



Fumigant Toxicity of the Essential Oils from Medicinal Plants Against Bean Weevil, *Acanthoscelides obtectus* (Say) (Coleoptera: Bruchidae)

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Fumigant activity of essential oil vapours from the aromatic plants such as, wild thyme (*Thymus serpyllum* L., Lamiaceae), origanum (*Origanum onites* L., Lamiaceae), rosemary (*Rosmarinus officinalis* L., Lamiaceae), basil (*Ocimum basilicum* L., Lamiaceae) and lemon balm (*Melissa officinalis* L., Lamiaceae) against the bean weevil *Acanthoscelides obtectus* (Say) adults under laboratory conditions. Volatile phase effects of different concentrations of the essential oils were used to determine insecticidal activities. Laboratory bioassay results indicated that all essential oils were found to cause 100 % adult mortality of *A. obtectus* in a dose-dependent manner. Essential oils of thyme and origanum had the highest insecticidal effects, causing high adult mortalities at the lower concentration ($10 \mu\text{g mL}^{-1}$ air) in comparison to other essential oils tested. Adult viability was totally affected by essential oils of rosemary, basil and lemon balm at the concentrations of 20, 30 and $30 \mu\text{g mL}^{-1}$ air, respectively. The estimated LC_{50} (lethal concentration 50) values obtained for each essential oil were calculated by using probit analysis. The lowest LC_{50} values were recorded for thyme essential oil ($1.12 \mu\text{g mL}^{-1}$) was followed by origanum ($1.31 \mu\text{g mL}^{-1}$), rosemary ($2.66 \mu\text{g mL}^{-1}$), basil ($3.10 \mu\text{g mL}^{-1}$) and lemon balm ($3.60 \mu\text{g mL}^{-1}$) respectively. The results of the present study concluded that plant essential oils could be useful in promoting research aiming at the development of new agent for pest control from the plants with medicinal values.

Key Words: *Acanthoscelides obtectus*, Stored pest, Essential oil, Insecticidal, Biopesticide.

INTRODUCTION

Stored products of agricultural and animal origin are attacked by more than 600 species of beetle pests, 70 species of moths and about 355 species of mites causing quantitative and qualitative losses¹ and insect contamination in food commodities is an important quality control problem of concern for food industries. Among the insects attacking stored products, members of Bruchidae have attracted the attention of many scientists not only because they can easily be manipulated but also because of the economic importance they have. Bean weevil, *A. obtectus* with its physiological adaptability, is one of the most destructive pests of *Phaseolus vulgaris* L. (kidneybean) which is one of the most important food pulses in the Mediterranean countries. Pest damage by this species has been estimated to cause 20-40 % loss of stored seeds per annum².

Fumigation plays a major role in insect pest elimination in stored products. Previously, phosphine and methyl bromide were the two common fumigants used for stored-product protection world wide. Insect resistance to phosphine is a global issue now and control failures have been reported in field

situations in some countries^{3,4}. Methyl bromide, a broad-spectrum fumigant, has been declared an ozone-depleting substance and therefore, is being phased out completely. In view of the problems with the current fumigants, there is a global interest in alternative strategies including development of chemical substitutes, exploitation of controlled atmospheres and integration of physical methods⁵.

Plant essential oils and their major components, various monoterpenoids, show a broad spectrum of activity against insects, mites and various arthropods^{6,7}. These compounds act as antifeedants and repellents while they may also affect some biological parameters such as growth rate, life span and reproduction⁷⁻¹². Their vapour action may also be very promising for pest control because of their insecticidal properties and the fact that they can act as fumigants.

The objective of the present work was to investigate the fumigant activity of essential oil vapours from the aromatic plants wild thyme (*Thymus serpyllum* L., Lamiaceae), origanum (*Origanum onites* L., Lamiaceae), rosemary (*Rosmarinus officinalis* L., Lamiaceae), basil (*Ocimum basilicum* L., Lamiaceae) and lemon balm (*Melissa officinalis* L., Lamiaceae) against *A. obtectus* adults.

TABLE-1
EFFECTS OF DIFFERENT CONCENTRATIONS OF VOLATILE PHASES OF
ESSENTIAL OILS ON PERCENT MORTALITY OF *A. obiectus* ADULTS

Concentrations ($\mu\text{g mL}^{-1}$ air)	Essential oils and % mortality				
	<i>T. speyllum</i>	<i>O. onites</i>	<i>R. officinalis</i>	<i>O. basilicum</i>	<i>M. officinalis</i>
0.0	0,0a	0,0a	0,0a	0,0a	0,0a
1.0	46,7bB	40,0bB	23,4bA	23,3bA	20,0bA
2.0	73,3cB	66,6cB	43,3cA	40,0cA	36,7cA
5.0	96,7dB	93,2dB	63,2dA	56,7dA	53,3dA
10.0	100,0dB	100,0dB	80,0deA	73,3eA	66,6eA
15.0	100,0dB	100,0dB	93,3efAB	86,6fAB	83,4fA
20.0	100,0dA	100,0dA	100,0fA	96,7gA	96,6gA
30.0	100,0d	100,0d	100,0f	100,0g	100,0g
LC ₅₀	1,12	1,31	2,66	3,10	3,60
LC ₉₀	3,19	3,83	11,96	16,67	19,14

Mean values (n = 3 replicates with ten female adult insects per replicate) followed by the same small or capital letters within the row or column, respectively, are not significantly different according to Duncan multiple range test ($P \leq 0.05$); The estimated lethal concentrations (LC₅₀ and LC₉₀) values ($\mu\text{g mL}^{-1}$) for each essential oil were estimated by using Probit analysis

EXPERIMENTAL

Acanthoscelides obiectus adults were obtained from laboratory cultures maintained on bean, *Phaseolus vulgaris* L., at $25 (\pm 1)^\circ\text{C}$, $65 (\pm 5) \%$ relative humidity and a photoperiod of 8:16 h light: dark. The insects were collected from a warehouse in Hatay province and the beans were purchased from the local market and maintained in a freezer at -20°C .

The plants used in this study were identified by Dr. I. Uremis. A voucher specimen has been deposited in the herbarium of the plant protection department, MKU (No. TspyYD7, OoAS56, RoBK40, ObRY15 and MelAM23). For the extraction of essential oils, plants were collected from Yayladagi, Serinyol, Reyhanli and Antakya districts of Hatay provinces situated in the Eastern Mediterranean region of Turkey. Air-dried plant material (200 g) was placed in a 5 L round-bottom distillation flask and 3 L double distilled water added. The essential oils were obtained by steam distillation for 3 h using Clevenger-type apparatus (Ildam, Ankara). The oils were separated, dried over anhydrous sodium sulphate and stored in an amber bottle at 4°C until used.

Toxicity tests: Transparent acrylic cups (6 cm height and 3.5 cm diameters, which offer 50 mL air space) were used as test chambers for determination of volatile phase effects of the essential oils. Ten bean seeds without insects were individually placed into the chamber. Ten adult insects were introduced in each cup and allowed to settle for half an hour before exposure to essential oil. The top of the insect chamber was covered by using the lids. The essential oils were applied with a micropipette on a filter paper strip (3 cm $\times$ 1 cm) attached to the lids. Different concentrations of essential oil were prepared by dissolving the requisite amounts in sterile dimethyl sulfoxide (DMSO) solution. Essential oils were diluted from the concentration of 15000 μg to 500 $\mu\text{g mL}^{-1}$ in DMSO and a 10- μL aliquot of each concentration was added on the inner surface of the lid of test chamber with a micropipette giving concentrations of 1, 2, 5, 10, 15, 20 and 30 $\mu\text{g mL}^{-1}$ air. Insect chambers were sealed immediately with parafilm to prevent loss of essential oils from the chamber. Three replications were made

for each concentration. As untreated control, three cups containing only 10 μL of DMSO were used.

The treated insect-chambers were returned to the incubator set at 25°C , 60-65 % relative humidity and a photoperiod of 16:8 (L:D) h. Mortality was determined under a dissecting microscope 24 h after treatment. Adult insects were assessed for 30 min for mobility and for 2 h for mortality, defined by lack of response to stroking with a paintbrush.

Data analysis: Mortality observations were analyzed by using the SPSS program, version 11.5, for ANOVA. Tukey's test was used to compare means. Probit analysis was used to determine lethal concentrations (LC₅₀), by using the SPSS program, version 11.5. Abbott's formula was used to correct mortality in controls.

RESULTS AND DISCUSSION

The volatile phase effects of different concentrations of essential oils on the mortality of *A. obiectus* adults are shown in Table-1. All essential oils were found to cause 100 % adult mortality of *A. obiectus* in a dose-dependent manner. Essential oils of thyme had the highest insecticidal effects, causing high adult mortalities at the lower concentrations in comparison to the other essential oils tested. Adult viability was totally affected by thyme, origanum, rosemary, basil and lemon balm at the concentrations of 10, 10, 20, 30 and 30 $\mu\text{g mL}^{-1}$ air, respectively. The LC₅₀ (lethal concentration 50) was the concentration of the essential oil, which kills 50 % of adult insect within 24 h following exposure. The estimated LC₅₀ values obtained for each essential oil were calculated by using probit analysis (Table-1). The lowest LC₅₀ values were recorded for thyme essential oil (1.12 $\mu\text{g mL}^{-1}$) was followed by origanum (1.31 $\mu\text{g mL}^{-1}$), rosemary (2.66 $\mu\text{g mL}^{-1}$), basil (3.10 $\mu\text{g mL}^{-1}$) and lemon balm (3.60 $\mu\text{g mL}^{-1}$).

Botanical insecticides have long been recommended as attractive alternatives to synthetic chemical insecticides for pest management because these chemicals pose little threat to the environment or to human health⁷. In previous studies, volatile compounds from essential oils were reported to possess antimicrobial activity against a variety of food borne, human

and plant pathogens and pest¹². Pyrethrum and neem are well established commercial pesticides based on plant essential oils and recently entered the marketplace.

In previous study, insecticidal efficacy of essential oils, derived from taxonomically different medicinal plants species, such as *Salvia officinalis*, *Cuminum cyminum*, *Anethum graveolens*, *Mentha spicata spicata*, *Micromeria fruticosa* and *Ocimum minimum* were investigated against three important stored product insects, including *A. obtectus*, *Sitophilus granarius* and *Sitophilus oryzae* under laboratory conditions¹³. In their results the lowest LC₅₀ value against *A. obtectus* was recorded for *C. cyminum* essential oil (0.029 µL mL⁻¹). In another study, the chemical composition and insecticidal action of the essential oils isolated from various parts of three aromatic plants (*Lavandula hybrida* Rev, *Rosmarinus officinalis* L and *Eucalyptus globulus* Labill) was determined on *A. obtectus* adults and their LC₅₀ values were estimated by Papachristos *et al.*¹⁴. Several authors have reported that all essential oils tested exhibited strong activity against *A. obtectus* adults, with varying LC₅₀ values depending on the essential oils. Greater susceptibility of *A. obtectus* to essential oils and monoterpenoids has also been reported^{2,14,15}. In another study, which was conducted by Karaborklu *et al.*¹⁶, the mean longevity of *A. obtectus* adults was significantly decreased depending on the increasing doses of savory, myrtle, marjoram, laurel, lemon, tansy and sage essential oils.

To our knowledge, our results is the first study demonstrating that the essential oils of *T. serpyllum*, *O. basilicum* and *M. officinalis* possess insecticidal activities against *A. obtectus*.

The number of compounds and their relative amount found in plant essential oils varied according to plant species and the particular compound. The major compounds found in essential oils of thyme, origanum, rosemary, basil and lemon balm were carvacrol, carvacrol, borneol, linalool and citronellal respectively¹⁷⁻²⁰. These compounds have previously been reported to have antimicrobial activity against a variety of insects, mites, weeds and plant pathogens^{12,21,22}.

Major components of essential oils such as monoterpenoids, carvacrol, 1,8-cineole, *p*-cymene, menthol, γ -terpinene, terpinen-4-ol and thymol were investigated against three stored product insects^{15,23-25}. Authors have reported that these individual components possess insecticidal, fecundity, antifeeding and oviposition deterrent effect against different major stored product insects^{26,27}. In this aspect, insecticidal activities of essential oils used in our study on adult bean weevil mortalities could be due to the fumigant toxicities of the major components of essential oils.

In previous studies, in addition to insecticidal activities, essential oils derived from medicinal plants possess antifungal, antibacterial, herbicidal and acaricidal activities against plant pathogenic fungi, bacteria, weeds and mites²⁸⁻³⁰.

In conclusion, essential oils from medicinal plants growing in the region can offer good control of *A. obtectus* adults. Natural insecticides are a desirable alternative to synthetic pesticides

because they have low toxicity in mammals, little environmental effect and wide public acceptance. In this aspect, essential oils of plants may play an important role in the stored-products protection and reduce the risks associated with the use of highly toxic fumigants. The essential oils tested in this study could be considered as potential alternatives for highly toxic fumigants with modification as their structures could lead to the development of new classes of insecticide compounds. However, further experiments are needed to elucidate the efficacy of these substances in large-scale applications, their fumigant properties.

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