggs, nymphs, and adults on a cotton leaf disk. The whitefly predators examined were: *Hippodamia convergens* Guérin-Méneville (Coleoptera: Coccinellidae), *Collops vittatus* (Say) (Coleoptera: Melyridae), *Geocoris punctipes* (Say) (Heteroptera: Lygaeidae), *Orius tristicolor* (Say) (Heteroptera: Anthocoridae), and *Lygus hesperus* Knight (Heteroptera: Miridae). These predators were selected for this study because assays of predator gut content by ELISA have shown that they readily prey on whitefly (Hagler and Naranjo 1994a,b). Moreover, *L. hesperus* was included because it has been shown that this omnivorous pest is also a prolific whitefly predator (Hagler and Naranjo 1994a). Our specific objectives were to: (1) determine if these predators prefer a particular whitefly life stage, (2) determine the amount of and rate of whitefly consumption, and (3) compare predator foraging patterns between species.

Materials and Methods

Predators

Adult predators of the five test species were collected with 36.0-cm diameter sweep nets from cotton, *Gossypium hirsutum* L. fields located at The University of Arizona’s Agricultural Research Station, Maricopa, Arizona. The entire contents of the sweep net samples (e.g., arthropods and plant debris) were placed in a large communal rearing cage containing a potted cotton plant and held at 27°C in our laboratory. The predators were allowed to feed freely on any available prey collected in the sweep nets. The day before behavioral observations, a single adult predator was removed from the communal rearing container, placed individually into a clean Petri dish (9.0-cm diameter), and provided with only water.

Feeding arena

Single ‘Delta Pine 5415’ cotton plants were grown in 15.2-cm diameter pots in a greenhouse. Plants that were four to five weeks old were infested with adult whiteflies each week. When the plants were approximately eight to nine weeks old, a single leaf was removed from a plant and cut to fit exactly into the bottom of a 3.5-cm diameter plastic Petri dish (the feeding arena). We selected leaves that contained numerous whitefly eggs and nymphs. The number of whitefly eggs and nymphs were counted on each leaf disk and placed abaxial side up into the bottom of the feeding arena. Adult whiteflies were then introduced into the arena and the Petri dish lid was placed over the top of the arena. A typical feeding arena contained a 3.5-cm diameter cotton leaf disk infested with an average (± SD) of 837.2 ± 67.8, 704.4 ± 60.0 and 43.1 ± 1.5 eggs, nymphs, and adults (males and females at a 1:1 sex ratio), respectively.

Behavioral observations

A single, randomly selected predator (each predator species had an approximate sex ratio of 1:1), that had been held overnight without food was placed into the feeding arena and continuously monitored for 1 h under a dissecting microscope. Preliminary observations of the predator’s behavior revealed several distinct behaviors that were common for each predator species. Subsequently, a behavioral ethogram (Lehner 1979) was developed and its components were programmed into The Observer®, a software program designed specifically for animal behavior research (Noldus Information Technology, Version 3.0, www.noldus.com). Descriptions of the predator behaviors that were observed and tabulated during this study are given in Table 1. For most of the 1 hour observations a new feeding arena was used for each observation. In a few instances, limitations in the number of available prey and plant material meant that a predator was removed from the arena after a 1 hour observation and replaced with another randomly selected predator of the same species. No more than two predators were observed consecutively in the same arena. All observations were made between 08:00 and 12:00 at 27°C. The feeding arenas were replaced daily with fresh plant and prey material.

Data analysis

The prey choice and the amount of time that each predator species spent feeding (i.e., handling time) on the various whitefly lifestages did not conform to the assumption of normality, so a Kruskal-Wallis one-way ANOVA on the ranked data was used to determine significant differences between prey choice and prey handling time for each predator species. For the prey choice analysis,

<table>
<thead>
<tr>
<th>Observed behavior</th>
<th>Prey choice</th>
<th>Description of behavior</th>
</tr>
</thead>
<tbody>
<tr>
<td>Walking</td>
<td>Predator moving forward across the leaf surface</td>
<td></td>
</tr>
<tr>
<td>Resting</td>
<td>Predator standing motionless</td>
<td></td>
</tr>
<tr>
<td>Grooming</td>
<td>Predator making rapid movements with its fore and hind legs across its body surface and antennae</td>
<td></td>
</tr>
<tr>
<td>Orienting</td>
<td>Predator pivoting on the leaf without advancing in any particular direction</td>
<td></td>
</tr>
<tr>
<td>Probing</td>
<td>Egg</td>
<td>Predator probing an egg, but not feeding</td>
</tr>
<tr>
<td></td>
<td>Nymph</td>
<td>Predator probing a nymph, but not feeding</td>
</tr>
<tr>
<td></td>
<td>Adult</td>
<td>Predator probing an adult, but not feeding</td>
</tr>
<tr>
<td>Feeding</td>
<td>Egg</td>
<td>Predator consuming an egg</td>
</tr>
<tr>
<td></td>
<td>Nymph</td>
<td>Predator consuming a nymph</td>
</tr>
<tr>
<td></td>
<td>Adult</td>
<td>Predator consuming an adult</td>
</tr>
<tr>
<td></td>
<td>Plant</td>
<td>Predator feeding on plant tissue</td>
</tr>
</tbody>
</table>

Table 1. Descriptions of the behavioral events recorded for the predators exposed to a cotton leaf disk containing the various lifestages of *Bemisia tabaci*. 

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a Student Newman-Keuls multiple comparison test was used to separate statistical differences in prey choice (P = 0.05, SigmaStat Ver. 2.03, SPSS Inc., www.spss.com). For the prey handling time analysis, Dunn’s multiple comparison test was used to separate statistical differences in handling time (P = 0.05).

Behavior transition matrices were determined for each predator species by transferring the observational data obtained from the Observer® from columns of sequences into matrices of preceding and succeeding behavioral events as described by Lehner (1979) and Isaacs et al. (1999). Thereafter, each first-order transition (Slater 1973) from one behavior to the other was analyzed by a $\chi^2$ test to identify those transitional behaviors that were significantly greater than expected by chance. This was done only for those transitions with a frequency greater than 1.0% of the total number of transitions in order to reduce the likelihood of making type II statistical errors (Isaacs et al. 1999). The critical P-value assigned to the $\chi^2$ test was 0.001.

**Results**

*Hippodamia convergens*

Adult *H. convergens* were most commonly observed eating whitefly eggs. On average, 18.3 ± 4.1 eggs, 6.2 ± 1.5 nymphs and 9.6 ± 1.5 adults were consumed per hour of observation (Figure 1A). The time spent feeding on individual whitefly eggs was significantly less than the time spent feeding on nymphs and adults. The average feeding durations were 5.0 ± 0.4, 9.2 ± 0.9, and 11.0 ± 0.4 seconds for eggs, nymphs, and adults, respectively (Figure 1B). *Hippodamia convergens* was rarely observed eating the cotton leaf.

Observations revealed that the proportion of time that *H. convergens* spent on each behavior was grooming (35%) > walking (19%), resting (18%), orienting (17%) > feeding (7%) > probing (4%) (Figure 1C). Even though *H. convergens* consumed twice as many eggs as adults, the majority of its feeding time (40%) was spent feeding on adults (Figure 1C).

*Hippodamia convergens* was very active, yielding a total of 3,464 behavioral events during 33 hours of observation (105.0 events per hour). The general sequence of behavioral events exhibited by *H. convergens* as revealed by the analysis of the transition matrix showed that there is a high likelihood that *H. convergens* will proceed from:

1. egg feeding to egg feeding or walking
2. adult feeding to orienting or walking
3. nymph feeding to nymph feeding or orienting
4. leaf feeding to walking
5. grooming to walking, resting or orienting
6. orienting to grooming, adult feeding or nymph feeding
7. adult probing to walking
8. egg probing to egg feeding
9. resting to grooming
10. walking to grooming, adult feeding, adult probing or leaf feeding.

The most common behavioral sequence recorded was egg feeding followed by another egg feeding 7.8% (e.g., 272 of the 3464 behavioral events) of the time (Table 2). If an individual ate more than one egg in succession, an average of 3.1 ± 1.2 (range 2 to 9) eggs were consecutively consumed. Interestingly, another of the significant transitions was nymph feeding followed immediately by another nymph feeding 1.0% of the time. If an individual ate more than one nymph in succession, an average of 2.7 ± 0.9 (range 2 to 5) nymphs were consecutively consumed.

*Collops vittatus*

Adult *C. vittatus* were most commonly observed feeding
Table 2. Frequencies of transitions between all observed behavioral events for adult *Hippodamia convergens* on cotton leaves that also contained whitefly eggs, nymphs, and adults. Transitions that were significantly more common than expected by chance are shown in grey boxes (chi-square, adjusted P<0.001).

<table>
<thead>
<tr>
<th>Preceding behavioral event</th>
<th>Feeding event</th>
<th>Probing event</th>
<th>Other behaviors</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Egg</td>
<td>Nymph</td>
<td>Adult</td>
</tr>
<tr>
<td>Egg feeding</td>
<td>272</td>
<td>35</td>
<td>4</td>
</tr>
<tr>
<td>Nymph feeding</td>
<td>28</td>
<td>36</td>
<td>2</td>
</tr>
<tr>
<td>Adult feeding</td>
<td>12</td>
<td>4</td>
<td>6</td>
</tr>
<tr>
<td>Leaf feeding</td>
<td>1</td>
<td>0</td>
<td>3</td>
</tr>
<tr>
<td>Egg probing</td>
<td>47</td>
<td>2</td>
<td>16</td>
</tr>
<tr>
<td>Nymph probing</td>
<td>0</td>
<td>10</td>
<td>0</td>
</tr>
<tr>
<td>Adult probing</td>
<td>1</td>
<td>5</td>
<td>6</td>
</tr>
<tr>
<td>Walking</td>
<td>141</td>
<td>55</td>
<td>134</td>
</tr>
<tr>
<td>Resting</td>
<td>9</td>
<td>3</td>
<td>19</td>
</tr>
<tr>
<td>Grooming</td>
<td>31</td>
<td>10</td>
<td>32</td>
</tr>
<tr>
<td>Orienting</td>
<td>63</td>
<td>48</td>
<td>94</td>
</tr>
</tbody>
</table>

on whitefly eggs. On average, 13.0 ± 5.6 eggs, 3.6 ± 1.2 nymphs, and 7.6 ± 1.1 adults were consumed per hour of observation (Figure 2A). The feeding duration for individual eggs was significantly less than for whitefly nymphs and adults (Figure 2B). The average feeding durations were 4.6 ± 0.5, 6.6 ± 1.0, and 14.3 ± 0.6 seconds for eggs, nymphs, and adults, respectively (Figure 2B). *C. vittatus* was rarely observed eating the cotton leaf (Figure 2B).

Observations revealed that the proportion of time that *C. vittatus* spent on each behavior was walking (52%) > grooming (23%) > orienting (11%) > resting (7%), feeding (6%) > probing (1%) (Figure 2C). Although *C. vittatus* ate significantly more eggs than adults, almost one half of its feeding time was spent consuming adult whiteflies (Figure 2C).

*C. vittatus* was very active yielding a total of 2,571 behavioral events during 30 hours of observation (85.7 events per hour). The behavioral sequence analysis revealed that there is a high likelihood that *C. vittatus* will proceed from:

1. adult feeding to walking and orienting
2. egg feeding to egg feeding, walking or orienting
3. nymph feeding to nymph feeding
4. leaf feeding to walking
5. grooming to walking
6. orienting to egg feeding
7. resting to walking and grooming
8. walking to grooming, adult feeding, resting, egg feeding or leaf feeding (Table 3).

The most common behavioral sequences recorded were walking followed by grooming 13.0% of the time and grooming followed by walking 12.6% of the time. As with *H. convergens*, egg feeding followed immediately by another egg feeding (8.9% of the time) and nymph feeding followed immediately by another nymph feeding (1.1% of the time) were significant transitional behaviors. On average, if an individual ate more than one particular prey type in succession, 3.5 ± 1.7 (range 2 to 15) eggs and 2.4 ± 0.5 (range 2 to 3) nymphs were consecutively eaten.

*Geocoris punctipes*

Adult *G. punctipes* were commonly observed feeding (9.8 ± 2.7 times per hour) on the cotton leaf for extended periods of time (98.0 ± 17.4 seconds per feeding event) (Figure 3A-B). The frequency of predation on adult whiteflies was significantly greater than on eggs and nymphs (Figure 3B). Very few whitefly eggs and nymphs were eaten during the 18 hours of observation. On average, 0.1 ± 0.1 eggs, 1.1 ± 0.3 nymphs, and 3.7 ± 1.0 adults, were consumed per hour of observation. The average feeding duration was 8.5 ± 0.5, 54.0 ± 11.8, and 111.5 ± 9.8 seconds for eggs, nymphs, and adults, respectively (Figure 3B).

Observations revealed that the proportion of time that *G. punctipes* spent on each behavior was feeding (40%) > resting (20%), walking (19%), grooming (18%) > orienting (2%) probing (1%) (Figure 3C). Overall, *G. punctipes* spent two-thirds of its feeding time eating the cotton leaf.

*G. punctipes* was an active predator yielding a total of 1,554 behavioral events during 18 hours of observation (86.3 events per hour). The general sequence of behavioral events exhibited by *G. punctipes* as revealed by the analysis of the transition matrix showed that there is a high likelihood that *G. punctipes* will proceed from:

1. leaf feeding to orienting
2. grooming to resting
(3) orienting to leaf feeding  
(4) adult probing to adult feeding  
(5) resting to walking or grooming  
(6) walking to resting.

The most common sequence recorded was walking followed by resting 13.4% of the time, and resting followed by walking 12.4% of the time (Table 4).

*Orius tristicolor*

Adult *O. tristicolor* were most frequently observed eating the cotton leaf (3.7 ± 2.0 times per hour) (Figure 4A). On average, 0.1 ± 0.1 eggs, 0.6 ± 0.4 nymphs, and 2.1 ± 0.5 adults were consumed per hour of observation. The feeding duration on adult whiteflies
Table 3. Frequencies of transitions between all observed behavioral events for adult *Collops vittatus* on cotton leaves that also contained whitefly eggs, nymphs, and adults. Transitions that were significantly more common than expected by chance are shown in grey boxes (chi-square, adjusted P<0.001).

<table>
<thead>
<tr>
<th>Preceding behavioral event</th>
<th>Succeeding behavioral event</th>
<th>Feeding event</th>
<th>Probing event</th>
<th>Other behaviors</th>
</tr>
</thead>
<tbody>
<tr>
<td>Egg feeding</td>
<td>Egg</td>
<td>228</td>
<td>21</td>
<td>7</td>
</tr>
<tr>
<td>Nymph feeding</td>
<td>Nymph</td>
<td>9</td>
<td>28</td>
<td>1</td>
</tr>
<tr>
<td>Adult feeding</td>
<td>Adult</td>
<td>1</td>
<td>1</td>
<td>5</td>
</tr>
<tr>
<td>Leaf feeding</td>
<td>Leaf</td>
<td>2</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td>Egg probing</td>
<td>Egg probing</td>
<td>12</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>Nymph probing</td>
<td>Nymph probing</td>
<td>0</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>Adult probing</td>
<td>Adult probing</td>
<td>0</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td>Walking</td>
<td>Walking</td>
<td>71</td>
<td>39</td>
<td>174</td>
</tr>
<tr>
<td>Resting</td>
<td>Resting</td>
<td>3</td>
<td>0</td>
<td>5</td>
</tr>
<tr>
<td>Grooming</td>
<td>Grooming</td>
<td>6</td>
<td>5</td>
<td>16</td>
</tr>
<tr>
<td>Orienting</td>
<td>Orienting</td>
<td>58</td>
<td>6</td>
<td>18</td>
</tr>
</tbody>
</table>

Table 4. Frequencies of transitions between all observed behavioral events for adult *Geocoris punctipes* on cotton leaves that also contained whitefly eggs, nymphs, and adults. Transitions that were significantly more common than expected by chance are shown in grey boxes (chi-square, adjusted P<0.001).

<table>
<thead>
<tr>
<th>Preceding behavioral event</th>
<th>Succeeding behavioral event</th>
<th>Feeding event</th>
<th>Probing event</th>
<th>Other behaviors</th>
</tr>
</thead>
<tbody>
<tr>
<td>Egg feeding</td>
<td>Egg</td>
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<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Nymph feeding</td>
<td>Nymph</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Adult feeding</td>
<td>Adult</td>
<td>0</td>
<td>0</td>
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</tr>
<tr>
<td>Leaf feeding</td>
<td>Leaf</td>
<td>0</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td>Egg probing</td>
<td>Egg probing</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Nymph probing</td>
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<tr>
<td>Adult probing</td>
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<td>44</td>
</tr>
<tr>
<td>Walking</td>
<td>Walking</td>
<td>1</td>
<td>3</td>
<td>9</td>
</tr>
<tr>
<td>Resting</td>
<td>Resting</td>
<td>1</td>
<td>5</td>
<td>10</td>
</tr>
<tr>
<td>Grooming</td>
<td>Grooming</td>
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<td>1</td>
<td>3</td>
</tr>
<tr>
<td>Orienting</td>
<td>Orienting</td>
<td>0</td>
<td>2</td>
<td>0</td>
</tr>
</tbody>
</table>

was significantly greater than for the egg and nymphal stages. The average feeding durations were 45.0 ± 12.0, 44.4 ± 9.7, and 455.1 ± 65.3 seconds for eggs, nymphs, and adults, respectively (Figure 4B).

Observations revealed that the proportion of time that adult *O. tristicolor* spent on each behavior was feeding (35%) > walking (26%) > resting (16%), grooming (14%) > orienting (8%) > probing (1%) (Figure 4C). Overall, *O. tristicolor* spent three-fourths of their feeding time eating adult whiteflies. Less than 3.0% of their feeding time was spent eating eggs and nymphs (Figure 4C).

A total of 1,489 behavioral events were recorded during 22 hours of observation (67.7 per hour). The general sequence of
behavioral events exhibited by *O. tristicolor* as revealed by the analysis of the transition matrix showed that there is a high likelihood that *O. tristicolor* will proceed from:

1. grooming to walking
2. leaf feeding to orienting
3. orienting to leaf feeding
4. adult probing to walking and adult feeding
5. walking to grooming, resting and adult probing.

The most common sequence recorded was walking followed by grooming 16.8% of the time and grooming followed by walking 16.5% of the time (Table 5).
Table 5. Frequencies of transitions between all observed behavioral events for adult *Orius tristicolor* on cotton leaves that also contained whitefly eggs, nymphs, and adults. Transitions that were significantly more common than expected by chance are shown in grey boxes (chi-square, adjusted P<0.001).

<table>
<thead>
<tr>
<th>Preceding behavioral event</th>
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<th>Probing event</th>
<th>Other behaviors</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Egg</td>
<td>Nymph</td>
<td>Adult</td>
</tr>
<tr>
<td>Egg feeding</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Nymph feeding</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Adult feeding</td>
<td>0</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Leaf feeding</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Egg probing</td>
<td>2</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Nymph probing</td>
<td>0</td>
<td>6</td>
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</tr>
<tr>
<td>Adult probing</td>
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<td>34</td>
</tr>
<tr>
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<td>3</td>
<td>8</td>
</tr>
<tr>
<td>Resting</td>
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<td>3</td>
<td>6</td>
</tr>
<tr>
<td>Grooming</td>
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<td>0</td>
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</tr>
<tr>
<td>Orienting</td>
<td>0</td>
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</tr>
</tbody>
</table>

Table 6. Frequencies of transitions between all observed behavioral events for adult *Lygus hesperus* on cotton leaves that also contained whitefly eggs, nymphs, and adults. Transitions that were significantly more common than expected by chance are shown in grey boxes (chi-square, adjusted P<0.001).

<table>
<thead>
<tr>
<th>Preceding behavioral event</th>
<th>Feeding event</th>
<th>Probing event</th>
<th>Other behaviors</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Egg</td>
<td>Nymph</td>
<td>Adult</td>
</tr>
<tr>
<td>Egg feeding</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Nymph feeding</td>
<td>0</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Adult feeding</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Leaf feeding</td>
<td>1</td>
<td>12</td>
<td>0</td>
</tr>
<tr>
<td>Egg probing</td>
<td>0</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Nymph probing</td>
<td>1</td>
<td>27</td>
<td>0</td>
</tr>
<tr>
<td>Adult probing</td>
<td>0</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Walking</td>
<td>1</td>
<td>22</td>
<td>1</td>
</tr>
<tr>
<td>Resting</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Grooming</td>
<td>1</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td>Orienting</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

Adult *L. hesperus* were most commonly observed feeding (5.33 ± 0.71 times per hour) for extended periods of time (235.3 ± 123.2 seconds per feeding event) on the cotton leaf (Fig. 5A-B). Adult *L. hesperus* were next most commonly observed preying on whitefly nymphs. Very few whitefly eggs (4) or adults (3) were eaten during the 27 hours of observation. On average, 0.2 ± 0.1 eggs, 2.4 ± 1.0 nymphs, and 0.1 ± 0.1 adults were consumed per hour of observation (Figure 5A). The average prey feeding durations were 28.0 ± 17.2, 93.0 ± 11.4, and 221.7 ± 26.7 seconds for eggs, nymphs, and adults, respectively (Figure 5B).

Observations revealed that the proportion of time that *L. hesperus* spent on each behavior was feeding (42%) > grooming.
spent about 7.0% of their overall time budget feeding, but
lifestages with a preference for eggs > adults > nymphs. Both species
spent eating the plant (Figure 5C). The majority (83%) of their feeding time was
on all whitefly stages.

L. hesperus was not as active as the other predators examined in this study yielding a total of 670 behavioral events during 27 hours of observation (28.8 events per hour). The general sequence of behavioral events recorded for L. hesperus, as revealed by the analysis of the transition matrix, showed that there is a high likelihood that L. hesperus will proceed from:

1. nymph feeding to nymph probing and nymph probing to nymph feeding
2. egg probing to nymph feeding
3. grooming to walking and walking to grooming.

The most common sequence recorded was walking followed by grooming 11.8% of the time and grooming followed by walking 10.4% of the time (Table 6).

Discussion

Generally, insect predators either stalk or ambush their prey (Murdoch et al. 1985; O’Brien et al. 1989; Sabelis 1992; Hagler 2000). Certain assumptions about prey selection have been made for some whitefly predators based solely on their predominant hunting strategy (Breene 1992; Breene et al. 1992; Nordlund and Legaspi 1996). For example, ambush predators that place themselves in a strategic location and wait until prey move into their field of attack are considered to be best adapted for capturing mobile prey such as adult whiteflies (Sabelis 1992; Breene 1992). Conversely, stalking predators that constantly roam their environment searching for prey are considered to be best adapted for capturing immobile prey such as whitefly eggs and nymphs (Breene et al. 1992, 1994; Nordlund and Legaspi 1996). The predators examined in this study are generally regarded as stalking predators. As a consequence, an overwhelming majority of studies examining whitefly predation have been restricted to studies with the sessile whitefly lifestages conducted in no-choice arenas containing either eggs or nymphs or in choice arenas containing both eggs and nymphs (Elbadry 1968; Gerling 1986; Butler and Henneberry 1988; Kapadia and Puri 1991; Hoelmer et al. 1993, 1994; Legaspi et al. 1994; Nordlund and Legaspi 1996; Guershon and Gerling 1999). To our knowledge, only a few no-choice experiments have been conducted to identify the sources of adult whitefly predation. Of these, various species of true bugs, flies, and spiders have been identified as predators of adults (Sussman 1988; Cohen and Byrne 1992; Breene et al. 1992, 1994). Furthermore, to date only one predator evaluation has been conducted in which a predator was simultaneously presented with a choice of whitefly eggs, nymphs and adults. In that study, Hagler (2002) showed that the predatory fly, Drosophila nr. divergens fed exclusively on adult whiteflies. Clearly, as Nordlund and Legaspi (1996) pointed out, more studies are needed to evaluate predation on all whitefly stages.

The results from our study revealed that the beetles, H. convergens and C. vittatus, readily and rapidly preyed on all whitefly lifestages with a preference for eggs > adults > nymphs. Both species spent about 7.0% of their overall time budget feeding, but H.

Prey selection by the true bugs, G. punctipes, O. tristicolor, and L. hesperus yielded some surprising results. For example, G. punctipes and O. tristicolor preyed on adults over three times more frequently than on eggs and nymphs combined. In contrast, L. hesperus, a notorious pest in many cropping systems, was observed feeding on nymphs over nine times more frequently than on eggs and adults combined. Our observations of G. punctipes and those of Cohen and Byrne (1992) indicate that adult whiteflies rarely escaped when attacked by true bugs. Further observations revealed that the true bugs did not feed on nearly as many whiteflies as the beetles and spent a considerably longer time (e.g., 5 to 25 times longer) eating their prey. The increased handling time exhibited by the true bugs when compared to the beetles is due primarily to differences in their feeding mechanisms. The chewing mouthparts of beetles enable them simultaneously to kill by crushing or macerating and to rapidly devour their prey (see Video 1 available at http://insectscience.org/4.1/). Conversely, the true bugs have piercing and sucking mouthparts. The stylects are used to pierce the prey, to disrupt the cell walls and cellular contents, and to deliver saliva containing potent digestive enzymes (see Video 2 available at http://insectscience.org/4.1/). Together, the mechanical damage and the enzymes break down the tissue into a slurry of small particles, which are ingested along with the saliva (Cohen 1998). Overall, the time budgets exhibited for each of the true bugs were similar with all three species spending approximately one-third of their time in the immobile resting and grooming behaviors, two common characteristics of an ambush predator. Furthermore, approximately 40% of their time was spent feeding on either plant or insect tissue. However, there was a sharp contrast between the amounts of time that G. punctipes and L. hesperus spent feeding on plant tissue compared to O. tristicolor. Analysis of the feeding times revealed that L. hesperus, G. punctipes, and O. tristicolor spent 83, 66, and 21% of their feeding time on plant tissue, respectively. The majority of feeding by O. tristicolor’s (77%) was spent eating adult whiteflies.

The prey choices observed in the feeding arena provided some insight to the results obtained from previous gut content examinations on these predators using a whitefly-specific ELISA (Hagler and Naranjo 1994a,b). In those studies, ~35% of all field collected G. punctipes and O. tristicolor contained whitefly egg protein in their guts (Hagler and Naranjo 1994a). However, as discussed above, a positive ELISA reaction can be obtained for a predator that eats a whitefly egg or a gravid adult female (Hagler et al. 1993). Originally we believed that these predators were feeding primarily on whitefly eggs. The results from this study suggest that the majority of positive responses yielded by field-collected G.
**punctipes** and *O. tristicolor* might be more likely due to predation on adult females. Prey selection by *L. hesperus*, *H. convergens*, and *C. vittatus* make it impossible to differentiate a positive ELISA response for the presence of whitefly between an egg and an adult female. For instance, while *L. hesperus* preyed almost exclusively on whitefly nymphs, their rare occurrence of predation on eggs and adults was about the same. Conversely, *H. convergens* and *C. vittatus* readily preyed on all whitefly life stages. The voracious appetite exhibited by the beetles for every whitefly life stage suggests that a positive whitefly ELISA response is probably due to the beetles feeding on a combination of eggs and adult females.

We would be remiss if we did not discuss some of the potential drawbacks associated with studies of predation conducted in confined feeding arenas. First, the whitefly population in each arena was very high. This could lead to an overestimation of predation because it minimizes the time needed for the predator to search for its prey. However, the whitefly densities in our arenas were not atypical of the densities often found late in the season in the cotton growing regions of the Southwestern United States (Ellsworth et al. 1993; Natwick et al. 1995). Second, the predators and whiteflies were fully enclosed in a Petri dish. The enclosed arena can reduce the ability of an adult whitefly to escape and the ability of the predator to move to a different plant or patch. This too would likely lead to an overestimation of predation rates. Moreover, no alternate prey species were made available to the predators in the enclosed arenas. The presence of alternate prey would likely reduce the number of whiteflies consumed by these generalist predators, especially if the alternate prey species were preferred or prey selection were random. Finally, in some instances, two predators were observed consecutively in the same feeding arena. While we did not observe any noticeable differences in the behaviors between the two individuals, it is possible that the behavior of the first predator could have indirectly affected the behavior of the second due to an induced response mediated by the plant, herbivore, and/or predator (Dicke et al. 1990, 2003; Dicke 1994, 2000; Dicke and van Loon 2000; Cortesero et al. 2000). The investigation of chemically induced responses between the trophic levels was beyond the scope of this investigation.

The major objective of this study was to identify prey selection by these five whitefly predators. Additionally, we identified the propensity of these predators to feed on cotton foliage. The frequency and duration of cotton feeding was highly variable between the beetles and the true bugs. The beetles were sometimes observed briefly nibbling or “skimming” the surface of the cotton leaf, probably to obtain water, sugar and other micronutrients. On the other hand, the true bugs were often observed inserting their piercing and sucking mouthparts deep into leaf veins and feeding for extended periods of time. It appears from this study and others that plant feeding is an important component to the nutritional ecology of many heteropterans (Naranjo and Gibson 1996; Wiedenmann et al. 1996; Coll 1996, 1998; Coll and Izraylevich 1997; Coll and Guershon 2002). For example, while it has been determined that *Geocoris* spp. can be reared in the laboratory exclusively on a meat diet (Cohen 1985), their fitness can be increased with the addition of plant nutrients to their diet (Dunbar and Bacon 1972; Tamaki and Weeks 1972; Naranjo and Gibson 1996). For instance, Tamaki and Weeks (1972) showed that *Geocoris bullatus* and *G. pallens* had increased longevity and fecundity and a faster rate of development when a diet of aphids was supplemented with plant material. Others have shown that facultative phytophagy by true bugs provides the insects with water, but also may provide some nutrients that supplement prey diet and help the predator survive periods when prey are scarce (Ridgway and Jones 1968; Stoner 1972; Salas-Aguilar and Ehler 1977; Kiman and Yeargan 1985; Wiedenmann and O’Neil 1991, 1992; Naranjo and Gibson 1996; Armer et al. 1998; Coll and Guershon 2002). Eubanks and Denno (2000) reported that the availability of high quality bean pods in the diet of *Geocoris* spp. resulted in a shift from carnivory to phytophagy.

Armer et al. (1998) state that a continuum exists among heteropteran feeding characteristics ranging from facultative phytophagous predators to facultative predaceous herbivores (Wiedenmann et al. 1996). While facultative herbivory is common among many predaceous true bugs (Polis et al. 1989; Rosenheim et al. 1995; Naranjo and Gibson 1996; Polis and Winemiller 1996), it is surprising that a greater research effort has not been made towards determining what advantages in fitness are derived by “predators” that feed on plants and “herbivores” that feed on insects. Insight into whether these true bugs are primarily predators or primarily herbivores might be determined by studies of their digestive enzyme complex. Specifically, researchers have determined that the presence of certain digestive enzymes indicates an insect’s ability to efficiently eat plant tissue, animal tissue, or both (Baptist 1941; Adams and McAllen 1956; Strong and Kruitwagen 1968; Tinge y and Pillemer 1977; Cohen 1995, 1998). For example, Agusti and Cohen (2000) conducted a study of the digestive enzyme complex of two omnivorous *Lygus* spp. They concluded that *L. hesperus* was better suited to carnivory than *L. lineolaris* because it has greater trypsinlike activity, and *L. lineolaris* was better suited to herbivory because it had greater α-amylase and pectinase activity.

In conclusion, our study provides insight into the host range and feeding characteristics of five common whitefly predators. The beetles preyed on all of the whitefly life stages with a preference for eggs and adults, whereas *G. punctipes* and *O. tristicolor* preferred adults, and *L. hesperus* preferred nymphs. Based solely on the results from this laboratory study and from previous predator gut content evaluations (Hagler and Naranjo 1994a,b), we conclude that adult *H. convergens* and *C. vittatus* will be more effective whitefly biological control agents than the adult true bugs, *G. punctipes* and *O. tristicolor* in the cotton system of the Southwestern United States. Specifically, *H. convergens* and *C. vittatus* fed on many more whiteflies in less time than *G. punctipes* and *O. tristicolor* and their prey choice was directed more against the egg stage than the other species tested. Furthermore, we have shown that the proportion of beetles feeding on whiteflies in the field is over twice as high as that of the true bugs. While *L. hesperus* is a predator of whitefly (Hagler and Naranjo 1994a) and many other pest species (Knight 1941; Lindquist and Sorensen 1970; Cleveland 1987; Bryan et al. 1976; Wheeler 1976; Hagler and Naranjo 1994a,b), it is not a viable biological control agent because of its propensity to feed on plant merismatic tissue.

This study and other laboratory studies indicate that predatory beetles possess many characteristics that make them superior biological candidates, but we caution that such laboratory
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