Behavioral responses of *Losiderma serricorne* and chemical composition of essential oil isolated from *Nicotiana tabacum* leaves

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Essential oils can have an impact on behavioural responses of cigarette beetle (*Lasioderma serricorne*). In this study, we provide evidence that some chemical compounds in essential oil of tobacco (*Nicotiana tabacum*) leaves, which were subjected to hydrodistillation and extracted with n-hexane, can strongly attract *L. serricorne* significantly. Two main chemical constituents, 6,8-nonadien-2-one, 8-ethyl-5-(1-methylethyl) and 1-(2,6,6-trimethyl-1,3-cyclohexadien-1-yl) 2-buten-1-one, in the essential oils were confirmed by GC-MS analysis. This result could be helpful to find some chemical compositions from *N. tabacum* as leading compounds for development of new bionic attractant.

Keywords: Lasioderma serricorne, Nicotiana tabacum, behavioural responses, essential oil, GC-MS

1. Introduction

Tobacco, *Nicotiana tabacum*, originated in South America. It is mainly distributed in South America, South Asia and China [1]. During storage, tobacco vulnerable to tobacco beetles infestation. Tobacco beetles leaving smell or holes in the cigarette and serious impact availability and quality of tobacco [2].

The cigarette beetles, *Lasioderma serricorne*, is one of the most serious pest of stored tobacco, tobacco products, cereal grains and processed foods and cause

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significant damage to the multibillion dollar food and tobacco industries worldwide each year[3-6]. *L. serricorne* is a pervasive pest with an international distribution mainly in tropical, subtropical and warm-temperate climates [7].

In this paper, we studied the attracting activity of essential oils from *N. tabacum* L. leaves on *L. serricorne*, and tried to find some chemical compositions from *N. tabacum* as leading compounds for development of new bionic attractant.

2. Materials and Methods

2.1. Plant materials

The dried leaves of *N. tabacum* were obtained from Hongyun Honghe tobacco (Group) Co., Ltd. Honghe Cigarette Factory, Mile, Yunnan Province and then kept in a dark and cold condition until used shortly after that for the experiments.

2.2. Insects

The cigarette beetles (*L. serricorne*) were obtained from laboratory cultures maintained for the last 2 years in the dark incubators at 28±0.5°C and 70±0.5% relative humidity. Unsexed adult beetles used in all the experiments were about 10 days old.

2.3. Extraction of essential oil

The powder of *N. tabacum* Leaves was subjected to hydro distillation using a modified Clevenger-type apparatus for 4 h and extracted with *n-*hexane. Anhydrous sodium sulphate was used to remove water after extraction. The obtained essential oil was kept in sealed glass vial at 4 °C until further use.

2.4. Behavioral responses test

Behavioral responses of the cigarette beetles (*L. serricorne*) were investigated with a glass Y-tube olfactometer of 1cm internal diameter, with a 14 cm stem and 15 cm arms at a 75° inner angle. Air was pumped through an activated charcoal filter and a humidifier through Teflon tubing and divided by a glass Y-junction. The two airflows then passed through two separate flow meters, which regulated the flow rate to 500 mL·min⁻¹. The air then passed into two spherical glass vessels (i.d. 35 mm), into which the essential oil was placed. From here, the air from the vessels flowed into the arms of the olfactometer. The essential oil of *N. tabacum*. Was diluted with *n-*hexane at two different concentration levels of 2.5 μg·μL⁻¹ and 5.0 μg·μL⁻¹, respectively. Ten microlitres
of essential oil diluents was added to a filter paper (diameter 3.0 cm). The solvent was allowed to evaporate for 30 s before the filter paper put into the odor vessel of olfactometer. *N-hexane* was used as control.

The Unsexed adult beetles about 10 days old were used for the tests. The beetles were individually introduced into the olfactometer. The beetles in the olfactometer chose one side of the two arms by chemotaxis at the Y-junction of the olfactometer. The observation ended when the beetles reached the finish line (3.5 cm from Y-junction) of one of the olfactometer arms, with maximum observation duration of 5 min per beetle. The treatment and control arms were reversed after half of the beetle tests had been accomplished. Experiments were conducted in a climate-controlled room (26 ± 1°C, 70 ± 5% RH) under one 40-W fluorescent lamps, fitted with a prismatic filter, to ensure a completely even distribution of light. This illuminated the Y-tube with a light intensity of 750 lux (measured by a digital luxmeter, model 7001, Germany). Ten different beetles were used for each replication and five replications were carried out for behavioral responses test. Between trials, all glassware was washed with acetone and distilled water and then baked at 210°C over night to remove any volatiles adhering to the glass.

2.5. Identification of essential oil

Essential oils were identified by gas chromatograph-mass spectrometry (GC-MS). Gas chromatographic analysis was conducted on a gas chromatograph (Agilent 7890, America) coupled to a quadrupole mass spectrometer (Agilent 5975N, America). Separation of compounds was performed on a HP-5MS capillary column (J&W Scientific, 30m×0.25mm i.d., film thickness 0.25μm). The GC oven temperature was programmed as follows: from 40 to 250°C at a rate of 5°C min⁻¹ and held for 10 min. The injector temperature was 260°C. Helium of high-purity was used as the carrier gas at a flow rate of 1.0 mL·min⁻¹. Split injection mode was used and split ratio was 10:1.

The mass spectrometer was fitted with an EI source operated at 70 eV with a source temperature of 230°C, and mass spectra were recorded in the range of m/z 50-550 amu in the full scan acquisition mode. The quadrupole temperature was 150°C, and the interface temperature was 280°C. Essential oils were identified by comparing the obtained mass spectra of the analytes with those of authentic standards from the NIST 08 library with a resemblance percentage above the ninety percent. The chemical components relative content was calculated based on the peak area of total ion current chromatogram.
2.6. Statistical analysis

Chi-square tests were used to test the hypothesis that the numbers distribution of side-arm choices between the essential oil and control. Independent-Sample T tests were used to determine significant differences between the mean times of individual beetle making a choice between the essential oil and control. All the data were analyzed with statistical package SPSS 17.0

3. Result

3.1 Responses of L. serricorne to the essential oil

The results obtained on the olfactory response of L. serricorne between different doses of the essential oil and blank control were showed in figure 1.

Figure 1 clearly illustrated that the beetles (L. serricorne) was attracted to essential oil from tobacco leaves significantly ($p<0.01$) when a dose of the essential oil was less than 500 $\mu$g. But when a dose of the essential oil was 1000 $\mu$g the beetles (L. serricorne) did respond significantly better to blank control.

![Fig. 1. Mean ± SD percentages of cigarette beetles in an olfactometer that moved upwind toward the essential oil or blank control. An asterisk above bars indicates they are significantly different by a paired t-test at $P<0.01$.](image)

3.2 GC-MS analysis of the essential oil

Total ion current (TIC) chromatogram of mass spectrometry for essential oil was showed in figure 2.

![Fig. 2. TIC of essential oil from leaves of N. tabacum.](image)
Based on the total ion current chromatogram of mass spectrometry, there were two main chemical constituents in the essential oil. They were 6, 8-Nonadien-2-one, 8-methyl-5-(1-methylethyl) (retention time 22.763 min) and 1-(2, 6, 6-Trimethyl-1, 3-cyclohexadien-1-yl)-2-buten-1-one (retention time 23.256 min). Result of quantitative analysis by peak area normalization showed that content of 6,8-Nonadien -2-one, 8-methyl-5-(1-methylethyl) and 1-(2,6,6-Trimethyl-1, 3-cyclohexadien-1-yl)- 2-buten-1-one in essential oils was up to 31.748% and 18.009%, respectively.

4. Discussion

*L. serricorne* is major pests on a wide variety of dry and durable stored agricultural products [8]. There are many different traps baits of *L. serricorne* were reported to attract tobacco beetles, such as sex pheromone serricornin, red chilli powder, chilli volatiles, combining pheromone with plant-derived volatiles and other synthetic sex pheromone [5, 8-11]. The essential oil was extracted by Soxhlet method from *Ailanthus altissima* Swingle bark could repelled and fumigate *L. serricorne* adults [12]. Essential oils isolated from *Cymbopogon citratus* and *Eucalyptus citriodora* could repel *Tribolium castaneum* [13].

The data from this study provide evidence that essential oils from tobacco leaves can attract *L. serricorne* significantly. It is similar to *L. serricorne* attracted by chilli volatiles [8]. But there are differences between tobacco essential oils and chilli volatiles, there are come from host plant and another plant, respectively. If we combine tobacco essential oils with chilli volatiles instead of sex pheromone, it not only enables build a green tobacco industry but also dramatic save prevention costs.

There are two main chemical constituents, 6, 8-nonadien-2-one, 8-ethyl-5-(1-methylethyl) and 1-(2,6,6-Trimethyl-1,3-cyclohexadien-1-yl)-2-buten-1-one, in the tobacco essential oils were confirmed by GC-MS analysis. They may be two of the food pheromones of tobacco beetles. According to the report, there are many different pheromones consisting of essential oil components of various plants and released by insects and arachnids (pests) [14]. Therefore, strategies for pest management by using pheromones seem to be cost-effective, and environment friendly compared to the use of insecticides.

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