Stereochemistry of two pheromonal components of the bumblebee wax moth, *Aphomia sociella*

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The bumblebee wax moth, *Aphomia sociella*, is a parasite of bumblebees. In this species, males produce sexual pheromone to attract females, while females produce an aphrodisiac pheromone that initiates male courtship. Both pheromones contain 6,10,14-trimethylpentadecan-2-one (TMPD-one) and the corresponding alcohol, 6,10,14-trimethylpentadecan-2-ol (TMPD-ol) in sex specific quantities. Male sex pheromone consists of 7 components with TMPD-one as a minor one and traces of TMPD-ol. In female aphrodisiac pheromone, TMPD-ol is the major component, while TMPD-one is present in traces. Here we report on the absolute configuration of TMPD-one in male sex pheromone and TMPD-ol in female aphrodisiac pheromone of *A. sociella*. The configuration was determined from GC/MS of prepared (S)-acetoxypropionyl esters of TMPD-ol. TMPD-one was first reduced to the alcohol that was then derivatized with (S)-acetoxypropionyl chloride. The GC/MS data of obtained diastereoisomers were compared with synthetic standards. The absolute configuration of TMPD-one in the male pheromone was (6R,10R). The configuration of TMPD-ol in the female pheromone was (2R,6R,10R). Electrophysiological experiments showed that TMPD-one and TMPD-ol are perceived by both sexes. The synthetic standards of naturally produced stereoisomers elicited higher responses than mixtures of all stereoisomers.

The stereochemistry plays an important role in the pheromone communication¹. Therefore, it is of great importance to determine the stereoisomeric composition of pheromonal components. *Aphomia sociella* L. (Pyralidae, Galleriinae) is a parasite of bumblebees. Unlike most Lepidoptera, the reproductive behavior of *A. sociella* is initiated by males and except chemical communication (sex pheromone) it includes also an ultrasonic part. The male sex pheromone attracts females from a distance, while ultrasonic signaling urges females to accept mating². Females produce aphrodisiac pheromone, which initiates male courtship behavior¹. Male pheromone is a multicomponent blend consisting of (Z)-nona-2,6-dien-4-olide and mellein as major components, and 1-hexanol, 2-phenylethanol, (Z)-nona-6-en-4-olide, and 6,10,14-trimethylpentadecan-2-one (TMPD-one, trivial names phytone or hexahydrofarnesylacetone) as minor components³⁴. In female aphrodisiac pheromone, 6,10,14-trimethylpentadecan-2-ol (TMPD-ol) represents the major component with TMPD-one and hexan-1-ol as minor ones³.

TMPD-one has been found earlier within male sex pheromone of other Galleriinae, such as *Galleria mellonella*⁵ and *Tirathaba mundella*. TMPD-ol plays a role in the communication of another species of this family, *Corcyra cephalonica*, where females utilize TMPD-ol for attracting males⁶. Both TMPD-one and TMPD-ol are chiral compounds, but in none Galleriinae species, the absolute configuration has been reported.

The stereochemistry of these two compounds was determined in other insects beside Galleriinae. TMPD-ol in *Bicyclus* butterflies has been determined after (S)-2-acetoxypropionyl derivatization and the formed

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diastereomers were separated by gas chromatography (GC)\textsuperscript{8,9}. The stereochemistry of TMPD-one found in Euglossa bees was reported by Eltz\textsuperscript{11}, where reduction of ketone was done prior to derivatization. This method was also used to determine the stereochemistry of TMPD-one extracted from Bicyclus butterflies\textsuperscript{10}. Here we present the stereochemistry of TMPD-one in male sex pheromone and TMPD-ol in female aphrodisiac pheromone of \textit{A. sociella} as well as the electroantennographic activity of these compounds. In addition, we also show for the first time that males produce minute amount of TMPD-ol and that female antennae respond to this compound.

Materials and Methods

Insects. Laboratory-started nests of buff-tailed bumblebee (\textit{Bombus terrestris}) were obtained from The Research Institute of Fodder Crops (Troubsko, Czech Republic)\textsuperscript{12}. The nests were exposed to colonization by \textit{A. sociella} L. (Pyralidae, Galleriinae) in the field (Prague area). In June/July, the bumblebee hives were situated in the garden and let parasitize by natural population of \textit{A. sociella}. In autumn, the infested hives with larvae were transferred to refrigerator and kept 1 month at 5 °C for the larvae to undergo hibernation. After hibernation, the hives were kept in the rearing room (24 °C, 40% relative humidity). Larvae were allowed to develop and pupate in the bumblebee hives. The newly emerged adults were collected and sexed. The sexes were kept separately first in continuous light under ambient laboratory temperature (20–23 °C). Adults (1–2 days old) were transferred to the dark and used for pheromone extraction. Moths (one day post-emergence) for electrophysiological experiments were kept in a refrigerator with the temperature set to 5 °C and in a high-humidity environment (90%) until used.

Sample preparation. Male sex pheromone. The pheromone is produced in wing glands that are located on the basal part of the forewing, analogically as in \textit{ Eldana saccharina} Walker (Lepidoptera, Galleriinae) published by Farine\textsuperscript{13}. Glands of 20 calling males were clipped out, pooled together and soaked in redistilled \textit{n}-hexane (Merck, 50 µL per gland). Samples were sonicated for 15 min, and extracted for 24 h. The extracts were filtered through purified glass wool, concentrated under argon flow to double the initial concentration, and stored at −20 °C.

Female aphrodisiac pheromone. Virgin females (1–2 days old) were placed individually in cages, and positioned next to caged calling males. Females (20 specimens) that elicited calling songs in males were cooled to 20 °C. Next to caged calling males. Females (20 specimens) that elicited calling songs in males were cooled to 20 °C.

Analysis of crude extract. Extracts from males and females were analyzed using a Hewlett-Packard 6890 N gas chromatograph (GC) equipped with a polar Varian FactorFOUR VF-23ms column (30 m × 0.25 mm i.d., δ 1 = 0.25 µm) and a HP 5973 MS-detector in electron ionization mode (EI). The carrier gas (1 mL min\textsuperscript{−1}) was helium; the sample (1 µL) was injected in the splitless mode, the injector temperature was 250 °C, the transfer line was maintained at 250 °C and the MS source was set to 280 °C. For identification of the alcohol and ketone in crude extracts, the full scan mode was used and the column temperature increased from 50 °C by 10 °C min\textsuperscript{−1} up to 230 °C and held at 230 °C for 10 minutes. Mass spectra of main extract components are in the Supplementary material (Figs. S1–S3).

Purification of insect extracts. Extracts were purified using liquid chromatography (LC). Prior purification, female extracts were washed with NaHCO\textsubscript{3} (sat. aq.) to remove free carboxylic acids. Afterwards, male and female extracts were treated in the same way. In LC purification, pre-conditioned solid phase extraction column (500 mg, SPE, Chromabond) were used. SPE conditioning was done with EtOAc (1.5 mL), 10% EtOAc in pentane (1.5 mL) and pentane (1.5 mL). Then the columns were gradually eluted in steps of 1% increase of ethyl acetate in pentane (0–15%) with each fraction consisted of a volume of 750 µL. TMPD-one was enriched in fractions 3–6 and TMPD-ol was enriched in fractions 9–11. In cases where the purity was not satisfactory, the combined fractions were subjected to the purification protocol again until no further improvement of purity was obtained. The combined TMPD-one fractions were then reduced to alcohols (described below). After reduction, the TMPD-ol fractions and those containing the reduced TMPD-one were derivatized and analyzed as described below (Fig. 1D).

Reduction of TMPD-one fraction. The collected ketone fractions were pooled and solvent evaporated under a stream of argon. The purified ketone was diluted in dry diethyl ether (1 mL) and 0.2 mL was reduced to alcohol with lithiumaluminumhydride in diethyl ether (1 M, 50 µL) at 0 °C for 1 hour (Fig. 1B). The reaction was quenched by addition of HCl (2 M, 100 µL) and H\textsubscript{2}O (100 µL). Phases were separated and the aqueous layer was extracted with Et\textsubscript{2}O (3 × 200 µL), the combined organic layers was washed with HCl (2 M, 100 µL) and brine (100 µL) and dried over Na\textsubscript{2}SO\textsubscript{4} (anhdr.) in a Pasteur pipette. The solvent was evaporated under a stream of argon. The sample was diluted with pentane (50 µL) and purified with the same SPE purification method as the crude extracts, prior to derivatization.

Derivatization. To determine the stereochemistry of TMPD-ol enriched from female extract and TMPD-one enriched from male extract of \textit{A. sociella}, both the pooled alcohol fraction and the reduced ketone fraction were derivatized with (S)–2-acetoxypropionyl chloride (SigmaAldrich) according to Bánó\textsuperscript{14}. Pooled extracted TMPD-ol and reduced TMPD-one were concentrated by means of evaporating solvent under a stream of argon. Dry cyclohexane (50 µL) was added to the vial containing the purified extract and subsequently was 1% pyridine in cyclohexane (65 µL) and 1% (S)-2-acetoxypropionyl chloride in dichloromethane (100 µL) added, the resulting mixture was heated at 80 °C for 30–60 minutes under argon atmosphere (Fig. 1D). A needle was inserted into the septum of the vial and the solvent was allowed to evaporate before allowing the vial to reach room temperature.
HCl (1 M, 200 µL) and pentane or cyclohexane (1 mL) was added, the organic layer was subsequently dried over Na2SO4 (anhydr.) in a Pasteur pipette. Solvent was either reduced to approximately 50 µL and the sample was analyzed as described below or the solvent was evaporated completely and sample dissolved in cyclohexane before analysis.

**Stereochemical analysis.** The derivatized TMPD-ol was analyzed using GC/MS as described above for the crude extracts. The temperature program was adjusted, starting at 50 °C, increased to 110 °C at a rate of 10 °C min⁻¹, further from 110 °C by 0.01 °C min⁻¹ up to 115 °C, and from 115 °C by 10 °C min⁻¹ up to 230 °C, and held at 230 °C for 10 minutes. Selected ion monitoring (SIM) (m/z 105, 115, 133, 210, 252) was used for identification of the (S)-2-acetoxypropionyl derivatives.

**Preparation of reference compounds.** (2R,S,6R,S,10R/S)-6,10,14-Trimethylpentadecan-2-ol (2E7/R/S,11R/S)-(3,7,11,15-Tetramethyl-2-hexadecan-1-ol ((2E7/R/S,11R/S)-phytol) (SigmaAldrich) was oxidized at room temperature by sodium periodate and catalytic amount of ruthenium (III) chloride (Fig. 1A), the obtained ketone was reduced with LiAlH₄ in diethyl ether resulting in a mixture of eight isomers of TMPD-ol (Fig. 1B) according to the supplemental material of Nieberding et al. The synthetic mixture was derivatized and analyzed according to the same method as the purified extracts.

(2S,6R,10R)-6,10,14-Trimethylpentadecan-2-ol and (2R,6R,10R)-6,10,14-trimethylpentadecan-2-ol (2E7/R,11R)-Phytol (TCI America) was treated as described above, resulting in a mixture of (2R,6R,10R)- and (2S,6R,10R)-TMPD-ol (Fig. 1B). The isomeric mixture was subjected to enzymatic resolution employing Candida antarctica lipase-B (CALB, Roche Diagnostics batch 90750729) as catalyst and vinylacetate (SigmaAldrich) as acyl-donor (Fig. 1C). With this method the (2R)-alcohol is acetylated and therefore the two isomers are easily separated by means of liquid chromatography. The obtained (2R)-acetate was reduced with litiumaluminum hydride in diethyl ether (Fig. 1B). A diastereomeric purity of 99.8% for (−)-(2R,6R,10R)-TMPD-ol and 97.8% diastereomeric purity of (+)-(2S,6R,10R)-TMPD-ol was obtained as described in the supplemental material of Nieberding et al. The stereoisomically pure isomers were derivatized and analyzed according to the same method as the purified extracts.

**Electroantennography (EAG).** Detailed description of EAG measurements on A. sociella is available in Kalinová et al. Briefly, isolated antennae, glass Ag/AgCl electrodes and Syntech EAG recording and stimulus delivery system were used. Male and female antennal responses to stimulation by 10, 100, and 250 ng of racemic mixtures and pure enantiomers of TMPD-one and TMPD-ol were investigated. Tested compounds were diluted in n-hexane. Aliquots of 10 µL of respective solutions were loaded onto the filter paper discs within Pasteur pipette odor cartridges. During stimulation, 0.5 mL of humidified air was flown through loaded odor cartridge and injected into the air stream directed onto the antennal preparation. For each sex and stimulus type, 6 replicates were performed.

The antennal responses from each series of stimulations were normalized to the stimulations by 10 ng of 1-hexenol using the Syntech software and expressed as percents of 1-hexenol responses. For statistical evaluation, the data were subjected to logarithmic transformation to correct for the heteroscedasticity inherent to EAG data. The resulting datasets complied with the assumption of equal variances (tested using Levene's test) and the
means were compared using ANOVA followed by Least Squares Differences post-hoc multiple comparison test, performed in Statistica 8.0.

**Dedication.** Dedicated to the memory of Prof. Kenji Mori and his significant contribution to the field of chemical ecology.

**Results and Discussion**

**Purification of extracts, derivatization, and analysis.** GC/MS has been previously used to determine the stereoisomeric composition of TMPD-ol and TMPD-one in wing extracts of *Bicyclus* butterfly species, stereoisomeric composition of TMPD-one in *Euglossa* bees and of 3,7-dimethylpentadecan-2-ol in female extracts of pine sawflies, where the stereoisomers of the chiral alcohols were separated as either (R)-trans-chrysanthemoyl or (S)-2-acetoxypropionyl esters. Here we used (S)-2-acetoxypropionyl derivatization method, because it is robust and more suitable for derivatization of compounds present at low concentrations.

Male gland extracts gave stronger samples than female body extracts obtained from one specimen (Fig. S4). TMPD-one and TMPD-ol were found in both sexes, but their ratio in the sexes differed. In males, the ketone was present in substantially higher amount than the alcohol while in females, the alcohol dominated over the ketone. In the male wing glands, the trace amount of TMPD-ol was on the detection limit and the compound was only found after pre-cleaning the extract with SPE. A rough estimate of TMPD-ol quantity produced by females and TMPD-one produced by males was done using dodecanol as internal standard. In average, males produce 2–7 ng TMPD-one and females produce 1–5 ng TMPD-ol per individual.

Obtaining pure TMPD-ol from male glands was complicated due to its co-elution with mellein, which is a major component of the male-produced pheromone (Fig. 2A, crude male extract, main peak). In the GC analysis of crude extracts TMPD-ol and TMPD-one co-elute, as a result of this a relatively large peak can be seen at the retention time of TMPD-ol/TMPD-one in the crude extract (Fig. 2A, dotted line). After the first run of purification, mellein still dominated the fraction containing TMPD-ol (Fig. 2B, fraction containing TMPD-ol and mellein; Fig. 2C, chromatogram B zoomed in on y-axis to show the presence of TMPD-ol in collected fractions).

**Figure 2.** Gas chromatogram of crude male extract, purified male TMPD-ol which is co-eluting with mellein during SPE purification. Dotted line indicates the retention time for TMPD-ol/TMPD-one. (A) Crude male extract. (B) Full chromatogram of fraction containing TMPD-ol and mellein. (C) Chromatogram B zoomed in on y-axis to show the presence of TMPD-ol in collected fractions.
stereochemistry. Thus, we were able to determine the stereochemistry of those pheromonal components present in higher amounts, i.e. of TMPD-one in the males and TMPD-ol in the females. The synthetic mixture of eight stereoisomers and pure isomers of (2R,6R,10R)- and (2S,6R,10R)-TMPD-ol were utilized for comparisons on GC/MS. The pure isomers were synthesized by Hedenström while working with Euglossa bees and Bicyclus butterflies. The eight-stereoisomers mixture of TMPD-ol was synthesized according to Nieberding (supplemental material). All 8 derivatized stereoisomers separated well as (S)-2-acetoxypropionyl esters on the highly polar VF-23ms column (Fig. 3A). The male-extracted, reduced and derivatized pheromonal component, matched with the authentic standard of derivatized (2R,6R,10R)-TMPD-ol (Fig. 3B) leading to a conclusion that males produce (6R,10R)-TMPD-one (Fig. 3C). During reduction of TMPD-one a new stereocenter is introduced in position 2, that is the reason to why two peaks appear in the chromatogram after derivatization (Fig. 3C). Due to co-elution during SPE purification of the male-produced TMPD-ol with melilin (Fig. 2) however, it was not possible to determine the stereochemistry of the native male TMPD-ol present in male wing glands in minute quantity. Female-produced TMPD-ol was determined as (2R,6R,10R)-TMPD-ol based on the match of the extracted and derivatized compound with the synthetic standard. The GC/MS analysis of female-produced and derivatized TMPD-ol showed one more peak at 397.5 min (Fig. 3D). However, retention time and mass spectral data did not match with any of the TMPD-ol isomers.

Schulz showed that the Large Cabbage White, Pieris brassicae, sequester (R,R)-phytol from food sources and via oxidative processes transform phytol to (6R,10R)-TMPD-one. Eltz reported on Euglossa bees carrying (6R,10R)-TMPD-one in their hind-leg pouch which is believed to be collected from a natural source, despite the source of TMPD-one has not been identified. This shows that some insects utilize the naturally derived (R,R)-phytol as a precursor. Since we kept A. sociella in the old bumblebee hives and not on an artificial diet of known composition, we cannot tell what the source of (2R,6R,10R)-TMD-ol was in this particular case.

The stereoisomers of TMPD-one found in males and TMPD-ol in females of A. sociella have the same configuration on chiral centres (2R,6R,10R) as 9 butterfly species of the genus Bicyclus studied earlier by Hedenström and as male orchid bees of the genus Euglossa (supplemental material). On the other hand, 3,7-dimethylpentadecan-2-ol, present in the female sex pheromone of the pine sawfly Neodiprion lecontesi, has the configuration (2S,3S,7S). The stereochemistry of TMPD-one and TMPD-ol in pheromones of related species of the subfamily Galleriinae (Galleria mellonella, Tirathaba mundella or Corcyra cephalonica) has not been studied so far. Thus, this paper is the first report on the configuration of the above pheromonal components within Galleriinae.

**Electroantennography.** To determine the biological activity of native stereoisomers of TMPD-one and TMPD-ol, antennal responses of males and females of A. sociella were recorded and compared to those of a mixture of eight stereoisomers for TMPD-ol and/or a mixture of four stereoisomers for TMPD-one. Three concentrations for each treatment were tested to see the dose-response dependence (Fig. 4). The lowest selected concentration (10 ng) was approximately of the same order as 2 gland equivalents of males (wing glands are pair organs) or 1 body equivalent of females (the source of pheromone is not known, therefore the extract of the whole body was used). TMPD-one and TMPD-ol were perceived by both sexes. The lowest concentrations of both compounds elicited responses that were not significantly different from the solvent (n-hexane). Higher doses elicited higher antennal responses that were statistically different from the solvent at the levels 100 ng or 250 ng. Female antennal reactions were higher in response to native stereoisomer of TMPD-one (100 ng) in comparison with TMPD-one stereisomeric mixture (Fig. 4A), while males responded in the same way on a higher concentration level (250 ng, Fig. 4C). On the other hand, both sexes responded significantly more strongly to the native stereoisomers of TMPD-ol than to the stereoisomeric mixture, both at the concentration level of 100 ng and 250 ng (Fig. 4B,D).
Conclusion

Based on the purification and derivatization of the extract of male and female pheromone of *A. sociella*, we have determined the absolute configuration of TMPD-one in the male sex pheromone as (6R,10R)-6,10,14-trimethylpentadecan-2-one and of TMPD-ol in the female aphrodisiac pheromone as (2R,6R,10R)-6,10,14-trimethylpentadecan-2-ol. The pure stereoisomers elicit significantly stronger antennal responses than the tested stereoisomeric mixtures of both TMPD-ol and TMPD-one, respectively. We are aware that in the tested stereoisomeric mixtures only one forth (TMPD-one) or one eights (TMPD-ol) of the naturally produced isomer is present. We are showing here that the antennal responses increased with dose, for both the pure enantiomers as well as the mixtures of all stereoisomers. EAG experiments suggest that the antennal perception may be enantioselective. The significant difference in the perception of the pure isomer and the mixture might mean that none of the non-natural stereoisomers is antennaly active, has synergic affect, or influences the antennal response in any way. However, further experiments with pure stereoisomers are needed to support this hypothesis.

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Figure 4. Electroantennographic responses of females (A,B) and males (C,D) of *Aphomia sociella* to either equimolar mixtures of all stereoisomers or to pure standards of natural isomers (dose-response diagrams). (A) Responses of females to the mixture of 4 isomers of TMPD-one compared to its (6R,10R)-isomer. (B) Responses of females to the mixture of 8 isomers of TMPD-ol compared to its (2R,6R,10R)-isomer. (C) Responses of males to the mixture of 4 isomers of TMPD-one compared to its (6R,10R)-isomer. (D) Responses of males to the mixture of 8 isomers of TMPD-ol compared to its (2R,6R,10R)-isomer. The data were normalized to the stimulations by 10 ng of 1-hexenol and expressed as percents of 1-hexenol responses (mean ± SD).
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Author contributions
E.W. did the synthesis reference compounds, purification of insects extracts, derivatization, chromatography, Figures. 1–3, writing the article. B.K. did electroantennography, Figures. 4 and writing the article. J.K. reared and dissected insects, made extracts. E.H. supervised and consulted the synthesis and participated in writing the article. I.V. headed the project, designed the study, made supplementary figures and wrote main part of the article.

Competing interests
The authors declare no competing interests.

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