Identification and field evaluation of a new blend of the sex pheromone of Hypsipyla grandella

Maria Carolina Blassioli-Moraes(1), Miguel Borges(1), Raul Alberto Laumann(1), Rafael Borges(2), Amanda Rodrigues Viana(1), Marcello Jose Thomazini(3) Cleonor Cavalcante Alves Silva(1), Márcio Wandre Moraes de Oliveira(1) and Mari Ines Carissimi Boff(2)

(1)Embrapa Recursos Genéticos e Biotecnologia, Parque Estação Biológica, W5 Norte (Final), Caixa Postal 02372, CEP 70770-917 Brasília, DF, Brazil. E-mail: carolina.blassioli@embrapa.br, miguel.borges@embrapa.br, raul.laumann@embrapa.br, amanda_2111@hotmail.com, rborges1977@hotmail.com, cleonor.silva@embrapa.br, marcio.wandre@embrapa.br (2)Universidade do Estado de Santa Catarina, Centro de Ciências Agroveterinárias, Departamento de Agronomia, Avenida Luiz de Camões, n.2 090, Conta Dinheiro, CEP 88520-000 Lages, SC, Brazil. E-mail: mari.boff@udesc.br (3)Embrapa Florestas, Estrada da Ribeira, Km 111, Guaraituba, Caixa Postal 319, CEP 83411-000 Colombo, PR, Brazil. E-mail: marcilio.thomazini@embrapa.br

Abstract – The objective of this work was to identify and carry out a field testing of the sex pheromone composition of the mahogany shoot borer, Hypsipyla grandella, from a population in Southern Brazil. Abdominal glands of H. grandella females were excised and extracted using n-hexane; the gland extracts were analysed by gas chromatography (GC-FID and GC-MS). Wind-tunnel and electrophysiology experiments were conducted to evaluate the role of gland compounds in the mating behavior of H. grandella males. In the field, pheromone traps containing the identified pheromone were tested. In addition to the two previously identified compounds – (9Z,12E)-tetradecadien-1-ol and (9Z,12E)-tetradecadienyl acetate –, in insects from Central America populations, two additional compounds were identified in the blend: (9Z)-tetradecen-1-ol and (9Z)-tetradecenyl acetate. Gas chromatography-electroantennographic analyses showed that these four components elicited antennal responses from conspecific males. Wind tunnel bioassays using different amounts of the components Z9-14:OH, Z9,E12-14:OH, Z9-14:OAc, and Z9,E12-14:OAc, elicited male responses similar to the response to conspecific calling females. When the binary and quaternary mixtures were tested in field conditions, males were attracted to the traps containing the quaternary mixture. The new pheromone blend identified in H. grandella males has a great potential to be exploited as a monitoring tool or control measure in the field.

Index terms: electroantennography, gland extract, mahogany shoot borer, pheromone redescription, semiochemical, wind tunnel bioassays.

Identificação e avaliação em campo de uma nova mistura do feromônio sexual de Hypsipyla grandella

Resumo – O objetivo deste trabalho foi identificar e testar em campo a composição do feromônio sexual da broca-do-ponteiro do mogno, Hypsipyla grandella, de uma população do Sul do Brasil. As glândulas abdominais de fêmeas de H. grandella foram excisadas e extraídas por meio de n-hexano; os extratos de glândula foram analisados por cromatografia gasosa (GC-FID e GC-MS). Experimentos de túnel de vento e eletrofisiologia foram conduzidos para avaliar o papel dos compostos das glândulas no comportamento de busca de acasalamento pelos machos de H. grandella. Armadilhas com o feromônio identificado foram testadas em campo. Além dos compostos previamente identificados – (9Z,12E)-tetradecadien-1-ol e acetato de (9Z,12E)-tetradecadienila –, em populações de insetos da América Central, dois novos compostos foram identificados: (9Z)-tetradecon-1-ol e acetato de (9Z)-tetradecenila. A análise electroantennográfica mostrou que estes quatro compostos provocaram uma resposta da antena dos machos coespecíficos. Os bioensaios em túnel de vento – com misturas binária, ternária e quaternária, com uso de diferentes quantidades dos compostos – Z9-14:OH, Z9,E12-14:OH, Z9-14:OAc e Z9,E12-14:OAc –, mostraram uma resposta do macho similar à resposta de chamamento das fêmeas coespecíficas. Quando as misturas binária e quaternária foram testadas em condições de campo, os machos foram atraídos para as armadilhas com a mistura quaternária. A nova mistura feromonal identificada em machos de H. grandella tem grande potencial para ser usado no monitoramento e controle desta praga no campo.

Termos para indexação: electroantennografia, extrato de glândula, broca-do-ponteiro do mogno, redescrição de feromônio, semioquímicos, bioensaios em túnel de vento.
Introduction

In several parts of the world, the establishment and cultivation of high-value timber species of the Meliaceae family like *Swietenia*, *Cedrela*, *Khaya*, and *Toona* species have failed due to the attack of the shoot borer moths, *Hypsipyla* spp. (Lepidoptera: Pyralidae), mainly *H. grandella* Zeller and *H. robusta* Moore (FAO, 2009; Plath et al., 2011). The larvae bore the terminal and lateral shoots, breaking the apical dominance, and producing stem ramifications, which can reduce the timber commercial value.

From the early 1970’s through the 1990’s, it is possible to find a number of studies with different approaches which aimed at controlling *Hypsipyla* spp., such as: the development of resistant cultivars; the use of mixed-species plantations to enhance the diversity of the plants, which can have negative effects on herbivorous by the reduction of the capacity to find host plants, and favors natural enemies, contributing to reduce pest population level; the biological control; the chemical control; the use of semiochemicals from meliaceous spp. that have a negative effect on *Hypsipyla* spp.; and the use of sex pheromones to control these pests (Assiri-Bosson & Gallois, 1982; Borek et al., 1991; Goulet et al., 2005; Lago et al., 2006; Plath et al., 2011). However, despite all the efforts to manage these insects, they are still a limiting factor to the success of Meliaceae plantations.

From all these mentioned studies, there is a consensus that the control of *Hypsipyla* will be successful by the use of an integrated approach, or a strategy that involves the use of resistant cultivars, rational silvicultural practices, and control methods for the adult insect (Lunz et al., 2009; Plath et al., 2011). In general, Lepidoptera pests are efficiently monitored and, in some cases, they can be controlled using sex pheromone traps (Witzgall et al., 2010). Mainly for *H. grandella*, the use of sex pheromones to control adults might be the unique viable alternative for directly controlling this pest, since the larvae penetrate the stem or leaf midrib as soon as they hatch, which hinders the control. Some previous studies tried to identify the sex pheromone of the two main pests of the meliaceous species, *Hypsipyla robusta* and *H. grandella* (Assiri-Bosson & Gallois 1982; Borek et al., 1991); however the identified blends in these works were not efficient to capture these moths under field conditions (Borek et al., 1991).

The objective of this work was to identify and carry out a field testing of the sex pheromone composition of the mahogany shoot borer, *Hypsipyla grandella*, from a population in Southern Brazil.

Materials and Methods

*Hypsipyla grandella* pupae were obtained from a laboratory colony maintained at Embrapa Floresta, in Colombo, PR, Brazil (25º17’S, 49º13’W). The laboratory experiments were conducted during 2011 and 2012. Pupae were sexed and placed inside 3 L plastic containers. The adults fed a sugar solution with 1,000 mL water, 50 g honey, 50 g sugar, 1 mg Nipagin, and 1 mg ascorbic acid, according to the methodology by Schmidt et al. (2001). Males and females were kept separately. For the bioassays, insects within the period of 24 to 72 hours after last molting were used. All insects were reared in acclimatized chambers with a reversed light: dark (L:D) cycle, and a 12:12 L:D photoperiod at 25±1°C, and 65±2% RH.

Sex pheromone glands were extracted from one to three-day-old virgin calling females. The gland was forced to extrusion through a gentle pressure at the tip of the abdomen; by using a small spring scissor (model 15003-08, FST- Canada), the glands were excised, and placed into a 0.5 mL conical vial containing 100 μL of hexane, and two to seven glands in each extract (n=6). Gland contents were extracted for 20 min; the extract was filtered using glass wool to remove solid residues and, then, it was concentrated to 50 μL under a pure N2 flow. The extracts were stored at -20°C until use.

The gland extracts were analysed by a gas chromatograph GC-FID Agilent 7890A, DB-5MS column, 60x0.32 mm ID, 0.25 μm film (Supelco, Bellefonte, PA, USA), with the oven temperature maintained at 50°C for 2 min, then increased at 5°C min⁻¹ to 180°C for 0.1 min, followed by a gradual increase of 10°C min⁻¹ to 250°C for 20 min. The column effluent was analysed with a flame ionization detector (FID) at 270°C. One microliter of each sample was injected in splitless mode with helium as carrier gas. The samples were also analysed using a DB-WAX column, and they were also subjected to the same temperature programs and flow conditions, in order to calculate the retention index (RI) of each compound. The data were collected with EZChrom Elite software and handled using Excel (Microsoft Office 2007, Microsoft Corporation, USA).
For the compound identification, selected extracts were analysed using an Agilent 5975C quadruple mass spectrometer, equipped with a DB-5MS column of 30x0.25 mm ID, 0.25 µm film (Supelco, Bellefonte, PA, USA), and a splitless injector, with helium as carrier gas. Ionization was achieved by electron impact (70 eV, source temperature 200°C), and the data were collected with ChemStation software. Identiﬁcations were made by comparison of spectra with library databases (NIST, 2008), or with published spectra, and using retention indices (El-Sayed, 2014; NIST..., 2016), and conﬁrmed by GC co-injection using authentic standards. The quantitation of the sex pheromone components was conducted using the internal standard (IS) method, with n-tetradecane as IS.

Gas chromatography-electroantennographic detection (GC-EAD) was used to determine the compounds within mixtures that were detected by the male antennae. Analyses were conducted following the same procedures described by Blassioli-Moraes et al. (2016). The GC was equipped with a DB-5 column of 30x0.25 mm ID, 0.25 µm film (J&W Scientific, Folsom, CA, USA), and a splitless injector with helium as the carrier gas (1 mL min⁻¹). The oven temperature was programmed to 50°C (2 min), then rised to 250°C at 15°C min⁻¹, and kept at this temperature for 10 min. Preparations were done in a continuous humidified air ﬂow (1 L min⁻¹) with a Stimulus Controller CS-55 detector (Syntech Inc., Hilversum, The Netherlands).

Antennae of H. grandella males were tested using conspeciﬁc female gland extracts (n = 5). Antennae of H. grandella males were also tested with synthetic solutions containing the four identiﬁed compounds – (9Z,12E)-tetracadien-1-ol (Z9,E12-14:OH), (9Z,12E)-tetracadienyl acetate (Z9,E12-14:OAc), (9Z)-tetracdecen-1-ol (Z9-14:OH), and (9Z)-tetracdecenyl acetate (Z9-14:OAc). A single antennal preparation was used for only one chromatography analysis.

The n-hexane compounds for HPLC (≥ 97% and redistilled) and n-pentane for HPLC (≥ 99%), and n-tetracosane for GC (≥99%) were purchased from Sigma Aldrich (Steinheim, Germany). Z9,E12-14:OAc (97%) and Z9-14:OH (99%) were purchased from Bedoukian Research Inc. (Danbury, CO, USA). Z9-14:OH and Z9,E12-14:OH were obtained by alkaline hydrolysis; their respective acetates were dissolved in 4 mL ethanol, and 2 mL of 10% sodium hydroxide solution was added. The reaction was kept under reflux for 1 hour. The solution was then neutralized with hydrochloric acid, the organic phase was extracted using dichloromethane, and dried using magnesium sulphate. After solvent evaporation, the obtained compounds were used without further puriﬁcation.

Purity and structural conﬁrmation of the chemicals were obtained by GC-FID, RMN, and GC-MS analysis. Z9-14OH and Z9,E12-14:OH showed the high purity of the obtained compounds, conﬁrmed by GC-FID and GC-MS analysis, as follows: Z9-14OH (98.7 % purity) ¹H NMR (300 MHz, CDCl₃) δ 0.89 (t, 3H J= 6.9, CH₃), 1.30 (b, 14H, CH₂), 1.56 (m, 3H, CH₂ + OH), 2.00 (m, 4H, CH₂), 2.71 (m, 2H, CH₂), 3.63 (t, 2H J= 6.6, OCH₂), 5.34 (m, 2H, CH=) MS (70 eV): m/z (abundance) 41 (53); 55 (100), 67 (72), 68 (45); 69 (45); 81 (69); 82 (76); 95 (49), 96 (49); 109 (21); 111 (19); 123 (9); 124 (7); 137 (4), 138 (4); 166 (2); 194 (7), 212 (0.2), and for Z9,E12-14:OH (99 % of purity) ¹H NMR (300 MHz, CDCl₃) δ 1.30 (b, 10H, CH₂), 1.50-1.61 (m, 5H, CH₂), 1.65 (d, 3H, J= 4.71, CH₃), 2.01 (q, 2H, J = 6.42, CH₂), 2.71 (m, 2H, CH₂), 3.63 (t, 2H J= 6.5, OCH₂), 5.39 (m, 4H, CH=) MS (70 eV): m/z (abundance) 41 (37); 55 (81), 67 (92), 68 (100); 79 (53); 81 (84); 82 (56); 95 (47), 96 (22); 107 (8); 109 (10); 121 (10); 123 (9); 135 (6), 149 (3); 192 (0.1); 210 (1.3).

Behavioral bioassays were conducted in a wind tunnel described in Blassioli-Moraes et al. (2016). Bioassays with H. grandella males were conducted using 0.5 m s⁻¹ airflow. Treatments were spotted on 1.5 cm long and 0.5 cm wide ﬁlter paper strips (Whatman Nº 1), which were placed inside a metal cage (2-4 mesh). The cage was placed on a support 15 cm above the wind tunnel ﬂoor, and 30 cm from the upwind end of the tunnel. Males were released individually and, before testing, they were allowed to acclimate for 5 min inside the wind tunnel. In the ﬁrst set of the bioassay, male responses were evaluated with the odour released by ﬁve conspeciﬁc calling females. In the second set of the bioassay, H. grandella male responses were evaluated with synthetic solutions containing the four identiﬁed compounds in the H. grandella female gland extracts: two binary blends named mix-1 – Z9,E12-14:OH and Z9,E12-14:OAc –, at 0.075 µg each, and mix-2 – Z9,E12-14:OH and Z9,E12-14:OAc –, at 0.05 µg each; one ternary mixture, named mix-3, containing Z9-14:OH, Z9,E12-14:OH and Z9,E12-14:OAc, at 0.05:0.05:0.025 µg; and two quaternary mixtures, named mix-4 – Z9-14:OH,
Treatment B, which contained only two pheromone components, was not used because the preliminary results obtained in Fraiburgo in 2012 showed a low capture of insects compared to the lure containing the four components. The traps were set in a completely randomized design, in a 5.5 ha area within a *Swietenia macrophylla* plantation of one-to-eight-year-old trees. The traps were placed 1.50 to 1.70 m above the ground, with 100 m of distance between them, and they were evaluated at the end of the field test 28 days later. To prepare the treatment C, rubber septa were impregnated following the methodology described by Aldrich et al. (2006). The septa were impregnated with 1 mg of the synthetic pheromones Z9-14:OH, Z9,E12-14:OH, Z9-14:OAc and Z9,E12-14:OAc, at the same ratio between the components quantified from female gland extract. The statistical analyses were performed with R 3.0.1 (R Core Team, 2012). Data from the wind tunnel bioassays were analysed by a GLM with a binomial distribution. The proportion and confidence interval (95%) of responding insects to each treatment were also calculated. Data from the field experiment were analysed using a GLM model, with Poisson distribution of error, and the deviance analysis with 95% of confidence errors level, using the number of insects per trap as dependent variable, and treatment as fixed effect. The mean number of insects captured by treatment was compared by contrast analysis.

**Results and Discussion**

The chemical analyses of gland extracts of *H. grandella* females showed the presence of four peaks consistently present in the chromatograms: (9Z)-tetradecen-1-ol (Z9-14:OH) (1), (9Z,12E)-tetradecadien-1-ol (Z9,E12-14:OH) (2) (9Z)-tetradecenyl acetate (Z9-14:OAC) (4) and (9Z,12E)-tetradecadienyl acetate (Z9,E12-14:OAc) (6), and two minor peaks that were occasionally present in the samples (Table 1, and Figure 1). The four major compounds are common in sex pheromones of several other families of Lepidoptera (Acín et al., 2010). The mass spectra and the RI calculates of these compounds suggest that they could be (9Z,12Z)-9,12-tetradecadien-1-ol (Z9,Z12-14:OH) (3), and (9Z,12Z)-tetradecadienyl acetate (Z9,Z12-14:OAc) (7) (Figure 1). The authentic standards of these compounds were not available for this study to confirm the identification; therefore,
the calculation of the retention index on DB-WAX column and the co-injection were not conducted for these compounds. The co-injection of the synthetic compounds with the female extracts and GC peak enhancement confirmed the identities of the four major peaks (Figure 2). The ratio of the components Z9-14:OH, Z9,E12-14:OH, Z9-14:OAc, and Z9E12-14:OAc averaged 0.27:1:0.05:0.2 (Table 1). The compounds Z9,Z12-14:OH and Z9,Z12-14:OAc could be by-products of pheromone biosynthesis (Jurenka, 2004), since there was no response of male antennae to these compounds. In a coupled GC-electroantennography, H. grandella male antennae responded to Z9-14OH, Z9,E12-14:OH, Z9-14:OAc, and Z9,E12-14:OAc, from the gland extracts of H. grandella females (Figure 3 A). It is interesting to notice, that even for very small amounts of the components Z9-14:OAc and Z9,E12-14:OAc, the male antennae showed a clear response of depolarization-polarization to these compounds. However, there is no response to the compounds Z9,Z12-14:OH and Z9,Z12-14:OAc, which could indicate that they do not play a function as sexual attractant, but they might play an important cue to other moths to discriminate species. However, since the antennae response is dose-dependent (Rouyar

### Table 1. Compounds identified in gland extracts of Hypsipyla grandella females(1), average amount±standard deviation per gland, and retention index calculated using DB-5MS and DB-Wax column.

<table>
<thead>
<tr>
<th>Compound</th>
<th>Amount per gland (ng)</th>
<th>Retention index DB-5MS</th>
<th>Retention index DB-Wax</th>
</tr>
</thead>
<tbody>
<tr>
<td>Z9-14:OH</td>
<td>6.82±3.79</td>
<td>1664</td>
<td>2239</td>
</tr>
<tr>
<td>Z9,E12-14:OH</td>
<td>25.88±12.22</td>
<td>1675</td>
<td>2314</td>
</tr>
<tr>
<td>Z9,E12-14:OH</td>
<td>traces</td>
<td>1686</td>
<td>-</td>
</tr>
<tr>
<td>Z9-14:OAc</td>
<td>1.43±1.31</td>
<td>1795</td>
<td>2144</td>
</tr>
<tr>
<td>Z9,E12-14:OAc</td>
<td>4.57±2.00</td>
<td>1805</td>
<td>2233</td>
</tr>
<tr>
<td>Z9,Z12-14:OAc</td>
<td>traces</td>
<td>1816</td>
<td>-</td>
</tr>
</tbody>
</table>

(1)Volatiles were collected from 7 to 10 females for each replicate (n=5).

Figure 1. Chromatogram profile from GC-MS obtained from a gland extract containing eight glands of Hypsipyla grandella females. 1, Z9-14:OH; 2, Z9E12-14:OH; 3, Z9Z12-14:OH; 4, Z9-14:OAc; 5, n-octadecane; 6, Z9E12-14:OAc, and 7, Z9Z12-14:OAc. X, contaminant peak.

Figure 2. Chromatographic profile of A, female gland extracts; B, synthetic solution containing the four components (1, Z9-14:OH; 2, Z9E12-14:OH; 4, Z9-14:OAc; 6, Z9E12-14:OAc); and C, co-injection of a synthetic solution with female gland extract.
et al., 2015; Badeke et al., 2016), further studies are necessary to verify this hypothesis. In the wind tunnel bioassays, the conspecific calling females used as odour source stimulated 42.8% of *H. grandella* males to display the reproductive behaviour, which consisted of antennation, genital exposure, and zig-zag flights towards the odour source; however, none of the tested males landed on the odour source (Figure 4). Therefore, a positive response was considered whether the tested males displayed reproductive behaviour and fly in direction to odour source. When stimulated with semiochemicals from plants or insects pheromones, both males and females present different behaviour patterns, like oriented flight, antennation, and genital exposure (von Arx et al., 2011). In the present study, *H. grandella* males, stimulated with gland female extract and synthetic mixtures of the sex pheromone, changed their foraging behaviour. The wind tunnel bioassays with males of *H. grandella* with the synthetic compounds identified in the conspecific female gland extracts induced the attraction of males to the odour source. When a binary mix containing Z9-14:OH and Z9,E12-14:OH at 0.075 µg (mix-1) was tested, 50% of males were induced to display mating behaviour. Mix-2 with the same ratio of these two components, but at lower amounts than mix-2 (0.05 µg), elicited responses in 48.5% of males. Mix-3 with the three components Z9-14:OH; Z9,E12-14:OH and Z9-14:OAc at 0.05:0.05:0.025 µg for each compound, respectively, induced responses in 59.6% of the *H. grandella* males. Mix-4 containing the four compounds Z9-14:OH; Z9,E12-14:OH; Z9-14:OAc, and Z9,E12-14:OAc, at 0.27:1:0.05:0.2 µg for each compound, respectively, induced 61% of males to display mating behaviour. And a mixture containing the four compounds with 0.05 µg of each component induced 76.9% of *H. grandella* males to display mating behaviour (Figure 4). Results show that in wind tunnel bioassays *H. grandella* males showed a higher-response level, when evaluated with synthetic solutions than when they were stimulated with calling live females. In general, the blends of female gland extracts of Lepidoptera from different families present redundancy in chemical composition, that is, there are some compounds that provide the same information (Bruce & Pickett, 2011). Therefore, these compounds can be removed from the blend without negatively or positively interfering on the male attraction (Bruce et al., 2010; Bruce & Pickett, 2011). For instance, in the female gland extracts of *Trichoplusia ni* Hübner (Lepidoptera:Noctuidae), six...

**Figure 3.** A, coupled gas chromatography-eletroantennogram (GC-EAD) of two male antennae of *Hypsipyla grandella* to gland extracts; and B, male antenna of *H. grandella* responding to a synthetic solution containing the four compounds: 1, Z9-14:OH; 2, Z9,E12-14:OH; 3, Z9-14:OAc; and 4, Z9E12-14:OAc

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components were identified, all of them are involved in the sex pheromone communication system of *T. ni*; however, when tested blends of five or four components were used as odour source, *T. ni* males showed an analogous response level to those observed with six components blends (Linn Jr. et al., 1984).

Similar results were found in other populations of Neotropical lepidopterans, as follows: *Spodoptera frugiperda* Smith (Lepidoptera: Noctuidae), with eight compounds in sex pheromone blends, three of which are attractive to males (Batista-Pereira et al., 2006); *Spodoptera eridania* Cramer, with five compounds in sex pheromone blends, four of which are attractive to males (Teal et al., 1985); *Elasmopalpus lignosellus* Zeller (Lepidoptera: Pyralidae), with 10 compounds in sex pheromone blends, four of which are attractive to males (Lynch et al., 1984); and *S. cosmioides* (Walk.) (Lepidoptera: Noctuidae), with seven compounds in sex pheromone blends, two of which are attractive to males (Blassioli-Moraes et al., 2016). In gland extracts of *H. grandella* females, six compounds were identified and, similarly to other moths, not all compounds appear to be necessary to elicit reproductive behaviour of conspecific males (Byers, 2013).

Previous studies partially identified the pheromone blend of *H. grandella*. Borek et al (1991) worked with a population from Matanzas (Cuba), and identified three components from female gland extracts: Z9,E12-14:OH, Z3-16:OAc, and 16:OAc. The compounds Z3-16:OAc and 16:OAc were not identified in gland extracts obtained from *H. grandella* females in the present study. These different blends identified in populations from Cuba and Brazil might indicate specific geographic sex pheromone blends originated by reproductive isolation of *H. grandella*, as it has been shown for other Lepidoptera, when geographical isolation conducted to variance of sex pheromone blends (Groot et al., 2009).

In the present study, three different treatments were tested in two field experiments, in Fraiburgo, SC, and one field experiment in Garça, SP, Brazil. In all fields, the treatment C (Z9-14:OH, Z9,E12-14:OH, Z9-14:OAc and Z9E12-14:OAc), with the four compounds, captured a higher number of *H. grandella* males than the other treatments, corroborating the results obtained in the wind tunnel bioassays. Over the three weeks of the first experiment conducted in Fraiburgo, 16 males of *H. grandella* were trapped in treatment C. The GLM and contrast analysis showed that the capture was different between the traps of different treatments (*t* = 35.15, df = 2, *p* < 0.001) (Figure 5 A). The septum impregnated with two components, and the control septum impregnated only with hexane did not capture any insect. In 2014, the second field test, conducted in Fraiburgo during fifteen weeks, captured 52 males of *H. grandella* in the treatment C, and 19 males in treatment B (Z9-14:OAc:Z9,E12-14:OAc), with the two major compounds; the septum impregnated with solvent did not capture *H. grandella* males. The effect of treatments on the captures (χ² = 73.52 df = 2, *p* < 0.001) (Figure 5 A) and the contrast analysis showed differences between the treatments C and B (χ² = 3.76, df = 93, *p* = 0.0003). The higher efficiency of the four component blends could be explained because the major components are used by male moths at a long distance, and generate an initial attraction response in males, while minor components should play a role in approaches and courtship (Zhang et al., 2012). In the field experiment set in Garça, SP, Brazil, the traps containing the four compounds captured 10 *H. grandella* males, over 28 days, and the control traps captured only 1 male (χ² = 15.25 df = 1, *p* < 0.001).

Figure 4. Proportion of male responses, in wind tunnel bioassays, for five females of *Hypsipyla grandella*, and for synthetic solutions. Mix-1, Z9,E12-14:OH and Z9,E12-14:OAc, at 0.075 µg each; mix-2, Z9,E12-14:OH and Z9,E12-14:OHAc, at 0.05 µg each; a ternary blend, mix-3, containing Z9,E12-14:OH, Z9,E12-14:OH and Z9,E12-14:OAc, at 0.05:0.05:0.025 µg; and two quaternary blends, mix-4 with Z9-14:OH, Z9,E12-14:OH, Z9-14:OAc, and Z9,E12-14:OAc (2.7:10:0.02:0.6 µg), and mix-5, with Z9-14:OH, Z9,E12-14:OH, Z9-14:OAc, and Z9,E12-14:OAc, at 0.05 µg each compound. Error bars show 95% confidence interval.

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The results of the field experiments corroborate those obtained by the wind tunnel bioassays. *Hypsipyla grandella* was more attracted to quaternary solutions – \( Z_9\text{-14:OH} \); \( Z_9\text{-12:14:OH} \); \( Z_9\text{-14:OAc} \); and \( Z_9\text{-12:14:OAc} \) – than to binary solutions containing only the two major compounds \( Z_9\text{-14:OAc} \) and \( Z_9\text{-12:14:OAc} \). It is well established that in Lepidoptera minor compounds identified from female gland extracts are important to the response specificity and signal recognition in males (Linn Jr. et al., 1984; Groot et al., 2009).

The results presented here suggest that the sex pheromone blend of the studied *H. grandella* population from Southern Brazil have a different composition of those populations from other parts of the world. Therefore, they should be studied before their use for population monitoring or control in field conditions (Borek et al., 1991). Additional field experiments are required to assess the efficiency of pheromone traps to be used as a monitoring tool in the field, to evaluate the relationships between trap catch and population densities, as well as to find the best ratio and amount of the components in the baits used in traps. The identification of a correct sex pheromone blend for this species is an important step in the developing of an integrated method for *H. grandella* management aimed at the commercial production of meliaceous trees and, consequently, at the preservation of trees in natural forests.

**Conclusions**

1. The sex pheromone of gland of mahogany shoot borer (*Hypsipyla grandella*) females plays an important role in the attraction of conspecific males.

2. The female sex pheromone of *H. grandella* of a Southern Brazil population is composed at least of four components (9\(Z\))-9-tetradecen-1-ol, (9\(Z\),12\(E\))-9,12-tetradecadien-1-ol, (9\(Z\))-9-tetradecenyl acetate and (9\(Z\),12\(E\))-9,12-tetradecadienyl acetate; the binary, ternary, and quaternary blends of these compounds are able to attract males in laboratory experiments.

3. The quaternary mixture is able to attract males under field conditions, which indicates that this blend has potential to aid in the field monitoring of mahogany shoot borer.

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**References**


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