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BIOEFFECTS OF THE ENTOMOPATHOGENIC FUNGI BEAUVERIA BASSIANA (BALS.) ON MUSHROOM FLY BRADYSIA OCELLARIS* (COMS.) (DIPTERA: SCIARIDAE)

[13]

Dawalibi¹, W.A.M.; S.M.T. Khoja¹; M.M. Abou- Shaar² and N.A. Kaake²

- 1. General Commission for Scientific Agricultural Research (Gcsar), Aleppo, Syria
- 2. Faculty of Agriculture, Aleppo University, Aleppo, Syria

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ABSTRACT

Laboratory experiments were carried out to determine the bioeffects of an isolate of entomopthogenic fungi Beauveria bassiana (Bals.) and Biofly a commercial product of B. bassiana, on the 1st instar larvae of mushroom fly. Bradysia ocellaris (coms). The bioactivity of B. bassiana was tested, using five concentrations of B. bassiana on some biological criteria of the mushroom fly, by calculating LC50 values after three and seven days of treatment. Results indicated that the mortality rates percentage increased with the increase of the concentrations used and the period after treatment. The highest percentage of mortality occurred within the first seven days following treatment. Statistical analysis of the obtained larval-pupal and adults period and weight revealed significant differences between treated and nontreated insects.

INTRODUCTION

In recent years, an increasing consumer demand for commercial mushroom products, as natural healthy foods has raised concerns about problems in mushroom production. Mushroom flies proved to be a serious pest facing mushroom growers (Finley et al 1984; Ishitani, et al 1997; Jess and Kilpatrick, 2000). The existing control measures against the fly still rely on chemical insecticides, which are not always appropriate. The use of pathogens may offer an environmentally

sound method for the management of insect pests. Hyphomycete fungi are the most promising candidates (Prior, 1990).

Entomopathogenic fungi have been intensively studied as potential microbial insecticides (Gillespie 1988; Ferron et al 1991; Malsam et al 1998). Beauveria spp. has been investigated because of their ability to infect a wide range of insects. There are more than 400 species recorded as entomopathogenic fungi, from which only about 20 species have the potential to be used in microbial control of insect pests (Zimmerman, 1986).

The present work was carried out with the endeavor of evaluating the bioassay of *Beauveria bassiana* against the mushroom fly *Bradysia ocellaris* and to evaluate the bioactivity effect of *Beauveria bassiana* on some biological criteria of mushroom fly *B. ocellaris*.

MATERIALS AND METHODS

Rearing mushroom fly

Mushroom fly *B. ocellaris* was reared under laboratory condition at 23=1°C in ventilated plastic boxes (28×16×9 cm) filled with peat to a depth of about 5 cm. About 10 ml of water was mixed into the peat so that the mixture will remain moist to the touch and 1g of wheat was distributed over the surface of peat as food source for the larvae. Adult mushroom flies *B. ocellaris* were put into the boxes and an additional 1g of wheat grain was added every 3-4 days together with a cotton plug soaked in a 20% glucose to boost the number of eggs laid by females (Gouge 1994).

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^{*}Bradysia ocellaris was identified at the Department of Entomology in the Natural History Museum. London

Two ELISA dishes were used for each treatment; every well of the bottom ELISA dish contained one wheat grain covered with spawn and one larvae of Bradysia ocellaris. The dishes were treated with different concentrations of B. bassiana $(10^2, 10^3, 10^4, 10^5, 10^6)$ conidia/ml, control treatment was treated with 0.1%Tween alone. In the well of the upper dish, a piece of cotton moistened with similar concentrations of B. bassiana and with sterilized water + 0.1%Tween 80 in the control treatment. Dishes were then put inside moist boxes containing moistened filter paper. The dishes were incubated at $23\pm 1^{\circ}$ C⁷ 70 ± 5 % RH and 24h (dark). Mortality was recorded daily for about 14 days. Dead larvae were removed and maintained in a moist chamber to allow for fructification of the relevant fungus

Bioassay experiments

In this study, the entomopathogenic fungus B. bassiana, (S1), isolated from adults of Eurygaster integriceps pest and (Biofly), a commercial product of B. bassiana, were assayed for their bioinsecticidal activities against 1st instar larvae of the B. ocellaris. The first isolate was grown and maintained on Potato Dextrose Antibiotic Agar (PDAA) and incubated in the dark for 20 days at 25±1 °C for the actual bioassay, conidial suspensions of each of the isolates were prepared by scraping conidia from the surface of 20 days old cultures in 0.1% Tween 80. The suspension was diluted in sterilized water ± 0.1% Tween 80 to get suspensions ranging from 1×10^2 to 1×10^6 conidia / ml. A total of 10 ml suspension was prepared for each dilution.

Larvae treatment

The experiments were carried out under laboratory conditions of $23\pm1^{\circ}$ C and 80 ± 5 RH. Five concentrations of B. bassiana 10^2 , 10^3 , 10^4 , 10^5 and 10^6 conidia/ml were used. In all experiments percentage mortality was calculated after 3, 5, 7, 10, and 14 days. Percentage of mortality of each treatment was corrected use using Abbotts formula (Abbott, 1925). The percentages of mortality were statistically computed according to (Finney, 1952) and computed mortality percentages were plotted versus log concentrations on logarithmic probit paper to obtain the corresponding regression mortality lines. The concentrations required to give 50% mortality (LC50) were estimated from the established regression lines (Jyoti

and Brewer, 1999). Each concentration test included 3 replicates each of 24 larvae (72 larvae/concentration, 432 larvae/treatment).

Biological studies: 1st instar larvae were treated in the same manner with the determined LC50 to study the effect of the entomopathogenic fungus *B. bassiana* and Biofly on certain biological aspects of the mushroom fly. Five replicates of 1st instar larvae (40 larvae/replicate) were used for each compound. Larval and pupal mortalities, their durations, weight and moth emergence were recorded. The experiments were carried out under laboratory conditions of 23±1C° and 80±5 RH.

Statistical analysis

ANOVA was performed using SAS software program.

RESULTS AND DISCUSSION

Bioeffect of an isolates of the entomopathogenic fungus *B. bassiana* and Biofly on mushroom fly *B. ocellaris* larvae

Results in **Table (1)** show the mortality percentage of 1st instar larvae of mushroom fly after treatment with different concentrations of *B. bassiana*. The tested *B. bassiana* had a great efficacy against mushroom fly larvae. Also results indicate that the mortality rates increased with the increase of the concentration used and the period after treatment.

The corrected mortality percentages after 3 days for S1 isolate treatment ranged from 8.9% using the lowest concentration (10² conidia/ml), to 38.82% using the highest concentration (10⁶ conidia/ml). As treatment, the corrected percentages of mortality ranged from 4.3 to 29.34 % for the lowest and highest concentrations, respectively for Biofly. These percentages increased to 50.6, and 19.06% after 7 days of treatment for S1 and bio fly, respectively. In contrast, the mortality percentages recorded for the highest concentration (10⁶ conidia/ml) were 82.06, 64.61% after 7 days, for the treatment with S1 and bio fly) respectively. Also the highest cumulative mortality (85.06, 69.6%) was recorded after 14 days of treatment with 10° conidia /ml of (S1, biofly) respectively Table (1). The concentration mortality lines are graphically illustrated in Figs. (1 and 2) and showed a positive relationship between larval mortality and the concentrations used. It is worth

Table 1. Corrected mortality percentages for the 1st instar of mushroom fly *Bradysia ocellaris* larvae treated with an isolate (S1) of the entomopathogenic fungus *Beauveria bassiana* and Biofly

	Mean of corrected mortality %							
Treatment	Concentration conidia/ml	l lays after treatment						
Isolate(S1)		3	5	7	10	14		
	·10 ⁺²	8.9	32.86	50.6	56.7	56.7		
	10 ⁺³	16.4	40.27	53.70	64.12	64.12		
	10 ⁺⁴	31.26	58.21	68.66	71.65	71.65		
	10 ⁺⁵	37.27	61.11	73.14	76.04	76.04		
	10+6	38.82	67.13	82.06	82.06	85.06		
LC50				0.95×10^{2}	0.9×10^{2}			
R				0.52	0.55			
Bio fly*								
	10^{+2} 10^{+3}	4.3	14.6	19.06	24.89	24.8		
	10^{+3}	7.30	17.58	26.37	29.3	29.34		
	10 ⁺⁴	14.6	26.37	30.8	33.79	33.79		
	10 ⁺⁵	13.18	35.16	45.5	48.4	48.4		
	10 ⁺⁶	29.34	47.03	64.61	69. 6	69.6		
LC50				5.7×10 ³	4.55×10 ⁵			
R				0.83	0.79			

^{*}Biofly: Commercial protect of B.bassiana

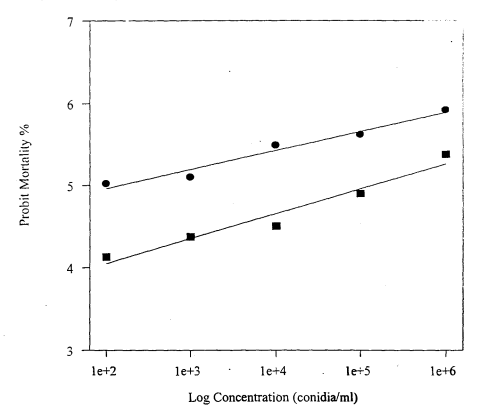


Fig.1. Toxicity lines of isolate S1 and Biofly on mushroom fly Bradysia ocellaris (Coms.) after 7 days following treatment

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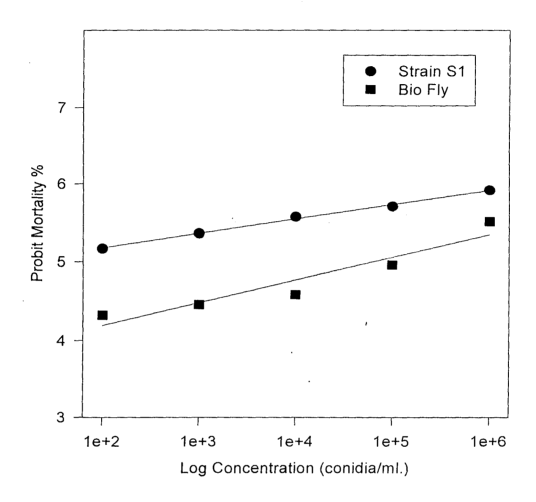


Fig. 2. Toxicity lines of isolate S1 and Bio fly on mushroom fly *Bradysia ocellaris* (Coms.) after 10 days following treatment

mentioning that the highest percentage of mortality occurred within the first 7days following treatment, while Larval mortality was relatively lower after 10 days (Table 1).

Effect of LC50 concentrations of an isolate of the entomopathogenic fungi *B. bassiana* and Biofly on larval, pupal and adult stages mortality of mushroom fly *Bradysia ocellaris*

Data in **Table (2)** show that the LC₅₀ concentrations of the entomopathogenic *B. bassiana* and

Biofly which killed 50% of the treated larvae (after 7 days of treatment) had great effects on larval-pupal periods and adult longevity. Statistical analysis of the obtained larval-pupal periods and pupal weight revealed significant differences between treated and non-treated insects. The larval, pupal periods and adult longevity were 11.2, 5, 6.1, days respectively (non-treated). Application of the entomopathogenic fungi *B.bssiana* and Biofly shortened larval and pupal development and adult longevity, which were 8.2, 4.2, 4.2- 8.4, 4.4, 4.4 days after treatment respectively.

Treatment	Mean larval duration period days	Mean larval weight (mg)	Mean pupal duration period (days)	Mean pupal weight (mg)	Mean adults long (days)	No of pupal*
S1	8.2 ^b	0.35 ^b	4.2 ^b	1.08 ^b	4.2 ^b	22.4 ^b
Biofly	8.4 ^b	0.29 ^b	4.4 ^{ab}	0.84 ^b	4.4 ^b	18.8°
control	11.2ª	1.54ª	5 ^a	3.55ª	6.1²	38.6ª
CV	6.8	21.1	12.7	12.9	13.4	9.4
F	35.2	105.7	2.6	204.7	12.6	88.7
SE =	0.4	0.02	0.33	0.06	0.43	6.2
LSD.50	0.87	0.21	0.8	0.32	0.91	3.4

Table 2. Effect of treating 1st instar larvae of the mushroom fly *Bradysia ocellaris* with LC₅₀ concentration of the *Beauveria bassiana* (Bals.) and Biofly on larval and pupal periods, pupal weight and adult longevity

• No. of larvae = 40 larvae.

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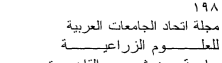
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التأثير الحيوي للفطر الممرض للحشرات (Bals.) على ذبابة الفطر Bradysia ocellaris (Coms.)

[14]

وجيه أحمد منير دواليبي' - سليم محمد طاهر خوجه' - محمد مصطفى أبو شعر' - نوال عبد القادر كعكة'
۱ - الهينة العامة للبحوث العلمية الزراعية - مركز البحوث الزراعية بحلب - سوريا
۲ - كليــة الزراعــة - جامعــة حلـب - حلـب - سوريــا

الموجيز

أجريت تجارب معملية لتحديد التأثير الحيوي الفطر الممرض للحشرات Beauveria bassiana وكذلك المستحضر التجاري منه Biofly في العمر الأول الذبابة الفطر الزراعي Biofly في العمر كما اختبرت الفعالية الحيوية للفطر على بعض الصفات الحيوية لذباب الفطر الزراعي وذلك بعد تحديد قيمة $1C_{50}$ بعد ثلاثة وسبعة أيام من المعاملة. واستخدمت في هذه الدراسة خمس تراكير للفطر الفطر على وجود علاقة

إيجابية ما بين التركيسز المستخدم ونسبة موت اليرقات، حيث لوحظت زيادة في معدل نسبة موت البرقات بزيادة التراكيز المستخدمة وطول الفتسرة بعد المعاملة. كما لوحظت زيادة في نسبة موت البرقات خلال الأيسام السبعة الأولى التسي تلت المعاملة، وقد أظهر التحليل الإحصائي وجود فروقات معنوية لطول فترة الطور البرقي والعسذراء وعمر الحشرة الكاملة وأوزانها، بين المعامل وغير المعامل.

تحكيم: ا.د مديحة ابو المكارم ا.د أحمد رؤوف حامد