Toxicity and Antifeedant Activity of Selected Plant Extracts Against Larval Obliquebanded Leafroller, *Choristoneura rosaceana* (Harris)

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**Abstract:** Several plant extracts were evaluated against obliquebanded leafroller larvae to determine potential toxicity and antifeedant effects. Two of the plant extracts exhibited contact toxicity; the LC₅₀ value of *Humulus lupulus* extract against 3rd instar larvae was 24.5 μg / insect. In a no choice context, larval survivorship was reduced 23 - 57 % after feeding on a meridic pinto bean diet treated with the extracts. Furthermore, average female and male pupal weight was reduced for leafrollers that fed on diet treated with each of the individually-tested extracts at a 4 % concentration (w/w). Incorporation of *Bifora radians* and *H. lupulus* into diet deterred larval feeding at a 1 % concentration by weight. *H. lupulus* and *Arctium lappa* extracts were deleterious to larvae in each of the bioassay methods conducted, exhibiting toxic, developmental, and antifeedant effects. These two extracts may be promising candidates for further development as botanical insecticides that could potentially substitute broad-spectrum synthetic neurotoxins for leafroller control.

**INTRODUCTION**

The Obliquebanded leafroller, *Choristoneura rosaceana* (Harris), is a tortricid moth with a wide host range including woody plants in the Rosaceae as well as *Ulmus, Populus, Quercus, Betula*, and *Tilia* [1]. Depending on the geographic region and climate, there are one or two generations of adults per year with an overwintering second or third-instar larval generation [2]. The polyphagous larvae are primarily foliage feeders, but are also known to feed on and damage apple fruit [3].

Obliquebanded leafrollers are major pests in apple growing regions throughout the U.S.A. and Canada [4]. Due to azinphos-methyl resistance, population densities of this pest have increased in commercial apple growing regions in the Eastern United States [5]. Resistance to organophosphates is also becoming prevalent in Canada [6, 7]. Physiological resistance to other broad-spectrum insecticides and newer chemistries such as insect growth regulators has been also identified for several compounds including chlorpyrifos, esfenvalerate, and tebufenozide [4, 8].

Botanical insecticides such as azadirachtin are often effective alternatives to organophosphates or other neurotoxins for pest control due to multiple modes of action. These include toxicity, antifeedant and anti-oviposition effects [9,10]. Natural products containing secondary plant compounds such as terpenes, steroids, alkaloids, phenolics and cardiac glycosides [11] affect insect behavior and are toxic in some cases [12-18]. Identification of plant extracts that exhibit the above-described deleterious effects on pest insect physiology and behavior represents a potential alternative strategy for development of biorational controls that could replace synthetic neurotoxins. Our intent was to identify potential botanical insecticides that may prove to be effective alternatives for controlling obliquebanded leafroller.

In the current investigation, we tested the effects of several plant extracts, known to produce secondary compounds such as monoterpenes, sesquieterpenes and triterpenes [19-22] on obliquebanded leafroller larvae. These extracts were also chosen based on their previously-documented insecticidal activity [17,18,23]. The specific objectives of this study were to determine whether the selected plant extracts: 1) exhibit contact and/or ingestion toxicity to obliquebanded leafroller larvae, 2) affect pupal weight following ingestion by larvae, and/or 3) deter larval feeding.

**MATERIALS AND METHODS**

**Insect Culture**

Obliquebanded leafrollers were drawn from a six-year-old laboratory colony originally collected from unsprayed apple orchards in Southwestern Michigan, U.S.A. Moths were collected from the field as 1st and 2nd generation pupae and reared continuously without diapause. Insects were reared at 24°C and 60 % RH on a meridic pinto bean-based diet [24] under a 16:8 (L:D) photoperiod.

**Plant Materials**

The plant materials used in this study are given in Table 1. The detailed experimental protocol for preparing plant extracts was described by Gökçe et al. [25]. All plant materials were collected during spring and summer seasons of 2002-2005. Dried and ground plant materials were treated with methanol for 24 h. Thereafter, the suspensions were sieved through cheese cloth and excess methanol was re-
Additional toxicity studies were conducted using of plant extract / insect. Insects were treated by applying 1 of Whatman # 1 filter paper in a 90 mm Petri dish. The dosages (see Results). For each replicate, 10 larvae were placed on obliquebanded leafroller larvae in the initial screening assay. Toxicity of the plant extracts / insect. This experiment tested the hypothesis that topically-applied plant extract solutions exhibit contact toxicity to obliquebanded leafroller larvae. Toxicity of the plant extracts was tested using 3rd instar larvae. For each replicate, 10 larvae were exposed to each plant extract. Each larva was treated topically with 20 μg of plant extract in 1 ml of acetone using a 50 μl Hamilton syringe. Control insects (N = 10 / replicate) were treated with 1 ml of acetone. Following application of treatments, insects were incubated as described above and mortalities were assessed after 48 h. Each treatment was replicated three times per trial and three trials (blocks) were completed on separate days for a total of 9 Petri-dishes of 10 larvae per dose.

Contact Toxicity of Plant Extracts

This experiment tested the hypothesis that topically-applied plant extract solutions exhibit contact toxicity to obliquebanded leafroller larvae. Toxicity of the plant extracts was tested using 3rd instar larvae. For each replicate, 10 larvae were treated with a Whatman No. 1 filter paper disc in a 90 mm disposable Petri dish. Three replicate Petri-dishes of 10 larvae were treated with each plant extract. Each larva was treated topically with 20 μg of plant extract in 1 ml of acetone using a 50 μl Hamilton syringe. In the control treatment, larvae were treated with 1 ml of acetone. After application of treatments, the larvae were allowed to dry for 10 min at 25 ± 2 ºC and were subsequently transferred individually into 240 ml soufflé cups loaded with 3.0 g of the meridic pinto bean diet described above. Following treatment application, larvae were maintained at 25 ± 2 ºC and on a 16:8 (L:D) photocyte for 48 h, after which mortality was assessed. The experiment was repeated on three different days, which were regarded as blocks. Plant extract stock suspensions described above were prepared immediately prior to each block of assays.

Toxicity of H. lupulus Extract

Additional toxicity studies were conducted using H. lupulus extract based on its substantial toxic effects on obliquebanded leafroller larvae in the initial screening assay (see Results). For each replicate, 10 larvae were placed on Whatman # 1 filter paper in a 90 mm Petri dish. The dosages of H. lupulus compared were 0.3, 1.5, 3.0, 15.0 and 30.0 μg of plant extract / insect. Insects were treated by applying 1 μl of HPLC grade acetone solution of plant extract to the thorax using a Hamilton syringe. Control insects (N = 10 / replicate) were treated with 1 μl of HPLC grade acetone. Following application of treatments, insects were incubated as described above and mortalities were assessed after 48 h. Each treatment was replicated three times per trial and three trials (blocks) were completed on separate days for a total of 9 Petri-dishes of 10 larvae per dose.

Ingestion Toxicity and Developmental Effects of Plant Extracts

This experiment tested the hypothesis that larval feeding on diet treated with plant extracts in a no choice context affects larval survival and pupal weight. Ten, 20.0 or 40.0 mg of each plant extract in 100 μl of HPLC grade acetone was incorporated per g of freshly prepared pinto bean diet (described above) to give 1, 2 or 4 % (w/w) concentrations. Each plant extract diet treatment was formulated separately. For the control treatment, 100 μl of HPLC grade acetone was added to the diet. Three g of each diet treatment was transferred into 240 ml soufflé cups. To avoid cannibalism, one freshly-moulted 3rd instar larva was transferred into each cup. The larvae were incubated at 25 ± 2 ºC and on a 16:8 (L:D) photocyte for 14 d. Larvae were checked daily until all had pupated. Pupation was considered complete following melanization. The pupae were sexed using a dissection microscope and pupal weight was recorded using a bench balance (Sartorious, CP124 S, Elk Grove, IL, U.S.A.). The experiment was repeated on three different days (blocks) and 30 larvae were exposed to each plant extract or control treatment per block.

Feeding Deterrence Assay

This experiment tested the hypothesis that larval feeding is deterred by plant extracts incorporated into the pinto bean diet in a choice context. Ten or 40.0 mg of each plant extract in 100 μl of HPLC grade acetone was incorporated per g of freshly prepared pinto bean diet to give 1 or 4 % (w/w) concentrations of each extract in diet. For the control treatment, 100 μl of HPLC grade acetone was added to the diet. For choice tests, larvae were presented with 1.0000 ± 0.0008 g of diet treated with a plant extract versus an equal amount of solvent-treated (control) diet in 50 mm round Petri dishes. Each plant extract and concentration treatment combination was tested individually versus the control. Larvae were inserted into Petri dishes individually on a piece of Whatman # 1 filter paper (1 x 1 cm) and placed centrally on portions of both treated and control diet pieces. Larvae were incubated for 72 h at 25 ± 2 ºC and on a 16:8 (L:D) photocyte. After incubation, the site of larval feeding was determined and scored as feeding on either extract-treated or control portions of the diet. Thereafter, larvae and visible frass were removed from diet with soft forceps and both extract-treated and control portions of the diet were weighed as described above. Remaining weights of treated and control portions of the diet were used to calculate an antifeedant index (AFI) modified from Isman et al. [26]. AFI = (C – T) / (C + T), where C and T are the weights of control and plant extract-treated diet consumed. Forty larvae were exposed to each plant extract or control treatment combination at both concentrations tested. Weight loss of diet due to water evaporation was quantified by establishing two positive control treatments of 1.0000 ±
0.0008 g of diet treated with a plant extract or solvent \((N = 20)\). These were incubated and weighed in a manner identical to that described above.

**Data Analyses**

For the initial screening bioassay, data were corrected for mortality in the controls using Abbott’s formula [27] and then normalized using an arcsine transformation [28]. Transformed data were submitted to a randomized complete block analysis of variance (ANOVA) \((P < 0.05)\) and differences between treatments were compared using Tukey’s test \((P < 0.05)\). Obliquebanded leafroller larval mortality data obtained from the dose-mortality bioassay utilizing \(H. lupulus\) were also corrected for control mortality with Abbott’s formula. The corrected mortality data were analyzed using POLO-PC [29] to estimate lethal concentration 10, 50, and 90 values \((LC_{10, 50, 90})\) and the regression line slope. For the ingestion toxicity experiment, data were corrected with Abbott’s formula and then normalized using an arcsine transformation [28]. Transformed data were submitted to a randomized complete block ANOVA \((P < 0.05)\) and differences between treatments were tested using Tukey’s test \((P < 0.05)\). Pupal weights of male and female leafrollers were analyzed separately and submitted to a randomized complete block ANOVA \((P < 0.05)\) followed by Tukey’s test \((P < 0.05)\). For the feeding choice assay, the numbers of larvae feeding on extract-treated versus control portions of the diet were compared by \(\chi^2\) tests. All statistical analyses were carried out using MINITAB Release 14 [30].

**RESULTS**

**Contact Toxicity of Plant Extracts**

Topically-applied \(H. lupulus\) induced the highest mortality of obliquebanded leafroller larvae (Fig. 1). Mortality of 3\textsuperscript{rd} instar obliquebanded leafrollers was significantly \((F = 9.8; \text{d.f.} = 5, 12; P < 0.05)\) greater for larvae treated with \(H. lupulus\) and \(A. lappa\) plant extracts compared with the solvent control (Fig. 1). There was no significant effect \((P > 0.05)\) of topical treatment with \(B. radians\), \(V. songaricum\), and \(X. strumarium\) extract solutions compared with the solvent control (Fig. 1).

**Toxicity of \(H. lupulus\) Extract**

The slope and intercepts of the dosage-mortality relationship for obliquebanded leafroller larvae treated with \(H. lupu-

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**Table 2. Average Survivorship ± S.E. of Obliquebanded Leafroller Larvae Reared on Pinto Bean Diet Containing Various Concentrations of the Plant Extracts**

<table>
<thead>
<tr>
<th>Plant species</th>
<th>Larval Survival (%) at Various Concentrations of Plant Extract in Diet (w/w)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1 %</td>
</tr>
<tr>
<td>Arctium lappa</td>
<td>98.85 ± 1.96 a</td>
</tr>
<tr>
<td>Bifora radians</td>
<td>98.85 ± 1.96 a</td>
</tr>
<tr>
<td>Humulus lupulus</td>
<td>93.10 ± 3.19 a</td>
</tr>
<tr>
<td>Verbascum songaricum</td>
<td>95.46 ± 1.96 a</td>
</tr>
<tr>
<td>Xanthium strumarium</td>
<td>100 ± 0.00 a</td>
</tr>
<tr>
<td>Solvent control</td>
<td>100 ± 0.00 a</td>
</tr>
</tbody>
</table>

*aMeans in a column followed by the same letter are not significantly different (ANOVA, \(P < 0.05\)) followed by Tukey’s mean separation \((P < 0.05)\).*
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Table 3. Average Pupal Weight of Obliquebanded Leafrollers Reared on Pinto Bean Diet Containing Various Concentrations of Plant Extracts

<table>
<thead>
<tr>
<th>Plant Species</th>
<th>Average Pupal Weight ± SE (mg) at Various Concentrations of Plant Extract in Diet (w/w)</th>
<th>Females</th>
<th>Males</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1 %</td>
<td>2 %</td>
<td>4 %</td>
</tr>
<tr>
<td>Arctium lappa</td>
<td>85.17 ± 2.65 a</td>
<td>80.37 ± 4.35 ab</td>
<td>61.03 ± 5.95 b</td>
</tr>
<tr>
<td>Bifora radians</td>
<td>83.83 ± 6.32 a</td>
<td>67.50 ± 6.88 ab</td>
<td>47.40 ± 8.46 cd</td>
</tr>
<tr>
<td>Humulus lupulus</td>
<td>79.20 ± 1.85 a</td>
<td>62.27 ± 3.50 b</td>
<td>41.80 ± 6.68 d</td>
</tr>
<tr>
<td>Verbascum songaricum</td>
<td>79.13 ± 1.93 a</td>
<td>56.40 ± 2.40 b</td>
<td>45.40 ± 5.19 cd</td>
</tr>
<tr>
<td>Xanthium strumarium</td>
<td>98.87 ± 7.06 a</td>
<td>63.97 ± 3.15 b</td>
<td>56.40 ± 6.40 bc</td>
</tr>
<tr>
<td>Solvent control</td>
<td>97.53 ± 2.96 a</td>
<td>96.97 ± 6.85 a</td>
<td>98.85 ± 7.97 a</td>
</tr>
</tbody>
</table>

*Means in a column followed by the same letter are not significantly different (ANOVA, P < 0.05) followed by Tukey’s mean separation (P < 0.05).

lusk extract were 0.93 ± 0.15 and 0.02 ± 0.02, respectively. The LC10, 50, and 90 values (Fiducial Limits %) were 1.03 (0.16 - 2.42), 24.52 (14.08 - 60.32), and 581.53 (164.20 to 11,724.49) µg / insect, respectively.

**Ingestion Toxicity and Developmental Effects of Plant Extracts**

There was no significant (F = 1.3; d.f. = 5, 12; P = 0.32) effect of any of the plant extracts on survivorship of feeding larvae at the 1 % concentration (Table 2). Significantly fewer (F’s = 7.1 and 4.8, d.f. = 5, 12, P < 0.05) obliquebanded leafroller larvae survived to pupation at the 2 and 4 % concentrations for every plant extract treatment except A. lappa (Table 2). Larval mortality appeared to increase in a dosage dependent manner (Table 2).

Feeding by obliquebanded leafroller larvae on the diet containing 1 % plant extracts (by weight) did not significantly affect pupal weight for both sexes (F’s = 0.7 and 3.1; d.f. = 5, 12; P > 0.05 for males and females, respectively) (Table 3). Increasing the concentration of plant extract in diet caused a reduction in pupal weight. Female pupal weight was significantly lower (F = 5.2; d.f. = 5, 12; P < 0.05) for larvae feeding on the 2 % concentration of H. lupulus, V. songaricum, and X. strumarium, while male pupal weight was significantly lower (F = 4.7; d.f. = 5, 12; P < 0.05) only for V. songaricum at that concentration (Table 3). At the highest concentration tested (4 % by weight), male and female pupal weight was significantly reduced (F = 141.7 and 75.8, respectively, d.f. = 5, 12; P < 0.05) for each plant extract tested (Table 3). At the 4 % concentration, B. radians, H. lupulus, V. songaricum, and X. strumarium had the greatest effect on male and female pupal weight.

**Feeding Deterrence Assay**

At the 1 % concentration, significantly fewer (P < 0.05) larvae were found feeding on B. radians compared with the solvent control; there was no significant (P > 0.05) effect for the other plant extracts tested (Table 4). However, significantly fewer (P < 0.05) larvae were found feeding on A. lappa, B. radians, and H. lupulus–treated diet compared with the solvent control at the 4 % concentration (Table 4).

At the 1 % concentration, B. radians and H. lupulus showed evidence of feeding inhibition on the pinto bean diet with AFI’s of 0.56 and 0.27, respectively (Fig. 2A). There was a high level of feeding inhibition for B. radians, H. lupulus and A. lappa—treated diet at the 4 % concentration with AFI’s ranging from 0.69 to 0.85 (Fig. 2B). There was no apparent antifeedant effect of V. songaricum or X. strumarium at either concentration tested (AFI’s ≈ 0) (Fig. 2A, B).

Table 4. Deterrence of 3rd Instar Obliquebanded Leafroller Larval Feeding by Plant Extracts Incorporated into Pinto Bean Diet

<table>
<thead>
<tr>
<th>Plant Species</th>
<th>Proportion of Larvae Feeding on Extract-Treated Versus Control Diet</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Treatment</td>
</tr>
<tr>
<td>Arctium lappa</td>
<td>0.45 a</td>
</tr>
<tr>
<td>Bifora radians</td>
<td>0.28 a</td>
</tr>
<tr>
<td>Humulus lupulus</td>
<td>0.35 a</td>
</tr>
<tr>
<td>Verbascum songaricum</td>
<td>0.48 a</td>
</tr>
<tr>
<td>Xanthium strumarium</td>
<td>0.55 a</td>
</tr>
</tbody>
</table>

*Values in a row followed by the same letter are not significantly different (P < 0.05).
DISCUSSION

Discovery of novel toxins and/or antifeedants from plant extracts has been recently emphasized as a potential method for the development of “ecologically safe pesticides” [31]. There is ample precedent for screening crude plant extracts for biological activity as botanical insecticides against Lepidoptera. For example, crude methanolic extracts of *Trichilia americana* and ethanolic extracts of *Annona squamosa* seeds reduced pupal weight and exhibited toxic activity against the Asian armyworm, *Spodoptera litura* (Fabr.) [31,32]. In addition, ethanolic seed extracts of *A. squamosa* and *A. muricata* reduced larval growth of *S. litura* and the cabbage looper, *Trichoplusia ni* (Hbn.) [33], while acetonic seed extracts of *A. squamosa* showed insecticidal activity against the cabbage head caterpillar, *Crocidolomia binotalis* Zeller [34]. Furthermore, crude extracts of *Melia volkensii* have been shown to inhibit larval growth of *T. ni* and the armyworm, *Pseudadetia unipunctata* (Haworth) [35].

Fig. (2). Antifeedant effects of plant extracts incorporated into pinto bean diet at 1 % (A) and 4 % (B) concentrations (w/w) on 3rd instar obliquebanded leafroller larvae. AFI = (C – T) / (C + T), where C and T are the weights of control and plant extract-treated diet consumed.
Our results demonstrated that two of the plant extracts evaluated (A. lappa and H. lupulus) exhibited contact toxicity to obliquebanded leafroller larvae while four of the extracts decreased larval survivorship and final pupal weight following ingestion (Table 2). H. lupulus yielded the highest contact toxicity with an LC₅₀ of 24.5 µg / larva. These results are in agreement with previous studies showing pronounced contact toxicity of H. lupulus to various development stages of Colorado potato beetle, Leptinotarsa decemlineata (Say) [17,18]. The contact and ingestion toxicities induced by the tested plant extracts, particularly H. lupulus, suggest the potential for their future use against obliquebanded leafroller as botanical insecticides. Further testing of biological activity on other related pest species such as redbanded leafroller, Argyrotaenia velutinana (Walker), and Pandemis leafroller, Pandemis pyrusana Kearfott, is warranted.

In addition to toxicity via contact or ingestion, plant extracts and allelochemicals have been screened for activity as insect antifeedants [36-38]. In some instances, the bioactivity of crude plant extracts on insects is comprised of both toxic and antifeedant effects [32,35]. Azadirachtin, for example, derived from the neem tree (Azadirachta indica), is both a toxicant and antifeedant and has been one of the most widely tested and successfully implemented botanical insecticides over the past two decades [39-41]. In the current study, H. lupulus and A. lappa exhibited antifeedant activity on obliquebanded leafroller larvae in addition to contact toxicity, while B. radians was an antifeedant and exhibited toxic effects when ingested. H. lupulus, A. lappa, and B. radians are also known to deter feeding of larval L. decemlineata [23]. Our results suggest that potential future application of these extracts or their active components for leafroller control may exploit more than one mode of action. However, future experiments should focus on determining whether prolonged exposure of obliquebanded leafroller larvae to these plant extracts decreases antifeedant effects over time due to habituation of response [42]. H. lupulus contains alpha and beta acids, prenyllavonoids, and proanthocyanidins [43,44]. The beta acid derivative of H. lupulus repels both chewing and sucking insect pests of plants [45] and the two-spotted spider mite, Tetranychus urticae, insect pests of plants [45] and the two-spotted spider mite, H. lupulus due to habituation of response [42]. Further testing of biological activity on other related pest species such as redbanded leafroller, Argyrotaenia velutinana (Walker), and Pandemis leafroller, Pandemis pyrusana Kearfott, is warranted.

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