Cannibalism in mosquito larvae during microbial larvicide potency tests

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Abstract: We observed instances of cannibalism (intraspecific predation) among intra-instar larvae of Culex pipiens Linnaeus, 1758 while performing a bioassay of Lysinibacillus sphaericus (formerly named Bacillus sphaericus) larvicide, when the larvae were exposed to the larvicide for 48 h in the absence of food. Larvae without symptoms of poisoning attacked and devoured those visibly affected. Cannibalism was more prevalent in 1st–2nd instar larvae than in 3rd–4th instar. This phenomenon should be taken into account when interpreting the results of larvicide bioassays, especially when the exposure lasts over 24 h. The necessity of creating optimal conditions for organisms tested is emphasised.

Key words: bioassay, larval feeding, larval starvation, intraspecific predation, Culex pipiens, Lysinibacillus sphaericus (Bacillus sphaericus)

Microbial larvicides are effective and widely used agents, which may at some point replace chemicals in controlling the aquatic stages of dipteran vectors of human pathogens, such as malaria parasites, filariae and numerous viruses (Margalit et al. 1983, Becker and Margalit 1993, Uspensky 1996, Becker 1998, Regis et al. 2000, Fillinger et al. 2003, Lacey 2007, Berry 2011, Dambach 2018). A variety of factors have been reported to affect the outcomes of laboratory-based bioassays of potential microbial larvicide (Davidson 1985, Becker and Margalit 1993, Lacey 1997, Skovmand and Becker 2000).

At the same time, no special schedule of larval maintenance during bioassay has ever been suggested. In our tests of Lysinibacillus sphaericus (formerly and best known as Bacillus sphaericus) against Culex pipiens Linnaeus, 1758 larvae, we noticed that in some experiments the final numbers of larvae (live and dead) in test dishes were significantly reduced relative to the initial numbers. Upon close observations it became clear that this reduction was caused by some larvae being cannibalised by others. This phenomenon is highly relevant to the design and interpretation of bioassays since it may skew the value of larvicide efficiency if not taken into account, but unfortunately the phenomenon has never been extensively described in the literature (Zaritsky and Khawaled 1986). To determine the prevalence and impact of larval cannibalism, we performed a series of tests under several bioassay conditions.

MATERIALS AND METHODS

Standard preparation of Lysinibacillus sphaericus 2362 larvicide was supplied by the Pasteur Institute, Paris, France. The laboratory colony of Culex pipiens was initiated from the eggs supplied by the Laboratory of Entomology, Ministry of Public Health, Israel. The colony originated in Lower Galilee, Israel, and has been maintained in the Laboratory of Entomology for over ten years. The maintenance of mosquitoes was as described earlier (Uspensky et al. 1998). Two groups of larvae were used in the experiments: the late 1st–2nd and the late 3rd–early 4th instars.

Freeze-dried larvicide was suspended in deionised water (5 mg/ml), homogenised on the Vortex mixer with glass beads (diameter = 3 mm), and diluted in deionised water to the desired concentration just before the experiment. All tests were carried out in Petri dishes (diameter = 7 cm). Larvae were removed from large jars where they were reared with abundant nutrition (powdered dog food supplemented with amino acids) and placed into test dishes with deionised water.

Special attention was paid to the conditions of the larvae, so that only specimens without any signs of morphological or behavioral abnormalities were selected for testing. The larvae were kept in Petri dishes without food for not more than 1 h before the introduction of larvicide. Following our standard procedure (Klein et al. 2002), we defined one larva per 1.0 ml of water volume as the ‘normal’ density, and used 20 larval specimens in 20 ml of water per test dish, unless specified otherwise. Six to eight concentrations of L. sphaericus larvicde were tested, each in five
replicates (i.e., a total of 100 larvae for each concentration). In the first series of tests with *L. sphaericus*, a drop of brewer’s yeast suspension (10% w/w) was added 24 h after toxin introduction and exposure was continued for another 24 h; in the second series, no food was added during the 48 h of exposure to the larvicide. Larval mortality was recorded after 48 h of exposure to the larvicide.

Additional tests were performed with *L. sphaericus* larvicide at a high density of larvae of the 1st–2nd instars (one larva per 0.33 ml of water volume, or 60 larvae in 20 ml per test dish) and no addition of food during the 48-h exposure. Four concentrations of the larvicide were used with four replicates for each concentration (a total of 240 larvae per concentration). Larval mortality was recorded after 48 h of exposure for *L. sphaericus*.

The mean values of larval mortality and cannibalism with standard deviation (SD) were calculated for each concentration of larvicide from all replicates.

Since we paid special attention to the conditions of the larvae selected for the tests, we did not observe appreciable mortality in the control dishes with feeding but without larvicides (the only larva of 1st–2nd instars, which died in a control dish with feeding [Table 1], appears to have been injured during transfer from a jar into the dish). Thus, Abbott’s correction was not applicable to our data. Cannibalised larvae have been counted as dead. The mean value of LC50 and standard deviation were calculated by the log-probit analysis using POLO-PC software.

In the second series of tests (without larval feeding) with normal and high larval density our main interest was connected with cannibalistic larval behavior and the values of LC50 were not calculated. The significance of differences between the values of larval mortality and cannibalism was estimated according to Student’s *t*-test. The difference between values was considered significant if *P* ≤ 0.05.

**RESULTS**

In test dishes with normal larval density (20 larval specimens in 20 ml per dish) where larvae received no food during the 48-h exposure to *Lysinibacillus sphaericus* larvicide, the final numbers of larvae were substantially reduced compared to their initial numbers, sometimes by as much as 50%. In addition to live and dead larvae, we observed significant numbers of fragments of sclerotised body parts (head capsule, thoracic segments), single hairs and setae. Continuous observation showed that after 24–25 hours of exposure, larvae without symptoms of poisoning began to attack the affected larvae ingesting the soft parts of their body (abdominal segments).

Both the number of cannibalised larvae of the 1st–2nd instars and their contribution to total mortality peaked around larvicide concentrations of LC5 to LC50 where they accounted for between 88% and 96% of total mortality (Table 1). At higher concentrations of larvicide, most larvae died within the first 24 h of the test, whereas at lower concentrations cannibalism was rare. We have not observed larvae without symptoms of poisoning falling prey to other larvae. The numbers of cannibalised specimens per dish were much lower among the 3rd–4th instar larvae than among those of 1st–2nd instar, and their prevalence in relation to larvicide concentration followed approximately the same pattern, with absolute numbers increasing from LC5 to LC50, and their contribution to total mortality peaked around LC5 to LC20 at 60% to 65% mortality, and decreased at higher concentrations (Table 1). In control dishes where the larvae had no food for 48 h, all larvae showing signs of starvation fell prey to the cannibals. Interestingly, cannibalism among control larvae in the test without food was higher than among larvae in dishes with a minimal (0.00125 µg/ml) concentration of larvicide (Table 1).

When the same assays were performed at a higher density (60 larval specimens in 20 ml of water per dish) of 1st–2nd instar larvae which received no food, cannibalism was less prevalent than at normal density (Table 2). In contrast to normal density conditions, the numbers of cannibalised specimens increased steadily with increasing the larvicide dose, while their contribution to total mortality remained relatively constant at about 65% in the range of larvicide concentrations higher than LC50. The difference between the number of cannibalised larvae in test dishes with the

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**Table 1. Mortality and cannibalism of larvae of Culex pipiens exposed to Lysinibacillus sphaericus 2362 larvicide under different feeding regimens (mean ± SD)**

| Concentration of larvicide (µg/ml) | 1st–2nd instars | 3rd–4th instars | 1st–2nd instars | 3rd–4th instars |
|-----------------------------------|-----------------|----------------|-----------------|----------------|
| 0.00125                           | 0               | ND             | 1±2.4           | ND             |
| 0.0025                            | 7 ± 7           | 1 ± 2          | 8 ± 5           | 0              |
| 0.005                             | 23 ± 12         | 4 ± 2          | 25 ± 7          | 5 ± 5          |
| 0.01                              | 49 ± 12         | 18 ± 10        | 48 ± 12         | 17 ± 5         |
| 0.02                              | 76 ± 12         | 39 ± 12        | 80 ± 5          | 42 ± 7         |
| 0.04                              | 89 ± 10         | 65 ± 12        | 87 ± 10         | 70 ± 7         |
| 0.08                              | 100             | 91 ± 7         | 99 ± 2          | 93 ± 10        |
| 0.16                              | ND              | 99 ± 2         | ND              | 100            |
| 0.32                              | ND              | 100            | ND              | 100            |
| Control (no larvicide)            | 1 ± 2.4         | 0              | 7 ± 5           | 5 ± 2          |

| Larval mortality (%) in tests with feeding after 24 h of exposure | Cannibalism (%) in tests without feeding for 48 h of exposure |
|------------------------------------------------------------------|-------------------------------------------------------------|
| 1st–2nd instars                                                  | from total number of larvae (mean)                           |
| 3rd–4th instars                                                  | from total mortality (mean)                                 |
| 0.01                              | 0               | 0              | ND             |
| 0.02                              | 3 ± 2           | 38             | 0              |
| 0.04                              | 22 ± 7          | 88             | 3 ± 2          |
| 0.08                              | 46 ± 12         | 96             | 11 ± 2         |
| 0.16                              | 32 ± 10         | 40             | 16 ± 5         |
| 0.32                              | 7 ± 5           | 8              | 15 ± 5         |

LC50 ± SD (µg/ml) 0.010 ± 0.002 0.029 ± 0.005

ND – not detected
different larval density per dish treated by similar concentrations of larvicides was significant (Tables 1 and 2).

**DISCUSSION**

Cannibalism, also known as intraspecific predation, occurs naturally in a wide variety of animals (Fox 1975, Polis 1981, Elgar and Crespi 1992, Toscano et al. 2017), including different taxa of arthropods (Schuausberger 2003, Wilder and Rypstra 2008, Santana et al. 2012). It has been documented in many aquatic animals (Sprules and Bowerman 1988, Hopper et al. 1996, Moksnes 2004), including larvae of mosquitoes of different genera (Mclver and Siemnicki 1977, Reisen and Emory 1976, Clements 1992, Merritt et al. 1992, Renshaw et al. 1993, Koenraadt and Takken 2003, Porretta et al. 2016, Mastrantonio et al. 2018). Larval cannibalism in mosquitoes deserves special attention because of their importance as vectors of dangerous human pathogens. The inter-instar cannibalism, i.e., predation of younger (and smaller) specimens by older (and larger) ones, is more typical of mosquito larvae than the intra-instar cannibalism observed in our experiments. The main factor promoting cannibalism (both inter- and intra-instar) is the absence or deficiency of food, and cannibalistic behaviour is modulated by the hormonal background (El Husseiny et al. 2018).

In our experiments where larvae of *Culex pipiens* were of the same instars, cannibalism was caused by the absence of food for more than 24–25 h and was directed at specimens with symptoms of poisoning who could not avoid or protect themselves from their siblings. Interestingly, the ingestion of visibly poisoned specimens by apparently healthy ones did not increase the total larval mortality in our experiments registered after 48 h of exposure (Table 1).

It would be interesting to observe the fate of predator larvae for a more prolonged time taking into account the phenomenon of *Lysinibacillus sphaericus* spore recycling in mosquito larval cadavers (Becker et al. 1995, Correa and Yousten 1995, Uspensky et al. 1998). In natural water bodies larval cadavers harbouring *L. sphaericus* spores can remain toxic for at least a year (Uspensky et al. 1998). Their persistence may aid the elimination of ‘fresh’ larvae of future generations if the latter feed on poisoned cadavers. The spore recycling was also observed in tests with larval cadavers of several mosquito species after poisoning by another microbial larvicide, *Bacillus thuringiensis israelensis* (Aly et al. 1985, Zaritsky and Khawaled 1986). While it is clear that this phenomenon depends on a variety of factors including larvicide concentration, larval age and stage, type of food and a number of abiotic components, we are not aware of any recent studies addressing the interaction among these variables.

In the 48-hour experiment with high density of the 1st–2nd *C. pipiens* instars treated with *L. sphaericus* larvicide in the absence of food, the frequency of cannibalism was lower than at normal density. Interestingly, if at normal density cannibalism peaked around LC50, decreasing in both directions, at the increased density cannibalism increased in parallel with larval mortality. We hypothesise that the restricted ability of predator larvae to maneuver in overcrowding conditions when the number of poisoned larvae is small limits their chances of encountering potential prey. The importance of encounters ‘favourable’ for cannibalistic attacks was shown by Mastrantonio et al. (2018). When many larvae are poisoned, the ‘healthy’ larvae and perhaps even the minimally poisoned ones have more opportunities to attack and cannibalise the more poisoned specimens even in overcrowded conditions.

Our findings have important methodological implications for assays testing the potency of microbial larvicides. We have shown that at the endpoint of larvicide exposure when the live and dead larvae are scored, some of the initially present larvae may be ‘missing’, having been cannibalised by their siblings. In order to calculate an accurate value of LC50 for a larvicide under testing, the number of any ‘missing’ larvae must be determined and added to the number of dead larvae. Accordingly, in assays of larvicide efficacy, it is essential that the total number of live and dead larvae at the endpoint be compared to their initial number, so that any ‘missing’ specimens could be taken into account. However, the most important point is that optimal conditions of larval maintenance must be supported during the laboratory bioassays. The optimum is specific for any test species and a pesticide tested and it should be determined before the beginning of bioassay.

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