Susceptibility of Field and Laboratory Strains of Cotton Leafworm, Spodoptera littoralis (Boisd.) (Lepidoptera: Noctuidae) to Spinosad Pesticide under Laboratory Conditions

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Abstract

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The susceptibility of field and laboratory strains against all instars larvae of *S. littoralis* to spinosad pesticide after a 24- and 48-h exposure and under laboratory conditions was investigated. As a result against $1^{\rm st}$ instar larvae, the LC $_{50}$ values after 24 h were 12 and 0.275 µg/ml for laboratory and field strain, respectively. In addition, the resistance ratio (RR) of $1^{\rm st}$ instar was 43.64-fold. In this interim, the 48 h LC $_{50}$ values were 8.7 and 0.18 µg/ml for laboratory and field strain, respectively and the RR was 48.33-fold, which revealed the field strain was more susceptible to spinosad than the laboratory strain. Distinctly similar trend was shown for later instar larvae stages. For instance, in $6^{\rm th}$ instar larvae, the LC $_{50}$ values after a 24-h exposure to spinosad were 1100 and 105 µg/ml for the laboratory and field strain, respectively, and the RR value was 10.48-fold. Furthermore, after a 48-h exposure, the LC $_{50}$ values for laboratory and field strains were 500 and 42 µg/ml, respectively, with RR value being 11.90-fold. On the other hand, according to relative tolerance values, the $6^{\rm th}$ instar larvae were the most tolerant instar of all the instars tested. The susceptibility of $6^{\rm th}$, $5^{\rm th}$, and $4^{\rm th}$ instar larvae was comparable and significantly lower than that of $3^{\rm rd}$, $2^{\rm nd}$, and $1^{\rm st}$ instar larvae. However, the $1^{\rm st}$ instar was the least tolerant. The results implied that spinosad may play a potential role in the control of *S. littoralis* and, therefore, it is considered a promising tool in integrated pest management program to control Cotton leafworm which is becoming resistant to conventional pesticides in Egypt.

Keyword: spinosad; Spodoptera littoralis; cotton; insecticide resistance; integrated pest management (IPM)

Cotton leafworm, *Spodoptera littoralis* (Boisd.), is a highly polyphagous pest with numerous hosts causing economically important loses. In Egypt, the Cotton leafworm, *Spodoptera littoralis* (Boisd), is considered one of the major pests attacking more than 112 host plants. Unfortunately, the rate of infestation may reach up to 119 048 egg-mass/ha, causing great damage to leaves, buds, flowers, and bolls (Temerak 2002; El-Sheikh 2012; El-Geddawy *et al.* 2014; Ahmed *et al.* 2015a,b).

The control of Cotton leafworm is complicated due to its high resistance to most of the currently used pesticides classes. Their widely indiscriminate use moreover results in set up into environmental contamination, threat to wildlife populations, and serious public health concerns over food safety (Funderburk et al. 1993; Ahmed 2014; El-Geddawy et al. 2014). Recently, the global occurrence of Cotton leafworm and its growing resistance problem have presented an area of great needs for more effective and acceptable control methods such as alternative safe pesticide with the advantage of its respect to the environment.

On the other hand, integrated pest management (IPM) strives to find the right tactics or combina-

tion of certain tactics to secure the main crop and to minimise the economic crisis. These control tactics include chemical, biological, genetic, culture, and physical controls (Pedigo 1996; Mesbah et al. 2007). In this trend, biopesticides have attracted attention and interest among those concerned to develop an environmentally friendly and safe tool towards the Integrated Crop Management (ICM). Moreover, biopesticides offer a unique opportunity in developing countries to explore and develop their own natural biopesticide resources in the field of crop protection. Such endeavours will assist in conserving foreign cash reserves, improve safety to applicators and consumers, and protect the environment (EL-GEDDAWY et al. 2014). Hence, spinosad is considered a promising biopesticide in controlling many pests regardless of its potent toxicity and low toxicological effects on the environmental components (NAN-NAN et al. 2000; HENDRIX et al. 2001; ARORA 2003; PINEDA et al. 2007).

In this study, we aimed to assess the susceptibility of field and laboratory strains against all instars larvae of *S. littoralis* to spinosad pesticide after a 24- and 48-h exposure and under laboratory conditions.

MATERIAL AND METHODS

Laboratory strain. The laboratory strain of Cotton leafworm, S. littoralis, has been reared in the laboratory of the Plant Protection Department Research, Faculty of Agriculture, Assiut University, Egypt for more than 25 years (without any exposure to chemicals). Insects were reared under controlled conditions in the incubator at $26 \pm 2^{\circ}$ C and $65 \pm 10\%$ relative humidity with 8 light: 16 h darkness photoperiod. Larval jars were supplied with fresh Castor leaves, *Ricinus communis* L., as a source of food which was provided daily. The adults were kept separately and mated on the third day of emergence in clean jars (250 g), fed on 10% honey solution, and fresh green leaves of Tafla, *Nerium oleander* (L.) were provided for egg laying.

Field strain. One field strain of all the eggs masses was collected from different localities in Assiut Governorate, Egypt and reared for one generation under the same laboratory conditions as the laboratory strain described above. However, this field strain was used for all experiments.

Pesticide. The formulation of spinosad used in the bioassay was Spintor[®] (24% SC, registration No. 1050)

R=H Spinosyn A
R=H₃ Spinosyn D

CH₃

CH

Figure 1. The structures of the spinosyn A and D

obtained from Dow AgroSciences Co., Cairo, Egypt. The product is a mixture of two active components, Spinosyn A and D produced by fermentation of the soil actinomycetes, *Sacharopolyspora spinosa* (Figure 1).

Bioassay test. Initially, a pilot test was conducted to choose the range of concentrations used for field or laboratory strain. Spinosad was dissolved in distilled water at different concentrations and leaves of castor bean (approximate radius (r) = 5 cm) were dipped in each concentration for 10 s and left to dry under laboratory conditions. Leaves were put on the bottom of plastic cans covered with a sieved lid. Then 10 larvae of 1st, 2nd or 3rd instar were added, and for 4th, 5th, and 6th instar 5 larvae were used. Four replicates were performed for each concentration and control (leaves dipped in distilled water only). The technique was performed for various instars of field and laboratory strains. The dead larvae were recorded 24 and 48 h after exposure and the percentage of mortality was estimated and corrected according to Abott's formula (ABOTT 1925).

Bioassay data were pooled and analysed (the LC_{50} , LC_{90} , and 95% confidence limit values) according to the methods described by LITCHFIELD and WILCOXON (1949) and SWAROOP *et al.* (1966). However, the resistance ratio (RR) was calculated by dividing the LC_{50} value of the laboratory strain by the LC_{50} value of the field strain. Further, relative tolerance (RT) was calculated by dividing the LC_{50} value of the $6^{\rm th}$ instar larvae of the laboratory strain by the LC_{50} value of the $1^{\rm st}$ instar larvae of the laboratory strain.

RESULTS

The LC_{50} and LC_{90} values for all instars after a 24-and 48-h exposure are shown in Table 1, and RT of various *S. littoralis* larval instars to spinosad after a 24- and 48-h exposure on the basis of LC_{50} values is shown in Table 2.

Initially, the LC_{50} and LC_{90} values for 1^{st} instar larvae after a 24-h exposure to spinosad were 0.275

14.58

3.38 5.92 4.38

2478.60 433.57 5531.40

1700

350

13.33

3.00

3301.05 920.70 9796.38 1308.72

2350

427.14

009 25

laboratory

2th

field

field

 4^{th}

26.39 741.10

76.71 1632.7

98.9

316.72

24

250

13.84 240.01

6.62

69.61 1165.98 144.15 2214.63 149.88

224.45

125

6.13

19.75 510.40 41.63 790.18 67.26

11

10.82

295.57 1672.96

986.58

540

13.68

45.67 842.82 11.90

384.31

240

3500

316.38

500 42

10.48

4.16

4446.54 539.15

0099 540

840

67.39

163.60

1100 105

laboratory

 e^{th}

field

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Table 1. Probit analysis parameters of various larval instars of field and laboratory strain of S. littoralis after being fed on spinosad treated leaves for 24 and 48 hours 12.18 48.33 23.75 20.91 RR_* slope 2.50 2.50 3.42 3.16 533.64 25.47 129.80 7.29 confidence limits 4.51 lower Treated leaves for 48 h 75.04 9.66 16.60 1026.16 249.61 upper (lm/gn) 11.0 62.0 180 7.40 33 9.9 confidence limits 165.88 lower of LC_{50} upper $\frac{11.27}{0.26}$ 23.00 3.62 318.95 2.42 79.04 (lm/gn) 8.7 0.18 19.0 1.56 230 57 2.4 43.64 9.84 20.93 15.20 RR^* slope 3.33 2.77 6.98 3.94 2.11 3.91 30.08 30.82 confidence limits 96.00 718.19 721.34 lower of LC_{90} Treated leaves for 24 h 64.37 75.48 71.67 50.01 535.10 386.30 upper (lm/gn) 1050 1000 58 1.8 0.199 confidence limits 9.221 123.08 274.11 lower of LC_{50} 0.379 15.617 upper 4.68 13.12 526.79 263.24 (lm/gh) 12 0.275 31.5 180 380 laboratory laboratory aboratory laboratory Strain field field field Instar 2nd 3^{rd} 1^{st}

Resistance ratio (RR) calculated by dividing the LC_{50} value of laboratory strain by the LC_{50} value of field strain

Table 2. Relative tolerance (RT) of various S. littoralis larval instars to spinosad after a 24-h and 48-h exposure on the basis of LC₅₀ values

			RT* after	er 24-h exposure	re			RT* a	RT* after 48-h exposure	ıre	
ınstar	nstar Strain —	1^{st}	2 nd	3^{rd}	4 th	5 th	1 st	2 nd	3^{rd}	4 th	5 th
6^{th}	laboratory field		34.92 32.8	6.11 12.12	2.89	1.83 2.33	57.47 233.30	26.32 26.92	8.77 17.50	2.17	1.43
5^{th}	laboratory 50.00 field 163.64		19.05 14.06	3.33 5.23	1.58		40.23 133.33	18.42 15.38	6.14	1.52 2.81	
$4^{ m rh}$	laboratory field		12.06 7.81	2.11 2.91			26.44 61.11	12.10 7.05	4.04 4.58		
3^{rd}	laboratory field		5.71				6.65	3.00			
2 nd	laboratory field	2.63					2.18				

*RT calculated, for example, by dividing the LC₅₀ value of 6th instar larvae of laboratory strain by the LC₅₀ value of 1st instar larvae of laboratory strain

and 0.18 μ g/ml for the field strain, and 12.0 and 58.0 μ g/ml for the laboratory strain, respectively.

However, comparing the LC $_{50}$ and LC $_{90}$ values after a 24-h action of spinosad tested against $1^{\rm st}$ instar of both strains, it was concluded that the field strain is more susceptible (43.64-fold on the basis of LC $_{50}$ value or 32.22-fold on the basis of LC $_{90}$ value) than the laboratory strain. At the exposure period of 48 h, the LC $_{50}$ value of the laboratory strain was 48.33-fold the LC $_{50}$ value of the field strain whereas on the basis of LC $_{50}$ value it was 27.5-fold.

Further, the 24-h LC $_{50}$ and LC $_{90}$ values of spinosad tested against $2^{\rm nd}$ instar larvae of field strain were 3.2 and 44.0 µg/ml, and for laboratory strain, the values were 31.50 and 120.0 µg/ml, thus the rate of resistance of laboratory strain was 9.84-fold on the basis of LC $_{50}$ and 2.73-fold on the LC $_{90}$ basis, as compared with field strain. Comparing the LC $_{50}$ value of $2^{\rm nd}$ instar larvae with that of $1^{\rm st}$ instar larvae, the $2^{\rm nd}$ instar is more tolerant (more than 10-fold in the field strain and more than 2-fold in the laboratory strain).

In regards to the LC $_{50}$ value of spinosad that was tested against field strain, the value was 8.6 µg/ml as compared with the LC $_{50}$ value of laboratory strain (180.0 µg/ml) indicating that laboratory strain is 20.73-fold more resistant than field strain. The LC $_{90}$ value for field strain was 47.0 µg/ml while for laboratory strain it was 1050.0 µg/ml. On the basis of LC $_{90}$ value, laboratory strain is 22.3-fold more resistant as compared with field strain. The 24 h LC $_{50}$ value of spinosad tested against $3^{\rm rd}$ instar larvae of field strain was more than two-fold than that for $2^{\rm nd}$ instar larvae, whereas in the laboratory strain LC $_{50}$ value for $3^{\rm rd}$ instar larvae was more than fold the LC $_{50}$ value of $2^{\rm nd}$ instar larvae.

Furthermore, the 24-h LC $_{50}$ and LC $_{90}$ values of spinosad action on $4^{\rm th}$ instar larvae of laboratory strain were 380 and 1000 µg/ml and for field strain the values were 25 and 540 µg/ml. Laboratory strain showed 15.2-fold higher resistance than field strain on the basis of LC $_{50}$ value, but it was only 1.85 µg/ml based on LC $_{90}$ value. After a 48-h exposure, the LC $_{50}$ value of spinosad against laboratory and field strain was 30 and 11 µg/ml, whereas the LC $_{90}$ value was 1026.16 and 284.45 µg/ml, respectively. The laboratory strain showed 20.9- and 5.92-fold resistance than the field strain according to LC $_{50}$ and LC $_{90}$ values.

On the basis of a 24-h LC $_{50}$ value of spinosad, $4^{\rm th}$ instar larvae of laboratory strain showed 31.67-, 21.06-, and 2.11-fold resistance as compared with $1^{\rm st}$, $2^{\rm nd}$, and $3^{\rm rd}$ instar larvae, respectively. For field strain the resistance was 90.91-, 7.81-, and 2.91-fold, respectively.

Interestingly, the slope values for laboratory and field strains after a 24- and 48-h exposure indicate that field strain is more homogeneous in response to spinosad than laboratory strain.

The 24-h LC $_{50}$ value was 600 and 45 µg/ml for laboratory and field strains, whereas the LC $_{90}$ value in respective was 2350 and 540 µg/ml. After a 48-h exposure, the LC $_{50}$ values were 350 and 24 µg/ml and LC $_{90}$ values were 1700 and 250 µg/ml. Based on the 24-h LC $_{50}$ value of laboratory and field strain, it might be concluded that laboratory strain was 13.33-fold more resistant than field strain.

The 24-h LC₅₀ value of spinosad tested against $5^{\rm th}$ instar larvae of laboratory strain was 50-, 19.05-, 3.33-, and 1.58-fold that of $1^{\rm st}$, $2^{\rm nd}$, $3^{\rm rd}$, and $4^{\rm th}$ instar larvae, respectively. For field strain, the values were 163.64-, 14.06-, 5.23-, and 1.8-fold, respectively (Table 2). The $5^{\rm th}$ instar larvae were significantly more resistant than $1^{\rm st}$ and $2^{\rm nd}$ instars, and slightly more tolerant than $3^{\rm rd}$ and $4^{\rm th}$ instars.

The relatively high slope value for field strain in both the 24- and 48-h exposure revealed a more homogeneous response of field strain to spinosad than laboratory strain. The 24-h LC $_{50}$ value of spinosad tested against $6^{\rm th}$ instar larvae was 1100 and 105 µg/ml for laboratory and field strain giving 10.48-fold RR. The 24-h LC $_{90}$ values were 6600 and 840 µg/ml for laboratory and field strains with 7.86 RR. The 48-h LC $_{50}$ values were 500 and 42 µg/ml for laboratory and field strains with resistance rate of 11.90, whereas the 48-h LC $_{90}$ values were 3500 and 240 µg/ml for the two strains with a 14.58-fold RR. Comparing the slope value after the 24- and 48-h exposure, it seems that $6^{\rm th}$ instars of both strains exhibit similar response to spinosad.

The 24-h LC $_{50}$ value of spinosad tested against $6^{\rm th}$ instar larvae of laboratory strain was 91.7-, 34.92-, 6.11-, 2.89-, and 1.83-fold the LC $_{50}$ value of $1^{\rm st}$, $2^{\rm nd}$, $3^{\rm rd}$, $4^{\rm th}$, and $5^{\rm th}$ instar larvae, respectively. For field strain, RT was 381-, 32.8-, 12.21-, 4.2-, and 2.33-fold, respectively. The same trend was observed in 48-h exposure results (Table 2). The $6^{\rm th}$ instar larvae of both strains showed a higher significant tolerance to spinosad than $1^{\rm st}$ and $2^{\rm nd}$ instars, a moderate tolerance compared to $3^{\rm rd}$ instar. However, in response to spinosad $5^{\rm th}$ instar was comparable with $6^{\rm th}$ instar.

DISCUSSION

Generally, spinosad showed great effects against *S. littoralis* larvae. However, the effect was significant

on field strain in comparison with laboratory strain. The LC₅₀ value of spinosad effect on 1st instar after a 48-h exposure was 0.18 and 8.7 μ g/ml for field and laboratory strain, respectively. Furthermore, the value increased in the successive instars to reach 500 and 42 μg/ml for 6th instar larvae of laboratory and field strains, respectively. For all instars, and at a 24- and 48-h exposure, the field strain was more susceptible than laboratory strain. Interestingly, the resistance rate was much higher in 1st, 3rd, and 4th instars. The reasons behind the high level of insensitivity of the laboratory strain to spinosad could be the lacking exposure of the laboratory strain to natural microorganisms or the lack of the exposure towards the severity of pesticides selective pressure in comparison to field strain which leads to a resistant trend against spinosad the active ingredient of which is a microorganism. Therefore, the laboratory strain may invalidate their ability to the microbial pesticides or biopesticides such as spinosad. Plus, the reasons could be xenobiotic metabolism changes or altered toxicokinetics and behaviour differences. Further, there are some correlations between the high level of resistance to spinosad and various kinds of insecticides, especially those having a similar mode of action (e.g. neonicotinoids, sulfoximine, and nereistoxin analogs) which lead to cross-resistance. However, in agreement with the present findings, TEMERAK (2002) found that spinosad was more active against field strain of *S. littoralis* larvae than laboratory strain. The LC₅₀ values of spinosad against 1st-6th instar larvae of field strain were 0.54, 1.19, 1.866, 18.17, 40.5, and 61.02 μg/ml, however, for laboratory strain they were 4.84, 10.9, 66.86, 1559.63, 2690.39, and 4013.23 µg/ml, respectively. The LC₅₀ values for the last three instars of laboratory strain (4th, 5th and 6th) were much higher than those recorded in the present study whereas; in field strain the reverse was noticed. This may be due to the variation in the strain and the environmental condition prevailing in the area. In contrast, AYADIN and GÜRKAN (2006) evaluated lethal dose bioassays of spinosad on 3rd instar larvae of S. littoralis using the leaf dip method. The LC₅₀ values for field and susceptible strains were 43.691 and 10.037 µg/ml, respectively. The field strain was approximately 4.4-fold less sensitive than the susceptible strain. SAUNDERS and BRET (1997) stated that spinosad undergoes photodegradation when exposed to sunlight and is rapidly metabolised when washed into soil. On the other hand, the variation in the result of spinosad toxicity may be due to the

application techniques used. REDDING and NEAD (1998) found that using hollow cone nozzles (TX6) with 41.3.7 kPa (60 psi) for the application of tracer provided better coverage control as compared with the same type of nozzles at lower pressure (27.3.5 kPa or 40 psi) or different nozzles (TX15 and 8003 flat fan) at the same or lower pressures. The higher tolerance of laboratory strain than of field strain was confirmed by many authors. During their research on natural product, ABO-ELGHAR et al. (1994) found that laboratory strain of cotton leafworm was more tolerant to Bacillus thuringiensis and Abamectin (fermentation of the actinomycete, Streptomyces avermitilitis) than field strain. In spinosad bioassays, Mascarenhas et al. (1998) demonstrated that field strains of Spodoptera exigua had significantly lower LC₅₀ values than reference strain. MOULTON et al. (1999) found that field population of 2nd and 3rd instar of Spodoptera exigua was 3- to 70-fold less susceptible to spinosad than a reference laboratory population. Against, field collected strains of the soybean pest, Pseudoplusia includens. MASCARENHAS and BOETHEL (1997) found in spinosad bioassay that field strain had lower LC₅₀ than the susceptible USDA reference strain.

In conclusion, spinosad showed variable degree of toxicity against *S. littoralis*. According to the present investigation and the available literature, spinosad proved to be the most active biopesticide against cotton pests. Its efficacy was comparable to that of synthetic insecticides.

One essential key of the pest management strategy is the use of safe and alternative products to ensure that continual selectivity will not occur and that any possibility of pest resistance is avoided, or at least significantly delayed. The naturalyte insect control class represented by spinosad provides the potential option because the members of this class show no cross-resistance to other product classes including pyrethroids, carbamates, organophsophates, and even newer classes, such as fipronils, imidaclopid, and avermectins. Biochemical and molecular biological investigation should follow to better elucidate the mode of action of spinosad on *S. littoralis*. Thus spinosad has gained a great interest especially after the establishment of organic farms in Egypt.

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