Hero Turned Villain: Identification of Components of the Sex Pheromone of the Tomato Bug, *Nesidiocoris tenuis*

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**Abstract**

*Nesidiocoris tenuis* (Reuter) (Heteroptera: Miridae) is a tropical mirid bug used as a biocontrol agent in protected crops, including tomatoes. Although *N. tenuis* predates important insect pests, especially whitefly, it also causes damage by feeding on tomato plants when prey populations decline, resulting in significant economic losses for growers. The pest is now established in some all-year-round tomato crops in Europe and control measures involve the application of pesticides which are incompatible with current IPM programs. As part of future IPM strategies, the pheromone of *N. tenuis* was investigated. Volatile collections were made from groups and individuals of mated and unmated, females and males. In analyses of these collections by gas chromatography coupled with electroantennographic (EAG) recording from antennae of male bugs, two EAG-active components were detected and identified as 1-octanol and octyl hexanoate. Unlike other mirids, both male and female *N. tenuis* produced the two compounds, before and after mating, and both sexes gave EAG responses to both compounds. Furthermore, only octyl hexanoate was detected in whole body solvent washes from both sexes. These compounds are not related to the derivatives of 3-hydroxybutyrate esters found as pheromone components in other members of the Bryocrinae sub-family, and the latter could not be detected in volatiles from *N. tenuis* and did not elicit EAG responses. Nevertheless, experiments carried out in commercial glasshouses showed that traps baited with a blend of the synthetic pheromone components caught essentially only male *N. tenuis*, and significantly more than traps baited with octyl hexanoate alone. The latter caught significantly more *N. tenuis* than unbaited traps which generally caught very few bugs. Traps at plant height caught more *N. tenuis* males than traps 1 m above or at the base of the plants. The trap catches provided an indication of population levels of *N. tenuis* and were greatly reduced following an application of insecticide.

**Keywords** GC-EAG · IPM · Mirid · 1-octanol · Octyl hexanoate · Hexyl octanoate

**Introduction**

*Nesidiocoris tenuis* (Reuter) (Heteroptera: Miridae) is an omnivorous mirid bug of tropical origin which is used as a biological control agent for protected crops in Mediterranean countries (Biondi et al. 2016; Pérez-Hedo et al. 2021). The bug predates a range of economically important crop pests such as whiteflies (family Aleyrodidae), aphids (superfamily Aphidoidea), thrips (order Thysanoptera), and moths (order Lepidoptera) (Kim et al. 2016). However, the use of *N. tenuis* as a biological control agent is controversial. Although this bug has a preference for prey, it can start feeding on the crop when prey is absent (Nakaishi et al. 2011; Pérez-Hedo and Urbaneja 2016; Sanchez 2008). Insertion of the stylet into the phloem to derive nutrients leads to brown necrotic rings around stems and petioles followed by the drying of flower stalks and flower abortion (Arnó et al. 2010; Calvo et al. 2009; Castañé et al. 2011; Raman and Sanjayan 1984). In some northern European countries *N. tenuis* has become invasive in glasshouses and the pest is threatening the 180 ha of tomatoes grown in the UK (Jacobson 2019).

In circumstances where *N. tenuis* needs controlling, traditional control options might not be suitable. Although the bug cannot complete development to adulthood if feeding exclusively on tomato plants, *Solanum lycopersicum* (Urbaneja...
et al. 2005), growers may resort to the use of chemical plant protection products (PPPs), incompatible with IPM programs, to prevent further crop damage. This can lead to a resurgence of whitefly populations and associated viruses, and disruption of other biocontrol systems such as use of predatory mites against spider mites. Moreover, a number of PPP’s have been shown to have lethal and sublethal effects on pollinators (Cloyd 2012; Feltham et al. 2014), and it is recommended that their use is avoided unless absolutely necessary (PAN Europe 2013).

Semiochemicals could provide tools for IPM-compatible management of N. tenuis. This species belongs to the Bryocorinae sub-family of mirids, and female-produced sex pheromones have been identified in several species of this sub-family. These include the cocoa mirids, *Distantiella theobroma* Dist. and *Sahlbergella singularis* Haglund (Mahob et al. 2011; Mahot et al. 2020; Sarfo 2013; Sarfo et al. 2018a, 2018b), and the aphidophagous mirid, *Macrolophus pygmaeus* Rambur (formerly *Macrolophus caliginosus* Wagner) (Gemeno et al. 2006). The components of these pheromones are derivatives of esters of 3-hydroxybutyric acid, rather different from those identified in mirid species from other subfamilies which are typically saturated or unsaturated, straight-chain esters or aldehydes (Zhang et al. 2020 and refs therein). For cocoa mirids, the synthetic pheromone has been used for monitoring (Mahob et al. 2011; Mahot et al. 2020; Sarfo 2013; Sarfo et al. 2018a) and mass trapping (Sarfo et al. 2018b).

To the best of our knowledge, the presence of a pheromone has not been demonstrated in *N. tenuis*. However, virgin females of the closely related *M. pygmaeus* attracted males (Gemeno et al. 2006). In this project we aimed to identify the female-produced sex pheromone of *N. tenuis* and demonstrate attraction of male *N. tenuis* to the synthesized pheromone in commercial glasshouses. This will provide a basis for development of pheromone traps for better detection of this invasive pest, for monitoring of pest numbers and more efficient use of control agents. The pheromone could also be used for control of *N. tenuis* by mass trapping or mating disruption and hence reduce or avoid the use of conventional pesticides against this pest on protected crops.

**Methods and Materials**

**Chemicals**

Unless otherwise stated, chemicals were purchased from SigmaAldrich (Gillingham, Dorset, UK) and were at least 99% pure. Hexyl (R)-3-hydroxybutyrate and hexyl (R)-3-[(E)-2-butenoyl]-butyrate were synthesized as described previously (Padi et al. 2002), and hexyl (R)-3-acetoxybutyrate was prepared by acetylation of hexyl (R)-3-hydroxybutyrate with acetic anhydride and pyridine (Gemeno et al. 2006).

Octyl hexanoate used in the work described here was synthesized by reaction of 1-octanol with hexanoic acid in dichloromethane. The product was purified by flash chromatography on silica gel eluted with 2% diethyl ether in petroleum spirit (bp 40–60 °C) followed by Kugelrohr distillation (100 °C/0.06 mmHg) in 90% yield. Hexyl octanoate was prepared similarly by reaction of 1-hexanol and octanoic acid in 93% yield. Both compounds were characterized by their mass spectra and 1H and 13C NMR spectra (details in Supplementary Material). Octyl hexanoate has subsequently been prepared on molar scale in 95% yield by reaction of hexanoyl chloride with 1-octanol and pyridine in dichloromethane.

**Rearing of Nesidiocoris tenuis**

Young tobacco (*Nicotiana spp.*), aubergine (*Solanum melongena*), tomato (*S. lycopersicum*), and cucumber (*Cucumis sativus*) were grown as host plants. Plants were sown separately in 11 × 11 × 11 cm pots containing standard compost and kept in a glasshouse at 26 ± 3 °C and 16:8 h L:D photoperiod. Organic dwarf beans used in bioassays and virgin adult cultures were obtained from a local supermarket (Sainsbury’s plc, London, UK).

*Nesidiocoris tenuis* adults were purchased from Bioline AgroSciences Ltd. (Nesline) and a culture established in a quarantine facility at NIAB-EMR. Insects were reared in BugDorms (50 × 50 × 50 cm; MegaView Science, Taichung, Taiwan) on the host plants supplemented with sterile *Ephestia kuehniella* eggs (Nutrimac; Biobest, Ashford, UK), pollen and sugar solution (5% dextrose). Cultures were maintained at 20–26 °C and 55 to 65% relative humidity on a 16:8 h L:D photoperiod. BugDorm floors were lined with a layer of damp tissue to maintain humidity.

Initially, male and female adults were not separated for up to seven days before collections of volatiles were made. Mating was observed and nymphs were produced. To obtain virgin *N. tenuis* adults, third to fifth instar nymphs were housed individually in ventilated Perspex rearing boxes (8 × 6 × 14 cm). Nymphs were reared on sterile *E. kuehniella* eggs, a dwarf organic bean and 2.5 ml of sugar solution (5% dextrose) under the same conditions as BugDorm cultures. Virgin adults were kept in the individual rearing boxes and were 3–7 days old when collections of volatiles were made. Adults were sexed by observation of the abdomen with gentle pressure if necessary to show the paramere in males and ovipositor in females (Kim et al. 2016).
**Pheromone Collection**

For collection of volatiles, virgin males and females were carefully placed in separate, silanized glass chambers (12 cm × 4 cm) with a glass frit at the upwind end (Hamilton Laboratory Glass, Margate, Kent, UK) containing an aubergine shoot or dwarf organic bean as food and maintained under the same conditions as the cultures above. Insects were left to settle in the chambers for at least 30 min prior to starting collection, and chambers containing food source only were used as controls. Air was drawn into the chamber with a vacuum pump (DA7C; Charles Austen, West Byfleet, UK) at 200 ml/min through a filter containing activated charcoal (20 cm × 2 cm, 10–18 mesh; Fisher Scientific, Loughborough, UK). Volatiles were collected on filters containing Porapak Q (50–80 mesh; 200 mg; Supelco, Gillingham, Dorset, UK) held between silanized glass wool plugs in a Pasteur pipette (4 mm i.d.). The Porapak was purified by Soxhlet extraction with chloroform and washing with dichloromethane before use. Volatiles were collected for 24 or 48 h.

Volatiles were first collected from groups of five (N = 3; aubergine as food) or 20 (N = 2) adult females and males which were presumed to be mated. Collections were then made from groups of 20 (N = 2) and individual (N = 9) virgin males and females. Collections were also made from chambers containing only a dwarf bean (N = 7). Volatiles were eluted from the Porapak with dichloromethane (1 ml; Pesticide Residue Grade), concentrated to approx. 100 μl under a gentle stream of nitrogen and stored at 4 °C before analysis.

Whole-body extracts were made by immersing individual *N. tenuis* adults in diethyl ether (0.5 ml; SLR grade containing butylated hydroxytoluene, BHT, as antioxidant) for 10 min, then removing the ether, drying with a few grains of anhydrous magnesium sulfate and storing at 4 °C before the analysis. Extracts were made at approximately 10.00 h from individual females and males which were presumed to be mated (N = 7 and 8, respectively) and from virgin females and males (N = 2). Diethyl ether was used because potential compounds of interest such as (E)-4-oxo-2-hexenal are poorly soluble in hexane and are relatively stable in diethyl ether containing BHT (Fountain et al. 2014). An internal standard of decyl acetate (5 μg) was added for quantitative analysis.

**Analysis by Gas Chromatography Coupled to Electroantennographic Recording (GC-EAG)**

GC-EAG Analyses were carried out on a HP6890 GC (Agilent Technologies, Manchester, UK) fitted with flame ionization detector (FID) and fused silica capillary columns (30 m × 0.32 mm i.d. × 0.25 μm film thickness) coated with DBWax and DB5 (Supelco). Injections onto the DBWax column were in splitless mode (220 °C), carrier gas was helium (2.4 ml/min) and the oven temperature was programmed from 50 °C for 2 min and then at 20 °C/min. to 250 °C for 3 min. The effluents of the two columns were combined with a glass push-fit Y-tube connector (Agilent Technologies) connected to a second Y-tube connector with deactivated fused silica tubing (10 cm × 0.32 mm i.d.). One arm of this connector was connected with deactivated fused silica tubing (50 cm × 0.32 mm i.d.) to the FID (250 °C) and the other to an equal length of deactivated silica tubing passing through a heated transfer line (250 °C; Syntech, Hilversum, The Netherlands, now Kirchzarten, Germany) into a glass tube (4 mm i.d.) through which air passed (500 ml/min) over the EAG preparation. Both the FID and EAG signals were collected and analyzed with EZChrom software (Elite v3.0; Agilent Technologies).

For EAG recordings, adult *N. tenuis* were anesthetized using carbon dioxide, and the head and one antenna removed under a dissecting microscope with a razor blade. A borosilicate glass capillary electrode (ID 0.86 mm, Warner Instruments, Hamden, CT 06514), pulled to a fine tip and filled with Beadle-Ephrussi Ringer containing 1% polyvinylpyrrolidone as electrolyte, was inserted into the back of the head. The electrode and head were then mounted onto a silver wire held within an electrode holder connected to the earth probe of a portable EAG amplifier (INR-2, Syntech). A similar electrode mounted onto the ×10 recording preamplifier was then brought into contact with the distal tip of the antenna. Straightening the antenna between the electrodes reduced background noise from the preparation.

**Analysis by Gas Chromatography Coupled to Mass Spectrometry (GC-MS)**

GC-MS Analyses were carried out on a CP3500 GC coupled to a CP2200 Saturn Ion Trap Detector (Varian, now Agilent Technologies). The GC was fitted with fused silica capillary columns (30 m × 0.25 mm i.d. × 0.25 μm film thickness) coated with DBWax (Supelco) and VF5 (Varian) with a switching device to select the column used. Carrier gas was helium (1 ml/min) and manual injections (1 μl) were made in splitless mode (220 °C), with oven temperature programmed from 40 °C for 2 min then at 10 °C/min to 240 °C. Compounds were identified according to their mass spectrum, retention index relative to retention times of n-alkanes, and co-chromatography with authentic compounds.

**Analysis by Gas Chromatography with Flame Ionization Detection (GC-FID)**

GC-FID Analyses were carried out on HP6850 instruments (Agilent) fitted with fused silica capillary columns (30 m ×
0.32 mm i.d. × 0.25 μm film thickness) coated with polar DBWax (Supelco) or non-polar HP5 (Agilent) and FID detectors (250 °C). Carrier gas was helium (2.4 ml/min), injection was splitless (220 °C and 250 °C, respectively) and the oven temperature was programmed from 50 °C for 2 min then at 10 °C/min to 250 °C. Data were collected and analyzed with EZChrom software (Elite v3.0; Agilent).

Controlled Release Dispensers

Two types of controlled-release dispenser for the pheromone components were investigated. The first were opaque, polypropylene pipette tips (1 ml; Fisher Scientific) with a 0.2 mm tip aperture, sealed with a Teflon-lined crimp seal (11 mm; Chromacol, Welwyn Garden City, UK) as developed for the pheromone of Lygus bugs by Fountain et al. (2014). The second were sealed low-density polyethylene vials (22 mm × 8 mm × 1.5 mm thick; Just Plastics, London, UK).

A blend of equal weights of 1-octanol and octyl hexanoate was formulated as the neat material (25 μl) or a 10% solution in sunflower oil (100 μl) and applied to cellulose acetate cigarette filters (14 mm × 6 mm; Swan, High Wycombe, Buckinghamshire, UK) in the dispensers. Two vials for each type of dispenser and loading were maintained in a wind tunnel (27 °C and 8 km/h windspeed) and release rates were measured by periodic trapping of volatiles on Porapak followed by quantitative GC-FID analysis using decyl acetate (5 μg) as internal standard, as described for collection of pheromone from insects.

Field Trials in Commercial Glasshouses

Two field trapping experiments were carried out on tomato in commercial glasshouses in Kent, UK. Varieties included Piccolo, Summer Sun, and Sun Stream. Temperature and other meteorological data were recorded every five minutes; during the period of the trapping experiments the mean temperature was 17.6 °C, with maximum 25.5 °C and minimum 13.0 °C. Traps were yellow “Drystick” (dry glue) sticky traps (25 cm × 10 cm; Koppert, Ashford, UK) with the lure attached to the center by a twist-tie. In preliminary tests, white sticky traps were used with standard polybutene glue, but laboratory and field studies indicated the bugs could escape from this.

In the first trial (19 December 2019–09 January 2020) there were 10 blocks of three treatments, with one replicate of each treatment per block. Treatments formulated on cigarette filters in polyethylene vials were (A) a 1:1 blend of neat 1-octanol and octyl hexanoate (25 μl), (B) neat octyl hexanoate (12.5 μl), (C) blank control of a cigarette filter only. In eight blocks traps were positioned at 6 m above ground, just above the crop canopy, and in the other two blocks traps were positioned at 1 m, within the crop canopy. Traps were spaced >10 m apart, with blocks in a continuous row, to minimize interference between treatments. Trap catches were recorded weekly on 27 December 2019, 3 January, and 9 January 2020. However, the glasshouse was treated with insecticide (Pyrethrum 5EC) on 4 January 2020 because of the high catches of N. tenuis. Catches were low in the following week and only data from the first two weeks were used in analyses.

In the second experiment (10 January – 30 January 2020), the three treatments were (A) and (C) as above, and treatment (D) containing a 1:1 blend of 1-octanol and octyl hexanoate as a 10% solution in sunflower oil, resulting in an approximately 10-fold reduction in release rate relative to that from (A). There were 15 blocks with one replicate of each treatment per block. Eight blocks were positioned immediately above the crop canopy, 4 blocks 1 m above the crop canopy, and 3 blocks just above the base of the plants. Catches were recorded on 18, 25 and 30 January 2020.

In all tests the same sticky trap was used throughout each experiment. N. tenuis captured were counted and sexed on the trap and circled with a marker pen to differentiate them from new catches the following week. Sticky traps were collected from the glasshouse at the end of experiment and sexing was confirmed under a microscope by dissecting open the posterior abdomen and identifying the paramere in the males and ovipositor in the females (Kim et al. 2016).

For each experiment, catches of males in each trap were summed over the course of the experiment and transformed to ln(x + 1) to normalize variance. Trap catch was entered as the dependent variable in a mixed model, with treatment and trap height as fixed factors, and block entered as a random factor (Bates et al. 2015). A treatment * trap height interaction was included to test whether numbers of insects caught by each treatment varied with height. Statistical significance of fixed factors was determined through χ² tests of changes in residual deviance following deletion from the model (Crawley 2013). Post-hoc comparisons between catches made by different treatments and at different traps were made using Tukey’s tests on estimated marginal means (Lenth 2019). All analyses were performed in R Version 3.3.3.

Results

Pheromone Identification

Initial GC-EAG analyses of volatile collections from mated female N. tenuis showed two peaks (A and B) eliciting consistent EAG responses from antennae of mated male N. tenuis. The same result was obtained in analyses of volatiles from a single unmated female with antennae of unmated males (Fig. 1).
In analyses of these collections by GC-MS, the two EAG-active components (A) and (B) were present in volatiles from both mated and unmated female \textit{N. tenuis} and in volatiles from both mated and unmated males, but not detectable in volatiles from green beans used as food source during the volatile collections (Fig. 2). The mass spectra indicated these two peaks corresponded to 1-octanol and octyl hexanoate, respectively (structures (I) and (II) in Fig. 3), and the corresponding synthetic compounds had identical mass spectra and retention times on both polar and non-polar GC columns (Table 1).

Detailed examination of the GC-MS traces failed to detect the cocoa mirid pheromone components, hexyl (R)-3-hydroxybutyrate (IV in Fig. 3) and hexyl (R)-3-[(E)-2-butenoyl]-butyrate (V), or the proposed pheromone component of \textit{M. pygmaeus}, hexyl (R)-3-acetoxybutyrate (VI) (Fig. 4).

The analogue of octyl hexanoate, hexyl octanoate (III) had identical retention times to octyl hexanoate on both polar and non-polar GC columns (Table 1), but clearly different mass spectrum. The mass spectrum of octyl hexanoate shows the anticipated diagnostic base peak at m/z 117, corresponding to protonated hexanoic acid (C$_5$H$_{11}$COOH$_2^+$), while that of hexyl octanoate shows a similarly diagnostic base peak at m/z 145 corresponding to protonated octanoic acid (C$_7$H$_{15}$COOH$_2^+$) (Supplementary Material, Fig. S1). In GC-EAG analyses, the analogue, hexyl octanoate, elicited a consistent EAG response from the antennae of male \textit{N. tenuis} (Supplementary Material, Fig. S2).

Quantification of Pheromone Components

Of the 31 collections of volatiles made from \textit{N. tenuis} adults, only two collections – one virgin female and one virgin male – contained no detectable pheromone components. The rest contained detectable amounts of both pheromone component.
(A), 1-octanol (I), and pheromone component (B), octyl hexanoate (II), as shown in Table 2.

In collections from groups of 5–20 insects, the ratio of octyl hexanoate/octanol was higher in those from females than in those from males for both mated (0.6 ± 0.2 SE and 0.1 ± 0.01 respectively), and unmated (0.9 and 0.1 respectively) insects. For single, unmated insects, the ratios were the same (1.1) although more of

![Fig. 2 GC-MS analyses on polar GC column of volatile collections from green bean (upper), single virgin male *Nesidiocoris tenuis* (middle), and single virgin female *N. tenuis*, showing peaks A and B corresponding to EAG responses](image)

![Fig. 3 Structures of pheromone components and analogs: (I) 1-octanol (component A); (II) octyl hexanoate (component B); (III) hexyl octanoate; (IV) hexyl (R)-3-hydroxybutyrate; (V) hexyl (R)-3-[(E)-2-butenoyl]-butyrate; (VI) hexyl (R)-3-acetoxybutyrate](image)
both components was produced by females than males (Table 2).

Unexpectedly, whole body extracts of mated and unmated males and females contained only octyl hexanoate, and 1-octanol could not be detected (<0.01%). For both females and males, amounts in unmated insects seemed to be higher than in mated, although only two unmated insects were extracted (Table 2).

### Table 1 GC Retention Indices of compounds on polar and non-polar GC columns (relative to the retention times of n-alkanes)

| Compound                                           | Polar GC-EAG | Polar GC-MS | Non-polar GC-MS |
|----------------------------------------------------|--------------|-------------|-----------------|
| EAG-active compound (A)                            | 1557         | 1566        | 1074            |
| EAG Active compound (B)                            | 1815         | 1820        | 1584            |
| 1-octanol (I)                                      | 1557         | 1566        | 1074            |
| octyl hexanoate (II)                               | 1815         | 1820        | 1584            |
| hexyl octanoate (III)                              | 1815         | 1820        | 1584            |
| hexyl (R)-3-hydroxybutyrate (IV);                  | 1912         | 1921        | 1338            |
| hexyl (R)-3-[(E)-2-butenoyl]-butyrate (V)          | 2232         | 2242        | 1709            |
| hexyl (R)-3-acetoxy-butyrate (VI)                  | 1942         | 1957        | 1487            |

Fig. 4 GC-EAG analyses of synthetic compounds (20 ng injected) with male (upper) or female (lower) *Nesidiocoris tenuis* antennal EAG preparation on polar GC column showing EAG responses to 1-octanol (I, component A) at 7.43 min and octyl hexanoate (II, compound B) at 8.93 min and also possibly at to hexyl (R)-3-acetoxybutyrate (VI) at 9.60 min but not to hexyl (R)-3-hydroxybutyrate (IV) at 9.44 min; or hexyl (R)-3-[(E)-2-butenoyl]-butyrate (V) at 10.97 min.
Table 2  Amounts and relative ratio of pheromone components in collections of volatiles and whole-body extracts from Nesidiocoris tenuis (analysis GC-FID on non-polar GC column against external standard for volatiles, internal standard 5 μg decyl acetate for extracts)

| Source                          | Component A octanol (I) | Component B octyl hexanoate (II) | Ratio octyl hexanoate/octanol |
|---------------------------------|-------------------------|----------------------------------|------------------------------|
| Volatile Collections (ng/h/insect) |                         |                                  |                              |
| 5–20 mated female (N=5)         | 1.3 (0.2)               | 0.8 (0.4)                        | 0.6 (0.2)                    |
| 5–20 mated male (N=5)           | 3.7 (0.5)               | 0.3 (0.04)                       | 0.1 (0.01)                   |
| 20 virgin female (N=2)          | 4.6 (2.3–6.9)           | 3.3 (2.7–3.9)                    | 0.9 (0.6–1.2)                |
| 20 virgin male (N=2)            | 3.7 (3.0–4.4)           | 0.4 (0.4–0.4)                    | 0.1 (0.1–0.1)                |
| single virgin female (N=8)      | 2.9 (1.1)               | 2.5 (0.8)                        | 1.1 (0.3)                    |
| single virgin male (N=5)        | 0.7 (0.1)               | 0.6 (0.2)                        | 1.1 (0.6)                    |
| Extracts (ng/insect)            |                         |                                  |                              |
| virgin female (N=2)             |                        | 1826 (1292–2360)                |                              |
| virgin male (N=2)               |                        | 1712 (993–2432)                 |                              |
| mated female (N=6)              |                        | 750 (197)                        |                              |
| mated male (N=6)                |                        | 821 (215)                        |                              |

Controlled Release Dispensers

Pipette tips and sealed polyethylene vials were investigated as controlled release dispensers for the synthetic pheromone components, using a 1:1 blend of 1-octanol and octyl hexanoate either neat or as a 10% solution in sunflower oil. Release of octyl hexanoate from the pipette tips was negligible and these were not investigated further. Polyethylene vials released both components for over six weeks at 27 °C and 8 km/h windspeed (Fig. 5). The release rates for the neat material were approximately ten times those for the solution in sunflower oil. The ratio of octyl hexanoate/ octanol during the first month of exposure was approximately 0.8 for the neat material and 0.5 for the sunflower solution.

Field Trapping

Two trapping tests were carried out in commercial greenhouses. In the first trial, a significant effect of treatment was found on numbers of male N. tenuis caught ($\chi^2 = 21.8$, df=2, $P < 0.001$). Traps baited with the two-component blend of 1-octanol and octyl hexanoate (Treatment A) caught significantly more male N. tenuis than traps baited with octyl hexanoate alone (Treatment B, Tukey’s test, $P < 0.05$). Treatments A and B caught more male N. tenuis than the blank control (Treatment C, Tukey’s test, $P < 0.05$) (Fig. 6). No evidence was found for an effect of trap height (6 m or 1 m above ground; $\chi^2 = 0.14$, df=1, $P = 0.71$) or an interaction between trap height and treatment ($\chi^2 = 2.7$, df=2, $P = 0.26$) on trap catches. Six female N. tenuis were captured across the experiment (Treatment A: two females, B: two females, C: two females). In this experiment, the glasshouse was treated with insecticide after two weeks and the total catch of male N. tenuis dropped from 56 and 58 in the first two weeks, respectively, to 11 in the third week. Only catches from the first two weeks were used in the analyses.

In the second experiment, significant effects of both treatment ($\chi^2 = 33.0$, df=2, $P < 0.001$) and trap height ($\chi^2 = 12.3$, df=2, $P < 0.01$) were found on catches of male N. tenuis. However, there was no evidence of a Treatment * Trap height interaction on catches ($\chi^2 = 2.5$, df=4, $P = 0.65$). Averaged over trap height, Treatments A and D caught significantly more male N. tenuis than control treatment C (Tukey’s post hoc test on estimated marginal means, $P < 0.05$), but lures with the higher release rate (A) did not catch significantly more bugs than those with a ten-fold lower release rate (D) (Fig. 7). Averaged over treatments, significantly more male N. tenuis were caught at the top of the plant compared to 1 m above or at the plant base (Tukey’s test, $P < 0.05$) (Fig. 8). Three female N. tenuis were caught in traps baited with treatment A, three with treatment C and seven with treatment D. Eight of the females were caught on traps positioned at the tops of the plants, three were caught 1 m above the plant, and 2 were caught at the plant base.

Discussion

Two components of the sex pheromone produced by adult female N. tenuis have been identified as 1-octanol and octyl hexanoate. In trapping tests carried out in commercial tomato glasshouses, traps baited with octyl hexanoate alone caught significantly more male N. tenuis than unbaited traps, but...
significantly less than traps baited with a 1:1 blend of the two components.

**Pheromone Identification**

Octyl hexanoate was reported as a minor component (5–7% of major component, hexyl hexanoate) in volatiles produced by females of another mirid bug, *Trigonotylus caelestialium* Kirkaldy, but was not important for attraction of males (Kakizaki and Sugie 2001). It is also reported to be involved in the attraction of females of *N. tenuis*.
in the chemical ecology of various stingless bees, e.g. of the genus *Trigona* (Hymenoptera; Apidae). For example, it is a component of a trail pheromone and labial gland secretion (Jarau et al. 2010), of a recruitment pheromone (Lichtenberg et al. 2011), and of the cephalic secretion (Francke et al. 2000). The isomer, hexyl octanoate has also been reported to be present in these stingless bee secretions, but 1-octanol has not previously been reported as an insect semiochemical. This compound was shown to be produced by female *Macrolophus pygmaeus*, but its role was not established (Gemeno et al. 2006).

*Nesidiocoris tenuis* belongs to the Bryocorinae sub-family of mirids, and female-produced sex pheromones have been identified in several species of this sub-family. These include the cocoa mirids, *Distantiella theobroma* (Dist.) and *Sahlbergella singularis* Haglund, (Mahob et al. 2011; Mahot et al. 2020; Sarfo 2013; Sarfo et al. 2018a, 2018b), and the aphidophagous mirid, *M. pygmaeus* (Gemeno et al. 2006). The components of these pheromones are derivatives of esters of 3-hydroxybutyric acid, and rather different from those identified in mirid species from other subfamilies which are typically saturated or unsaturated, straight-chain esters or aldehydes (Zhang et al. 2020 and refs therein). The pheromone components identified in *N. tenuis*, 1-octanol and octyl hexanoate, are more like the latter than the former, and the cocoa mirid and *M. pygmaeus* pheromone components could not be detected in volatiles or extracts from female or male *N. tenuis*. Furthermore, the synthetic compounds did not elicit a consistent EAG response from antennae of male *N. tenuis*.

It was also unexpected that both pheromone components are released by both mated and unmated male and female *N. tenuis* when in groups or as isolated individuals. For many species of mirid, the same compounds typically serve as pheromone components and also as defensive compounds (Aldrich 1988; Fountain et al. 2014; Staddon 1986). Thus, in four *Lygus* species, Fountain et al. (2014) showed that three compounds could be extracted from both females and males, and when volatiles were collected from groups of insects, all three compounds were produced in a similar ratio from females and males. However, when volatiles were collected from individual, undisturbed virgin insects, the three compounds were only produced by females and in a species-specific ratio that constituted the sex pheromone and was different from that produced by groups of insects. Collections from *N. tenuis* were rather different. In those from insects in groups, the ratio of octyl hexanoate/1-octanol was higher from females than males whether mated (0.6 and 0.1 respectively) or unmated (0.9 and 0.1 respectively). In collections from individual insects the ratios from females and males were essentially the same (1.1), although results from individuals were very variable at 0.4–3.0 in females and 0.2–3.4 in males.

Even more surprising was the observation that, although the two compounds were found in volatiles from mated and unmated *N. tenuis* groups and individuals of males or females, whole body extracts in diethyl ether of mated and unmated males or females contained only octyl hexanoate with no 1-octanol detectable. This result is difficult to explain unless the 1-octanol is contained in some gland not extracted by the solvent. Significant crushing of the bodies was avoided in order not to extract extraneous body materials that could damage the GC column used for analysis, but light compression of the bodies failed to change the result. The same technique has been used on several other mirid species and gave extraction of all pheromone components. One possible factor is that the extracts here were all made at a similar time in the diurnal cycle. Volatile collections were made for complete 24 h or 48 h periods, and analyses of whole body extracts made at different times could provide different results.

It is also possible that production of 1-octanol was induced by herbivore feeding on the green beans used as food for the bugs during collections of volatiles. However, in our studies, no 1-octanol was detected in collections from intact beans in the absence of insects, or from beans punctured with a dissection needle. Both 1-octanol and octyl hexanoate were obtained in initial collections from bugs on aubergines in similar absolute and relative amounts to those obtained in collections from bugs on green beans. Induction of production of volatiles by *N. tenuis* has been reported for tomatoes (Pérez-Hedo et al. 2018) and sweet peppers (Bouagga et al. 2018), but 1-octanol was not reported in these studies. Both 1-octanol and octyl acetate were detected in whole body extracts of female *M. pygmaeus*, along with proposed pheromone components hexyl and octyl 3-acetoxybutanoate (Gemeno et al. 2006). In collections of volatiles from *M. pygmaeus* with or without green beans, octyl acetate was the major component with much smaller amounts of the other compounds detected (Gemeno et al. 2006). Thus it is concluded that 1-octanol is most likely produced by *N. tenuis* adults.

**Field Trapping**

In the initial studies of volatiles released by mated insects, the amount of 1-octanol was relatively higher than that of octyl hexanoate. Nevertheless, it was thought that 1-octanol was possibly a biosynthetic precursor and that octyl hexanoate was the “major” pheromone component. Thus, in the first trapping experiment, because of the limited resources available, only lures containing the two-component blend or octyl hexanoate alone were tested. Traps baited with the latter did catch more male *N. tenuis* than unbaited traps, but those baited with the two-component blend caught significantly more. It was not possible to test the 1-octanol alone during the short timeframe of this project, but obviously this should be done in future work.

Polyethylene vials provided convenient dispensing systems for the two pheromone components. The release rate could be adjusted by diluting the components in an involatile
with damage thresholds in crops. Given the ready availability of the pheromone, the possibilities for using it in control of *N. tenuis* through mass trapping or mating disruption could be investigated, as well as the potential of the pheromone isomer, hexyl octanoate, as an inhibitor or repellent.

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Availability of Data and Material Raw chromatographic and trapping data are available on request from the authors.

Code Availability N/A

Authors’ Contributions DRH managed the chemical and analytical work, carried out chemical synthesis and was lead author writing the paper; SJH carried out chemical and analytical work; DPB carried out EAG studies, and designed and analysed the trapping studies; DIF assisted in pheromone collection and carried out analysis and release rate studies, RJ organised and managed trapping experiments; CXS reared the insects and made the pheromone collections; MTF managed the project and assisted with insect rearing, pheromone collection and trapping experiments. All authors contributed to writing the paper.

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Declarations

Ethics Approval Not required.

Consent to Participate Not required.

Consent for Publication All authors agreed to submission of the final manuscript.

Conflicts of Interest/Competing Interests None

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