Reproductive performance and longevity of female European corn borer, Ostrinia nubilalis: effects of multiple mating, delay in mating, and adult feeding

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Abstract

In a recent study on the pheromone-mating disruption of the European corn borer, Ostrinia nubilalis (Lepidoptera: Crambidae), we recorded a significant reduction in mating frequency, as well as a marked delay in mating in feral females captured in disruptant-treated fields. In order to be able to accurately interpret the results in terms of effective population control, the current study was undertaken on the effects of multiple matings and a delay in mating on reproductive performance. Female O. nubilalis that mated at least twice had significantly higher fecundity and fertility, compared with once-mated females. In addition, multiple-mated females deposited a significantly larger portion of their egg complement, relative to single-mated or unmated females. Females that experienced a 3-day delay in mating showed a significant reduction in fecundity compared with females that mated soon after emergence. A 1-week delay in mating resulted in a further reduction in fecundity and a near zero fertility. The effect of sugar feeding on reproduction was not significant. In general, unmated females lived longer than mated females, and sugar-fed mated females had a higher longevity than water-fed mated females. © 1999 Elsevier Science Ltd. All rights reserved.

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1. Introduction

Polyandry, a reproductive strategy in which insects acquire several mates during their reproductive life is widespread in the Lepidoptera (Drummond, 1984; Royer and McNeil, 1993; Ward and Landolt, 1995; Delisle and Hardy, 1997; Landolt, 1997). Several possible reasons have been proposed for multiple matings in female moths. The most important potential benefits to the female include: enhancement of fecundity and longevity by obtaining non-sperm resources from males; achievement of enough sperm supply to fertilize her full egg complement; or increase in the genetic diversity of progeny (Drummond, 1984; Marshall and McNeil, 1989; Ramaswamy et al., 1997).

The European corn borer, Ostrinia nubilalis (Hübner) (Lepidoptera: Cranbidae) is a major pest of corn, Zea mays L. in the United States. Although the female has been classified as normally monandrous (Caffrey and Worthley, 1927; McNeil et al., 1997), several workers have reported multiple matings in feral and laboratory-reared O. nubilalis females. Pesho (1961) recorded 8–43% multiple matings in feral females, while Showers et al. (1974) reported that 10% of females captured in light traps had mated twice. This was in the same range as the 11% multiple matings recorded by Loughner (1971). Drecktrah and Brindley (1967) recorded a higher proportion (46.5%) of multiple mated females, and Elliot (1977) reported that 29–37% of a feral population in southwestern Ontario had mated twice, and up to 13% three times. In their study on the reproductive performance of females reared on different diets, Onokogu et al. (1980) recorded a maximum of six spermatophores with multiple matings in the range of 30–70%.

In a recent study conducted in Iowa on the potential of pheromone mating disruption of the European corn borer, we recorded a maximum of four matings per female (Fadamiro et al., unpublished data). Of the 327 feral free-flying females net-captured in untreated
(check) plots during the first flight, about 70% had mated more than once. Also, 75% multiple matings \((n = 274)\) was recorded in the check plots during the second flight. The incidence of multiple mating was more preponderant during the peak than at the beginning of each flight. While the mean number of spermatophores averaged about two per female in the check plots, a significant reduction in mating frequency was recorded in the treated plots \((1.33-1.58\) spermatophores per female) (Fadamiro et al., unpublished data). Furthermore, our data showed that a higher level of mating disruption was achieved during the beginning of the first flight: about 50–60% of females captured in pheromone-treated plots were unmated during the first 10 days of the first flight, compared with the 90–100% mated females captured in the check plots during the same period.

In order to be able to accurately interpret our results, we need to know what influence mating frequency, as well as a delay in first mating might have on the reproductive success of \(O. nubilalis\) females. Because we are not aware of any studies conducted to directly address these pertinent questions, the current study was commenced to try to begin to understand why European corn borer females mate more than once.

In addition, previous studies on the effect of adult feeding on reproductive fitness in European corn borer have achieved variable results. Kira et al. (1969) suggested that only drinking water was essential for egg production by females. In recent studies, however, sugar-containing diets were shown to increase fecundity and longevity (Miller, 1988; Leahy and Andow, 1994). Thus, our experiment was designed to test the effect of sugar feeding on reproductive output and longevity of female European corn borer, as well as possible interactions with mating frequency.

2. Materials and methods

2.1. Insect rearing

The Iowa strain of \(O. nubilalis\) is bivoltine, with a female sex pheromone blend of the \((Z)\) and \((E)\) isomers of 11-tetradecenyl acetate in ratio 97:3 (Klun et al., 1973). Adults used in this experiment were from wild (1996) populations reared in a laboratory for 19 generations. Adults were either sexed at the pupal stage, or on the first day of emergence to minimize labor. Sexing of adults on the first day of emergence was sufficient for this study, since no matings by European corn borer were recorded during the first 12 h of emergence (Royer and McNeil, 1991; personal observation). An adult was considered 0 day old on the day of eclosion.

2.2. Experimental protocol

In order to maximize oviposition, the experiment was conducted under cyclic photoperiod and temperatures \((L:D, 16:8; 27.23 \pm 1^\circ\text{C}; 90 \pm 5\% \text{ relative humidity})\) (Binder and Robbins, 1996). Newly emerged virgin females were placed in individual \(7.6 \text{ cm} \times 7.6\)-cm-diameter cylindrical screen cages made of 18-mesh copper wire (Kira et al., 1969). The cages were then placed in plastic trays containing a thick layer of white sterile absorbent cotton (Acco Absorbent Cotton Co., MI) dampened with water or 8% sucrose-water. The cotton was replaced every 5 days.

There were six groups of females:

(i) female supplied with no males throughout its entire life;
(ii) female paired with a fresh male for 24 h only on day 1 post-emergence;
(iii) female paired with a fresh male each day for 3 days starting from day 1 post-emergence;
(iv) female paired with a fresh male each day for up to 5 days starting from day 1 post-emergence;
(v) female paired with a fresh male each day for up to 5 days, starting from day 4 post-emergence;
(vi) female paired with a fresh male each day for up to 5 days starting from day 7 post-emergence.

Females in groups (iv), (v) and (vi) were paired with fresh males for up to 5 days, or until death, whichever came first. Early mortality of adult European corn borer was minimal under these experimental conditions. Twenty-eight females were placed in each group.

To determine the effect of food on reproductive output and longevity, half of the females in each group were fed on water while the other half were fed on 8% sucrose-water (Flavorite Sugar, Preferred Products Inc., MN). Where applicable, a fresh (1 or 2-day-old) virgin male was paired with a female. The male was removed after 24 h, and where applicable, replaced with another fresh virgin male.

A sterile disposable polystyrene plastic Petri dish \((100 \text{ mm} \times 15 \text{ mm upper lid part}, \text{ Fisher Scientific, PA})\) was placed on top of the cylindrical cage as an oviposition site, and was changed daily. Eggs were incubated for 5 days on the oviposition site, under the experimental conditions described above.

To determine the fecundity of a female, the total area of an egg mass deposited on a Petri dish was measured on a digital analyzer (Decagon Industries Monochrome Ag Vision Imaging System, Pullman, WA), a standard method of estimating fecundity in European corn borer (Binder and Robbins, 1996). Measurement of egg mass was done on the second day following deposition. Areas measured were later converted to number of eggs using a calculated conversion factor of 0.20 mm per egg. Singly-laid eggs were counted manually. Preoviposition
3.1. Mating patterns

The general mating pattern of female European corn borer under different conditions is shown in Table 1. Although 28 females were placed in each mating group at the start of the experiment, only females that made it to the seventh day post-emergence ‘cut-off’ were included in the data (Table 1). Thus, the difference between the starting sample size (28 females per group) and sample size reported in Table 1 reflects loss of females due to early mortality or escape.

Only 28% of the 1-day-old virgin females paired with a male for 24 h had mated, and none more than once. A higher percentage of mating (68%) was recorded for females paired with a fresh male each day for 3 days, with multiple matings of about 29%. Of the 27 1-day-old virgin females paired with a fresh male each day for up to 5 days, 78% had mated at least once. This is in the same range as the 68% mating recorded for the females who gained their first access to male on their fourth day of emergence. Virgin females whose first access to a male was delayed for about a week mated the least (20.8% mating), and none more than once (Table 1). About 38% of the 1-day-old virgin females paired with a fresh male each day for up to 5 days mated more than once, compared with 10.5% multiple mating recorded for 4-day-old virgin females with the same length of access to males. Only one female mated three times, and this was in the treatment where newly emerged females had access to a fresh male for up to 5 days. Very few females that experienced a 7-day delay in mating mated (20.8%), and consequently this group was excluded from statistical analyses of the effects of mating frequency and food on reproductive performance.

3.2. Effect of mating frequency

Significantly higher fecundity ($P = 0.0001$) and fertility ($P = 0.0001$) were recorded for multiple-mated females, compared with single-mated, or unmated females (Table 2A). Multiple-mated females laid virtually all of their egg complement and had a significantly lower index of remaining eggs. Unmated females had a significantly lower fecundity, longer preoviposition period, higher longevity, and higher index of remaining eggs ($P = 0.001$) than both classes of mated females (Table 2A). Most of the eggs deposited by unmated females were laid as scattered, single eggs, rather than as egg masses, and all were infertile. No significant interactions were recorded among mating frequency, food and all the fitness parameters ($P > 0.05$).

Similar results were obtained when females were classified according to the number of ruptured spermatophores in their bursa (Table 2B), with the differences in reproductive output among the three groups of females becoming more pronounced. Females that had two ruptured spermatophores in their bursa had significantly higher fecundity ($P = 0.001$), fertility ($P = 0.0001$) and a lower index of remaining eggs ($P = 0.0001$), than females that had one or no ruptured spermatophores (Table 2B). Paired females having no ruptured spermatophores in their bursa had a longer preoviposition period ($P = 0.001$), and a higher longevity ($P = 0.005$) than females that had one or two ruptured spermatophores (Table 2B).

3.3. Effect of delay in first mating

Since few multiple matings were recorded in this study, and virtually none for females that had delayed access to males, comparisons were made among once-mated females with their first access to a male on 1-, 4-, or 7-days post-emergence. Significant differences in
Table 1
Mating patterns of female *O. nubilalis* under different conditions

<table>
<thead>
<tr>
<th>Female category</th>
<th>No. females paired</th>
<th>% mated</th>
<th>No. females with 1–3 spermatophores</th>
<th>% multiple matings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Paired for 1 day on day 1 post-emergence</td>
<td>25</td>
<td>28.0</td>
<td>7</td>
<td>0</td>
</tr>
<tr>
<td>Paired for 2–3 days from day 1 post-emergence</td>
<td>25</td>
<td>68.0</td>
<td>12</td>
<td>5</td>
</tr>
<tr>
<td>Paired for 4–5 days from day 1 post-emergence</td>
<td>27</td>
<td>77.8</td>
<td>13</td>
<td>7</td>
</tr>
<tr>
<td>Paired for 3–5 days from day 4 post-emergence</td>
<td>28</td>
<td>67.9</td>
<td>16</td>
<td>2</td>
</tr>
<tr>
<td>Paired for 3–5 days from day 7 post-emergence</td>
<td>24</td>
<td>20.8</td>
<td>5</td>
<td>0</td>
</tr>
</tbody>
</table>

An insect was considered 0 day old on the day of emergence. A female was paired daily with a fresh 1- or 2-day-old virgin male. With the exception of the first two categories of females that were paired with males for a maximum duration of 1 to 3 days, a female was paired with a fresh virgin male every day for a maximum of 5 days or until death, whichever came first.

*The difference between the starting sample size for each group (28 females per group) and the number of females reported reflects female loss due to early mortality or escape.

Table 2
Effect of mating frequency on reproductive performance and longevity of female *O. nubilalis* categorized by: (A) total number of spermatophores in the bursa, and (B) number of ruptured spermatophores in the bursa

<table>
<thead>
<tr>
<th>Female category</th>
<th>No. females observed</th>
<th>Fecundity (no. of eggs deposited)</th>
<th>% fertility</th>
<th>Pre-oviposition period (days)</th>
<th>Longevity (days)</th>
<th>Index of remaining eggs</th>
</tr>
</thead>
<tbody>
<tr>
<td>A. Total number of spermatophores</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Unmated</td>
<td>53</td>
<td>163 ± 18c</td>
<td>0 ± 0c</td>
<td>7.1 ± 3.6a</td>
<td>15.1 ± 3.8a</td>
<td>3.8 ± 1.1a</td>
</tr>
<tr>
<td>Single-mated</td>
<td>48</td>
<td>540 ± 292b</td>
<td>64.9 ± 34.8b</td>
<td>3.9 ± 1.6b</td>
<td>13.0 ± 3.6b</td>
<td>1.7 ± 1.4b</td>
</tr>
<tr>
<td>Multiple-mated</td>
<td>15</td>
<td>630 ± 181a</td>
<td>77.5 ± 29.9a</td>
<td>3.3 ± 1.5b</td>
<td>11.8 ± 2.6b</td>
<td>0.7 ± 0.9c</td>
</tr>
<tr>
<td>B. Number of ruptured spermatophores</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0</td>
<td>32</td>
<td>133 ± 170c</td>
<td>0 ± 0c</td>
<td>7.0 ± 3.4a</td>
<td>14.8 ± 3.7a</td>
<td>4.0 ± 0.9a</td>
</tr>
<tr>
<td>1</td>
<td>55</td>
<td>538 ± 274b</td>
<td>63.9 ± 34.5b</td>
<td>3.7 ± 1.6b</td>
<td>12.8 ± 3.6b</td>
<td>1.6 ± 1.4b</td>
</tr>
<tr>
<td>2</td>
<td>7</td>
<td>742 ± 193a</td>
<td>96.3 ± 4.7a</td>
<td>3.3 ± 1.8b</td>
<td>11.4 ± 1.3b</td>
<td>0.4 ± 0.5c</td>
</tr>
</tbody>
</table>

Values (means ± SD) in the same column, under the same classification, having no letters in common are significant at *P* < 0.05.

*Females with a 7-day delay in mating were excluded from the analysis.

Effect of food

In order to detect possible interactions among food, mating status and the different fitness parameters, unmated and mated females were grouped separately and the effect of food was determined. Single-mated females with access to sugar-water showed a slight but not sig-
4. Discussion

In addition to enhancing fecundity, multiple mating was shown to increase the fertility of female European corn borer in this study. Results from other studies on various lepidopteran species have recorded similar positive correlation between mating frequency and fecundity (Wilkund et al., 1993; Svärd and McNeil, 1994; Ward and Landolt, 1995; Landolt, 1997), fertility (Marks, 1976; Delisle and Hardy, 1997), or both (Svärd and McNeil, 1994; Delisle and Hardy, 1997). Simmons (1988) also recorded increased fecundity and hatching success for multiple-mated female crickets, Gryllus bimaculatus. Alternative hypotheses that have been proposed regarding the role of multiple mating in female insects include: increase in genetic diversity of offspring, sperm competition, prevention of possible genetic defects due to long-term storage of sperm, minimization of time and energy costs of resisting males, and achievement of factors that stimulate further egg development (Parker, 1970; Drummond, 1984; Halliday and Arnold, 1987; Marshall and McNeil, 1989; Ramaswamy et al., 1997).

The observed increase in the fecundity of multiple-mated females has been attributed to the role in egg production, of male-derived nutrients transferred along with sperm during mating (Boggs, 1981; Svärd and McNeil, 1994). Male accessory gland secretions transferred with spermatophore during copulation have been shown to play an important role in reproduction (Leopold, 1976; Boggs and Gilbert, 1979; Watanabe, 1988; Boggs, 1990). Female insects may acquire resources necessary for egg production (Boggs, 1990), or achieve factors that stimulate egg maturation (Ramaswamy et al., 1997), from the male at the time of mating. However, it is not
clear whether the increased fecundity recorded for multiple-mated *O. nubilalis* females in the current study was due to an increase in egg production in vivo, or simply an increase in egg deposition by multiple-mated females.

The significant reduction in index of remaining eggs recorded for multiple-mated females suggests that remating enhanced egg deposition as Watanabe (1988) proposed for the yellow swallowtail butterfly, *Papilio xuthus*. Herman and Barker (1977) reported that male-derived factors may induce oogenesis through hormonal effects. Our results, therefore, suggest that male-derived factors may also be playing a role in reproduction by inducing or enhancing oviposition (egg deposition), perhaps through similar hormonal effects.

Male-derived nutrients can contribute to egg production only in species that eclose as adults with little or no eggs yolked (Boggs, 1990). For these species also, adult feeding may play an important role in egg production (Boggs, 1990). Adult European corn borers have been observed to exhibit feeding-related behaviors in the field (Showers et al., 1976; Cantelo and Jacobson, 1979). However, results of work on the effect of adult feeding on the reproductive performance of female *O. nubilalis* have not been consistent. Caffrey and Worthley (1927) first reported that sugars in imbibed water did not improve reproduction, and Kira et al. (1969) concluded that only drinking water was essential for egg production by female European corn borer. Miller (1988) reported that females that were fed on 15% honey-water showed a significant improvement in four reproductive performance attributes, but not in total fecundity, than females that were fed on water alone. In contrast, Leahy and Andow (1994) recorded a significant increase in fecundity for European corn borer females that were fed on diets containing sugar. Although direct comparisons cannot be made among the different studies due to differences in experimental design and moth colonies, it is possible that the differences in these results could be due to the different classes and concentrations of sugar used in the different studies. While Miller (1988) utilized 15% fructose-water in his experiment, Leahy and Andow (1994) used both fructose (42% solution) and sucrose at very high concentrations. Since only 8% sucrose-water was presented to adults in the current study, it is possible that the use of a higher concentration of sucrose may lead to a significant improvement in fecundity. That adult feeding may contribute to egg production in *O. nubilalis*, as shown for some closely related pyralid species (Waldbauer et al., 1980; Taylor and Sands, 1986), seems not unlikely in light of the report that adult European corn borer females eclose with an average of only 90 mature oocytes (Miller, 1988).

Although multiple-mated females generally produced significantly more progeny than once-mated females in the current study, some (about 10%) of the females that mated once had over 90% of their eggs fertilized by the single mating. It thus appears that a female’s need to remate in order to acquire enough sperm to fertilize her full complement of eggs may depend on the size of her egg complement, as well as the size of the spermatophore transferred by her first mate. Indeed, Royer and McNeil (1993) have recorded a negative correlation between successive matings and the size of spermatophores transferred by male *O. nubilalis*. This decrease in spermatophore volume with subsequent matings translated to a decrease in the fertility of successive mated of a given male. A study in which both the size of a European corn borer female’s full egg complement (potential fecundity) and the volume of spermatophores transferred by males are controlled may be necessary to examine possible interactions among these factors and remating tendency/frequency.

In the current study, a 3-day delay in mating was shown to significantly reduce fecundity, while a 1-week delay resulted in near zero fertility. This considerable reduction in fertility in female *O. nubilalis* that experienced a delay in mating would offset any adaptive value of the ability of these females to prolong their preoviposition period. Knight (1997) recorded a 40% reduction in fecundity for *Cydia pomonella* females that experienced a 2-day delay in mating. No significant reduction in reproductive output, however, was recorded for female Spiny bollworm, *Earias insulana* whose mating was delayed up to about the 9th night (Kehat and Gordon, 1977). It thus appears that whether or not a delay in mating would affect reproduction depends on the biology of the species. For instance, in the case of *E. insulana*, the preoviposition period was greatly prolonged in unmated females, enabling the female to delay oviposition until after mating. Also, female *E. insulana* had a higher natural longevity and longer oviposition period (Kehat and Gordon, 1977) than female *O. nubilalis* (present study). It is, perhaps, reasonable to suggest that longer-lived females with a longer oviposition period might also have a longer period of receptivity to the male, compared with short-lived species. Invariably, species that possess an efficient mechanism of this type that ensures reproductive success, will be relatively unaffected by a delay in mating. The present results suggest that the female European corn borer had either stopped calling, or perhaps was no longer receptive to the male after the first week of emergence.

Female secretions are known to be involved in the liberation of sperm from spermatophores (Mordue et al., 1980). It may, therefore, be that the mechanism(s) associated with the process of spermatophore depletion/sperm utilization, including the production of the necessary female secretions, was no longer functional after the 7th day of emergence, since the few females that mated at this age (about 20%) did not deplete spermatophores transferred by males.

The lower multiple matings (30–38%) recorded in the
current laboratory study, compared with our field study (70–75%) (Fadamiro et al., unpublished data) may be due to the fact that females were only supplied with virgin males in the current study. In light of the decrease in spermophore volume with subsequent matings in male O. nubilalis (Royer and McNeil, 1993), we suggest that pairing of females with virgin males, as done throughout the present study, might have reduced the tendency of females to remate. Furthermore, females were allowed continuous access to males in this study, only for up to 5 days. It is, therefore, possible that some females after their first mating were still in their refractory period during the remaining window of access to male, and only became receptive again after the 5-th day window. Thus, it is possible that such females could have remated had they been allowed longer access to a male.

Another interesting observation in the current study was that some females depleted only one spermophore, even though they had two spermophores in their bursa at death. It therefore seems unlikely that O. nubilalis females have been selected to mate multiply in order to increase the genetic diversity of their progeny, since one would expect all available spermophores in the bursa to be utilized equally if genetic diversity was being enhanced. Taylor (1967) reported that multiple matings did not contribute significantly to genetic diversity in Atteva punctella.

Unmated females generally lived longer than mated females, and multiple-mated females tended to show reduced longevity compared with single-mated O. nubilalis females. A similar decrease in longevity was recorded for multiple-mated female cabbage looper moth, Trichoplusia ni, compared with once-mated females (Ward and Landolt, 1995). The widespread reduction in the longevity of mated female insects may likely be related to a cost associated with the utilization of female’s resources during reproduction. Pheromone production, calling, courtship and oviposition are processes that require significant energy consumption. The increase in the longevity recorded for sugar-fed mated female O. nubilalis compared with water-fed mated females in the present study is in agreement with the results of Leahy and Andow (1994), and suggests that sugar feeding may significantly replenish females from the depletion of resources associated with reproduction. The higher longevity recorded for females that experienced a 1-week delay in mating was probably due to the fact that such females still retained within them at least half of their estimated total egg complement at death. Hence, their longevity was similar to that of females that had no access to males throughout their lifetime. These results suggest that the resources necessary for somatic maintenance are, perhaps, being utilized during oviposition. Ramaswamy et al. (1997) hypothesized that the increase in longevity recorded for virgin female moths may be due to egg resorption caused by the absence of a male-derived endogenous gonadotropic signal.

In summary, our results suggest that O. nubilalis females may mate multiply in order to enhance reproduction. Therefore, it seems reasonable to expect a significant reduction in the deposition and fertility of eggs with a pheromone-mating disruption program in which a significant decrease in mating frequency of females is achieved. Furthermore, the significant reduction in reproduction in females that experienced a 3- or 7-day delay in mating suggests that an economic control of O. nubilalis population may be achieved by effectively disrupting mating at the beginning of the season. Whether or not such mating reductions can lead to significant economic suppression of damage from this species remains to be seen.

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