ANTIFEEDANT EFFECTS OF VITEX NEGundo L. LEAF EXTRACTS ON THE STORED PRODUCT PEST, TRIBOLIUM castaneum H. (COLEOPTERA: TENEBRIONIDAE)

P. HARIDASAN, M. GOKULDAS, A. P. AJAYKUMAR
Insect Physiology and Biochemistry Laboratory, Department of Zoology, University of Calicut, Kerala 673635, India

Keywords: Vitex negundo, Tribolium castaneum, Nutritional index, Antifeedancy

INTRODUCTION

Search for safer and more congenial alternatives for insect pest control using chemical insecticides has resulted in more attention getting focused on botanicals. Most of the plants contain different types of secondary metabolites that have various effects on many insect species. The active components from plants act on insect pests as toxins, repellents, antifeedants, ovicidal etc. Antifeedants prevent insects from feeding on vegetation, grain or other products leading them to starvation and subsequent death. Antifeedant activity is one of the principal bioactivities brought about by plant constituents on insects, which lend impetus to the incorporation of feeding deterrence strategy in integrated pest management (IPM) programmes, which ultimately lead insects to die of starvation. Since most of these antifeedant principles, derived from plants are non-toxic in nature, their environmental compatibility is an important advantage. Antifeedants of plant origin have the additional advantage of quick biodegradability and are friendly to non-target organisms. Hence, insect antifeedants have become the subject of considerable interest.

Many plant materials have been effectively used against stored product insects. Some studies have been carried out to investigate the antifeedant effect of pithuraj, Aphanamixis polystachya on T. castaneum [1] and certain secondary plant compounds isolated from the sub-fractions of A. polystachya seed extract and the subfraction A-142 contained strong feeding deterrents that acted against T. castaneum [2]. Similarly, antifeedant properties of the rhizome extract of Acorus calamus have been investigated against T. castaneum [3]. Four sesquiterpene alkaloids isolated from Celastrus angulatus were also showing antifeedant activities on T. castaneum [4]. Similarly, antifeedancy of alkaloid-containing fraction isolated from the leaves of Nicotiana tabacum against the larvae of T. castaneum has been reported [5].

Vitex negundo is an aromatic shrub, found throughout the greater part of India. Almost all parts of this plant are valuable in medicine. Various preparations of this plant were reported to have anti-inflammatory [6], antiallergic [7], hirudicidal [8], antiasthmatic [9] and other biological properties. Even though many biological effects of V. negundo have been identified, its antifeedant effect has not been studied on T. castaneum until the present study.

In the light of abovementioned facts, the present investigation was carried out to investigate the nutritional and feeding deterrence indices (‘no choice’ feeding bioassay) for the stored grain pest, T. castaneum adults to V. negundo leaf extracts prepared in petroleum ether and methanol.

MATERIALS AND METHODS

Materials

The experimental plant, Vitex negundo Linnaeus (B0485 CALI; Family: Verbenaceae) was used for the purpose of preparing extracts of leaves in petroleum ether (Glaxo BP 60-80°C) and methanol (Merck) and for evaluating nutritional and feeding deterrent indices for the stored product pest, Tribolium castaneum Herbst (Coleoptera: Tenebrionidae) in the laboratory. The equipment/apparatus used in the study were electronic balance (Anamed, Sartorius), automatic shaker (Toshniwal), hot air oven (Kemi), vacuum pump and Büchner funnel.

Methods

Insect culture

Adults of the rust-red flour beetle, Tribolium castaneum were collected from local granaries and reared in the insectary in...
plastic jars (13.5 x 10 cm) containing wheat flour. The culture was maintained at 26 ± 2°C and 70-80 % R H. After 2 w, larvae, pupae and adults were separated by sifting the wheat flour with a fine meshed sieve and were transferred into clean jars containing fresh food. Freshly emerged adults were used for the bioassay.

Preparation of plant extracts

Leaves of *V. negundo* (Vellonchi-Malayalam) were collected from the Calicut University campus, Kerala, India (11.1339° N, 75.8940° E). Fresh leaves were thoroughly washed in distilled water and shade dried at room temperature for a week and they were further dried for one day in a hot air oven maintained at 35-38°C. The dried leaves were then powdered using a domestic grinder and sifted through a fine meshed sieve. The leaf powder was stored in air tight glass bottles at about 4°C. The leaf powder was used for the preparation of *V. negundo* petroleum ether extract (VPE) and *V. negundo* methanol extract (VME).

Fifty grams of the powdered plant material was mixed with 200 ml of petroleum ether taken in a conical flask and the mixture was agitated on an automatic shaker for 24 h at room temperature keeping the flask tightly covered. The extract was then filtered through Whatman No. 1 filter paper by negative pressure using a Büchner funnel and a suction pump. The residue was re-extracted in another 150 ml of petroleum ether and filtered after 24 h. The filtrates were combined and allowed to dry in a hot air oven maintained at 40°C. The weight of the dried residue was determined. After ascertaining the final weight of the residue, 10% stock solution was prepared in petroleum ether. Other required dilutions were made using petroleum ether. Similarly, VME was prepared using methanol as the extracting solvent instead of petroleum ether.

Nutritional and feeding deterrence indices bioassay

Nutritional and feeding deterrence indices for *T. castaneum* adults to the VPE and VME were studied by the method of 'no choice' test using extract-incorporated diet in the chamber. It is a modified method of flour disc bioassay [10]. Testing of this nutritional effect was conducted in the 'no choice' chambers. This consisted of separate plastic vials with perforated lids (3.2 cm diameter x 4.5 cm height) containing diets incorporated with three concentrations of VPE (treated) and petroleum ether alone (control). The vials were labeled properly.

The extract-incorporated diet was prepared by mixing 150 µl each of 1.25, 2.5 and 5% concentrations of VPE added drop-wise into 150 mg of wheat flour taken in separate watch-glasses. Similarly, the same volume of petroleum ether alone was added into same quantity of wheat flour in a fourth watch glass (control). The contents of all the watch glasses were thoroughly mixed using separate glass rods and the solvent was allowed to evaporate completely at room temperature for about 3 h. The 4 'no choice' chambers were weighed separately. Then 100 mg each of the diet treated with different concentrations of VPE (1.25, 2.5 and 5%) were added into the four vials (3 treated and 1 control).

Forty healthy adults, pre-starved for 24 h before the test, were weighed as 4 groups of 10 and released into the 4 pre-weighted vial ('no choice' chambers) containing the diet. They were kept in the insectary at room temperature for one day for consumption of the diet. After 24 h, all insects were removed from the 'no choice chambers' and the live insects were weighed separately as 4 groups. Mortality of insects, if any, was recorded.

Also, each vial with lid plus flour diet was weighed to determine the decrease in weight of flour diet in control and the treated. Six replicates were used for each treatment and control. The experiment was repeated with the other extract, VME, in the same pattern as with VPE. The diet used for control was treated with methanol alone.

Nutritional indices were calculated as described by Manuwoto and Scriber 1982 [11]; Farrar et al., 1989 [12] with some modifications.

\[
RGR = \frac{(A - B)}{B} \times \text{day}
\]

\[
RGR = \text{Relative Growth Rate, where } A = \text{weight of live insects on one day (mg)/number of live insects on one day. B = initial weight of insects (mg)/initial number of insects.}
\]

\[
RCR = \frac{D}{B} \times \text{day}
\]

\[
RCR = \text{Relative Consumption Rate, where } D = \text{biomass ingested (mg)/number of live insects on one day.}
\]

\[
ECI(\%) = \frac{RGR}{RCR} \times 100
\]

\[
ECI(\%) = \text{Percentage of Efficiency of Conversion of Ingested food.}
\]

Antifeedant activity (feeding deterrence activity) was calculated as Feeding Deterrence Index (Isman et al., 1990) [13].

\[
FDI(\%) = \left[1 - \frac{(C - T)}{C}\right] \times 100
\]

\[
FDI(\%) = \text{Percent of Feeding Deterrence Index, where } C = \text{Consumption of control diet.}
\]

\[
T = \text{Consumption of treated diet.}
\]

Statistical analysis of the data

The results are expressed as means±SEMs for 10 insects per group. The effects of different concentrations of the both the extracts on relative growth rate and relative consumption rate for *T. castaneum* were subjected to ANCOVA and efficiency of conversion of ingested food and feeding deterrence indices for the insects were analysed and compared by ANOVA.

RESULTS

**Nutritional indices for *V. negundo* petroleum ether extract (VPE)**

The relative growth rate (RGR) of insects for 24 h were showing a slight reduction in treated samples compared with the control and the RGR decreased with increase in the concentration of the extract. Thus, 1.25, 2.5 and 5% concentrations afforded the RGR of 0.040, 0.033 and 0.028 mg/mg/d respectively whereas in the control the value was 0.052 mg/mg/d (table 1).

Similarly, the relative consumption rates (RCR) of insects were significantly reduced in a dose-dependent manner. The RCR of the insects were 0.161, 0.079 and 0.067 mg/mg/d for 1.25, 2.5 and 5% concentrations of petroleum ether extract of *V. negundo* respectively. In the case of the control, the RCR was 0.23 mg/mg/d (table 1).

On the other hand, the percentage of efficiency of conversion of ingested food (ECI %) of the insects increased with an increase in the concentration of VPE. Higher ECI % was shown by 2.5% (ECI = 43%) and 5% (ECI = 41%) concentrations of the extract. Values of ECI calculated for a concentration of 1.25% of VPE was almost equal to the value (25%) obtained for the control (23%) (table 1).

**Feeding deterrence index for VPE**

Significant feeding deterrent activity or antifeedant activity against *T. castaneum* adults was exhibited by VPE at different concentrations in a dose-dependent manner. The feeding deterrence indices (FDI %) increased gradually from about 34% to 69% and 74% respectively with VPE of 1.25, 2.5 and 5% concentrations (table 1).
The highest concentration of the extract (5%) was also showing a similar effect. There was only a very slight increase in the RGR (0.027 mg/mg/d) (table 2). Similarly, relative consumption rate (RCR) also significantly decreased with increasing concentrations of the extract. A concentration of 1.25% of VME brought about only a small decrease in the RGR (0.027 mg/mg/d) (table 2). Similarly, relative consumption rate (RCR) also significantly decreased with increasing concentrations of the extract. A concentration of 1.25% of VME brought about only a small decrease in the RGR (0.027 mg/mg/d) (table 2).

However, the RCR showed only a slight decrease when the concentration was further increased to 5%. The value was 0.063 mg/mg/d (table 2). In the case of the percentage of efficiency of conversion of ingested food (ECI %) of the insects, there was an increase in the case of 1.25% of VME (39%) from the control value of 28%. This value was seen to drop to 31% when the concentration of the extract was increased to 2.5%. However, this drop was overcome when the concentration was increased to 5%. The ECI value was 48% in this case (table 2).

Feeding deterrence index for VME

Like VPE, VME also exhibited significant feeding deterrent activity against T. castaneum. It was seen that here also, the feeding deterrence index (FDI %) increased in a dose-dependent manner. The FDI % were 37, 58 and 66%, respectively, for 1.25, 2.5 and 5% of the extract (table 2).

Comparison of the effects of VPE and VME on R. castaneum

The effects of VPE and VME on R. castaneum were compared (fig. 1). The data showed that the effects on RCR of the insects were similar with both the extracts (ANOVA, P=0.80). However, there were showing a dose-dependent decrease of RCR (ANOVA, P<0.05) of the insects with both the extracts. Analysis of ECI (% for VPE and VME showed (fig. 2) similar activity (ANOVA, P=0.75) and the extracts showed an increase in the ECI (%) of the insects with an increase in the concentrations (ANOVA, P<0.05).

The data of FDI (%) showed (fig. 3) similar activities (ANOVA, P=0.07) of the two extracts. On the other hand, there was a dose-dependent increase (ANOVA, P<0.001) of FDI (%) of the insects with each extract. Comparison of RGR of the insects was showed similar activity (ANOVA, P=0.49) with both the extracts.
The data represent the pooled values for the effects of VPE and VME for different concentrations used. The difference between the activities of the two extracts was not significant (ANCOVA, $P=0.80$).

**Fig. 2:** Categorised plot for variable of ECI % for *T. castaneum* for different concentrations of VPE and VME

The data represent the pooled values for the effects of VPE and VME for different concentrations used. The difference between the activities of VPE and VME was not significant (ANOVA, $P=0.75$).

**Fig. 3:** Categorized plot for variable of FDI % for *T. castaneum* for different concentrations of VPE and VME

The data represent the pooled values for the effects of VPE and VME for different concentrations used. The difference between the activities of VPE and VME was not significant (ANOVA, $P=0.07$).

**DISCUSSION**

The reduction of RGR and RCR of *T. castaneum* by both the extracts observed in this study are in agreement with similar studies of various other workers. Reduction of growth and food consumption of *T. castaneum* and *S. zeamais* was brought about by essential oils of nutmeg seeds [14] and by cinnamaldehyde from *C. aromaticum* [15]. Allyl disulphide, a volatile compound from garlic, *Allium sativum*
[16], and 1,8-cineole from A. annua [17] were found to cause significant reduction of RGR and RCR of T. castaneum. The reduction of growth and food consumption of T. castaneum observed in this study (fig. 1) may presumably be due to the antifeedant action of the extract incorporated into the diets. This study also reveals that the quantity of food material converted to body with both the extracts (P<0.05). It was seen that the difference between the activities of these two extracts on ECI was not significant (P=0.75) (fig. 2). Significantly higher ECI values over the control were obtained with all concentrations of the extracts in the present study. Moreover, there appeared to be a dose-dependency for this effect. Similar observations were reported in Crocidolomia binotalis [18] and Spodoptera litura [19]. Similarly, Joseph reported [20] higher ECi % from his studies of antifeedancy and growth inhibitory effects of neem seed kernel extract on Alanthus defoliator, Elyma nus nucius indic. Present study using V. negundo against T. castaneum show higher ECI values which presumably reflect the compensation for antifeedant effect as suggested by Fagoonee who reported [18] the neem related high ECI and ECD values in Crocidolomia binotalis.

It was also found that the extracts exerted significant feeding deterrence (antifeedancy) on this insect (fig. 3). The feeding deterrence was found to be dose-dependent and increased by all the concentrations of the two extracts were found to be significant (P<0.001). However, the activity was similar with both the extracts (P=0.07). Antifeedant effects of V. negundo have been reported earlier. Premeeea and Muraliedheran reported [21] that certain phytochemicals in the extracts of V. negundo caused a significant reduction in the levels of all the three digestive enzymes (midgut protease, invertase and amylase) and thus inhibited food digestion in the red cotton bug, Dytiscus cingulatus. Extracts of V. negundo, [18] and essential oils derived from this plant [22] have been reported to exhibit antifeedant activities against Spodoptera litura.

Several essential oil constituents contained in the petroleum ether and methanol extracts of V. negundo leaves may be responsible for the antifeedant activity against T. castaneum. GC and GC-MS analysis of the essential oils of V. negundo leaves [23] revealed that this oil contain 65 known compounds, including sabine (p-cymene, beta- phellandrene, gamma-terpinene, terpinene-4-ol, beta-caryophyllene, alpha-guaiene, spathulenol, beta-caryophyllene oxide, globulol, viridiflorol, bis [1,1-dimethyl]-methylphenol, abeta-7, 13-diene and several minor unidentified compounds. Jirovetz et al. investigated [24] the essential oil components of the leaves of V. negundo var. negundo and V. negundo var. purpureascens by GC-FID, GC-MS and olfactometry. The major monoterpenes (terpinene-4-ol, p-cymene, a-terpineol, sabine) and sesquiterpenes (beta-caryophyllene, globulol, spathulenol, beta-farnesene and bis [1,1-dimethyl] methyl phenol) from the leaves of the two varieties of V. negundo. Hebbalkar et al. reported [25] that oils from V. negundo leaves when analysed by column chromatography and IDCC technique revealed that it contained several compounds such as a-terpine, p-terpine, p-cymene, a mixture of sesquiterpene hydrocarbons, 4-terpineol, monoterpenes, sesquiterpenes and a mixture of monoterpenes and sesquiterpene alcohols. The majority of components present in the fractions exhibited repellent activity against A. aegypti. These essential oil constituents present in the leaf extracts presumably are responsible for antifeedant activity against T. castaneum. Several volatile constituents of V. negundo leaves were analysed by GC-MS [26]. The main components detected were viridiflorol, b-caryophyllene, sabine, 4-terpineol, g-terpinene, caryophyllene oxide, 1-octen-3-ol and globulol. These components in the leaves may also act as antifeedants against T. castaneum.

The mechanism of perception of inhibitory chemicals in insects has been reviewed by Chapman [27] and Schoonhoven [28]. On the basis of behavioural and electrophysiological studies, parts of the body on which the relevant receptors are located and the mechanisms by which insects differentiate inhibitory and stimulatory materials have been established [27]. Antennae in insects are generally considered to be the main organ for the perception of smell in insects [29] and they would be expected to be involved in responses to olfactory repellents. The experiments of Haskell and Mordue [30] suggested that sensilla on the tip of the labrum have a primary role in the detection of inhibitory chemicals in S. gregaria. These mechanisms of perception of inhibitory chemicals in feeding may also occur in T. castaneum adults in response to V. negundo leaf extract incorporated wheat flour diets. Various chemical components extracted from V. negundo in petroleum ether and methanol presumably inhibit the feeding of the insect and perception of these chemicals is effected by the receptors like sensilla on the antennae, labrum, maxillary and labial palps of T. castaneum.

Electrophysiological studies have indicated that inhibitory phytochemicals may be involved in the antifeedancy in insects. These chemicals may either directly inhibit the input from the phagostimulant receptor, or the signals may be interpreted as inhibitory at the central nervous system [31, 32 and 33]. Similar electrophysiological mechanism of perception of inhibitory chemicals may also be active in T. castaneum with regard to the response to the extract incorporated diets. Active phytochemicals in the extracts presumably suppress the activity of the receptor cells of these insects or input signal from the receptor neuron may be perceived by the central nervous system in which it is interpreted as inhibitory and may cause an antifeedant effect.

CONCLUSION

The present study thus reveals that both petroleum ether and methanol extracts of V. negundo show significant reduction of consumption and utilisation of food and thereby bring about antifeedancy to T. castaneum. Since antifeedant action offers a valuable weapon against this pest, further study on the effect of these extracts on other stored products insects and phytophagous insects should be carried out. This also suggested that there may be different chemical constituents in both the extracts responsible for antifeedancy in a dose-dependent manner. So, further analytical studies should be carried out to determine the exact chemical structure of the bioactive components in the extracts of V. negundo to understand how each of the constituent influences the physiology and behaviour of the insects. Moreover, it might be worthwhile if the insect antifeedant components of V. negundo are produced commercially so that their potential in controlling stored product pests can be fully exploited.

ACKNOWLEDGEMENT

P. Haridasan is grateful to the University Grants Commission (UGC), New Delhi, for the award of a teacher fellowship under FIP.

CONFLICT OF INTEREST

Declared none

REFERENCES


How to cite this article