

Sulfur volatiles from *Allium* spp. affect Asian citrus psyllid, *Diaphorina citri* Kuwayama (Hemiptera: Psyllidae), response to citrus volatiles

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Abstract

The Asian citrus psyllid, *Diaphorina citri* Kuwayama, vectors *Candidatus Liberibacter asiaticus* (Las) and *Candidatus Liberibacter americanus* (Lam), the presumed causal agents of huanglongbing. *D. citri* generally rely on olfaction and vision for detection of host cues. Plant volatiles from *Allium* spp. (Alliaceae) are known to repel several arthropod species. We examined the effect of garlic chive (*A. tuberosum* Rottl.) and wild onion (*A. canadense* L.) volatiles on *D. citri* behaviour in a two-port divided T-olfactometer. Citrus leaf volatiles attracted significantly more *D. citri* adults than clean air. Volatiles from crushed garlic chive leaves, garlic chive essential oil, garlic chive plants, wild onion plants and crushed wild onion leaves all repelled *D. citri* adults when compared with clean air, with the first two being significantly more repellent than the others. However, when tested with citrus volatiles, only crushed garlic chive leaves and garlic chive essential oil were repellent, and crushed wild onions leaves were not.

Analysis of the headspace components of crushed garlic chive leaves and garlic chive essential oil by gas chromatography-mass spectrometry revealed that monosulfides, disulfides and trisulfides were the primary sulfur volatiles present. In general, trisulfides (dimethyl trisulfide) inhibited the response of *D. citri* to citrus volatiles more than disulfides (dimethyl disulfide, allyl methyl disulfide, allyl disulfide). Monosulfides did not affect the behaviour of *D. citri* adults. A blend of dimethyl trisulfide and dimethyl disulfide in 1:1 ratio showed an additive effect on inhibition of *D. citri* response to citrus volatiles. The plant volatiles from *Allium* spp. did not affect the behaviour of the *D. citri* ecto-parasitoid *Tamarixia radiata* (Waterston). Thus, *Allium* spp. or the tri- and di-sulphides could be integrated into management programmes for *D. citri* without affecting natural enemies.

Keywords: *Diaphorina citri*, *Tamarixia radiata*, garlic chive, dimethyl trisulfide, dimethyl disulfide, citrus

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Introduction

Asian citrus psyllid, *Diaphorina citri* Kuwayama (Hemiptera: Psyllidae), is one of the most serious pests of citrus worldwide because it vectors the bacterial pathogen causing huanglongbing (HLB) disease. In addition, both nymphs and adults are obligate phloem feeders that cause chlorosis on infested leaves and excrete honeydew that promotes the growth of sooty mold. HLB affects plant phloem, causing yellow shoots, mottling, chlorosis and twig die back, which cause rapid tree decline and may ultimately cause tree death. Fruit on diseased trees do not colour properly and can be bitter tasting and misshapen as well as reduced in size (Capoor, 1963; Halbert & Manjunath, 2004; Bové, 2006). Where HLB is present, infected citrus trees decline and may die within a few years and never produce usable fruit (Halbert & Manjunath, 2004). The disease is presumed to be caused by either of three species of phloem-limited, noncultured, Gram-negative bacteria: *Candidatus Liberibacter asiaticus* (Las), *Candidatus Liberibacter africanus* (Laf) and *Candidatus Liberibacter americanus* (Lam) (Bové, 2006). The bacteria are efficiently vectored by two species of citrus psyllid (*D. citri* and *Trioza erytreae* (del Guercio)).

Direct damage caused by psyllid feeding is restricted to young and tender leaves; therefore, control measures were previously only justified for non-bearing citrus which often produce shoot flushes throughout the growing season. The discovery of HLB disease in Florida in 2005 changed the status of *D. citri* to a major pest requiring management in all stages of citrus production (Hodges & Morse, 2009). Currently, a broad range of vector and disease control methods is being employed to control this disease in Florida (Qureshi & Stansly, 2007; Rogers & Timmer, 2007). Current management practices are incomplete because of a lack of known resistant cultivars (Halbert & Manjunath, 2004), effective biological control agents (Qureshi & Stansly, 2007) and cultural control options (Childers & Rogers, 2005; Powell *et al.*, 2007). Classical biological control of *D. citri* with the introduced parasitoid, *Tamarixia radiata* (Waterston), was initiated in Florida in 1999 (Hoy & Nguyen, 2001). The parasitoid established and dispersed quickly, but parasitism rates are generally low and variable (Qureshi *et al.*, 2009). Primary control of the vector has relied on broad spectrum insecticides (Rogers, 2008). However, insecticide use is known to negatively affect populations of natural enemies and may lead to development of insecticide resistance. Therefore, novel management strategies, such as insect repellents, attractants and antifeedents, may serve as useful alternatives or supplements to insecticides.

All known host plants of *D. citri* belong to the family Rutaceae and comprise a wide range of species including ten from genera other than *Citrus* (Aubert, 1990). *D. citri* generally rely on olfaction and vision for detection of host cues (Moran & Brown, 1973; Kristoffersen *et al.*, 2006; Onagbola *et al.*, 2009; Wenninger *et al.*, 2009). Interplanting guava, *Psidium guajava* (L.), has been reported to reduce *D. citri* populations in citrus orchards in Vietnam, possibly due to volatile chemicals (Beattie *et al.*, 2006; Hall *et al.*, 2008; Zaka *et al.*, 2010). Recently, Rouseff *et al.* (2008) identified dimethyl disulfide (DMS) from guava as a possible defensive volatile that may explain guava's repellency to *D. citri*. Plants in the family Alliaceae also have been shown to produce sulfur compounds known to repel arthropods (Auger *et al.*, 1989; Dugravot *et al.*, 2002, 2003, 2004,

2005; Amarawardana *et al.*, 2007). Chemicals from *Allium sativum* (L.) have been reported to repel cotton bollworms (Gurusubramanian & Krishna, 1996), codling moth (Landolt *et al.*, 1999), aphids (Assis *et al.*, 2007), whiteflies (Salas, 2001), cabbage root flies (Prowse *et al.*, 2006), mosquitoes (Snow & Cutler, 2006) and phlebotomine sand flies (Valerio & Maroli, 2005). Furthermore, garlic chive, *A. tuberosum*, has been shown to repel green peach aphid, *Myzus persicae* (Sulzer), in sweet pepper (Amarawardana *et al.*, 2007). In addition, the leek, *A. porrum* L., has been reported to produce sulfur-based chemicals in response to *Acrolepiopsis assectella* (Zeller) attack (Dugravot *et al.*, 2002, 2005). Based on the repellent properties of these plants, certain *Allium* spp. products such as garlic barrier[®] (Garlic Research Labs, CA), Garlic shield[®] (Nutritek Inc., OR) and Biorepel[®] (JH Biotech Inc., CA) have been commercially formulated for pest management. In this series of experiments, we examined the effect of *Allium* spp. as well as sulfur volatiles identified from them on the behaviour of *D. citri* and its parasitoid, *T. radiata*. Our objective is to identify possible effective repellents for *D. citri* that have minimal effects on its parasitoid.

Methods and materials

Insects

Adult *D. citri* used in behavioural bioassays were obtained from a laboratory culture at the University of Florida Citrus Research and Education Center (Lake Alfred, USA). The culture was established in 2000 from field populations in Polk Co., FL, USA (28.0°N, 81.9°W) prior to the discovery of HLB in FL. The culture is maintained without exposure to insecticides on sour orange (*Citrus aurantium* L.) and 'Hamlin' orange (*C. sinensis* (L.) Osb.) seedlings at 27 ± 1°C, 63 ± 2% RH and under a L14:D10 photoperiod. The sexual maturity of psyllids was confirmed by placing adults for at least seven days on citrus seedlings in Plexiglass[®] cages prior to use in experiments (Wenninger & Hall, 2007). *Tamarixia radiata* adults were collected from newly expanded 'Hamlin' orange leaf flush infested with *D. citri* from mature trees in an unsprayed grove in Polk Co., FL. The flush was maintained in 40 × 40 × 40 cm Plexiglas cages at 26 ± 1°C, 60 ± 5% RH and under a photoperiod of 14:10 (L:D) h. Parasitoids emerging from psyllid nymphs were collected daily and maintained as described in Onagbola *et al.* (2009).

Plant material and chemical samples

Garlic chive (*A. tuberosum*), wild onion (*A. canadense*), white guava (*P. guajava*) and Valencia orange (*C. sinensis*) plants were grown in 1.0 gallon pots in a temperature controlled green house. *Allium* spp. samples were composed of 10-week-old whole plants. Since fresh leaves are known to contain higher proportions of secondary compounds (Hrutfjord *et al.*, 1974) and *D. citri* are exclusively associated with new growth (Catling, 1970), we harvested approximately 2.0 g of fresh leaf flush (immature leaves of the growing shoots (Hall & Albrigo, 2007)) for *D. citri* behavioural assays. Dimethyl disulfide (DMS) (≥98% purity), dimethyl trisulfide (DMTS) (≥98.5% purity), allyl methyl sulfide (AMS) (≥98% purity) and Chinese garlic chive essential oil (unknown purity) were obtained from Sigma-Aldrich Inc. USA. Allyl methyl disulfide (AMDS) (≥98%

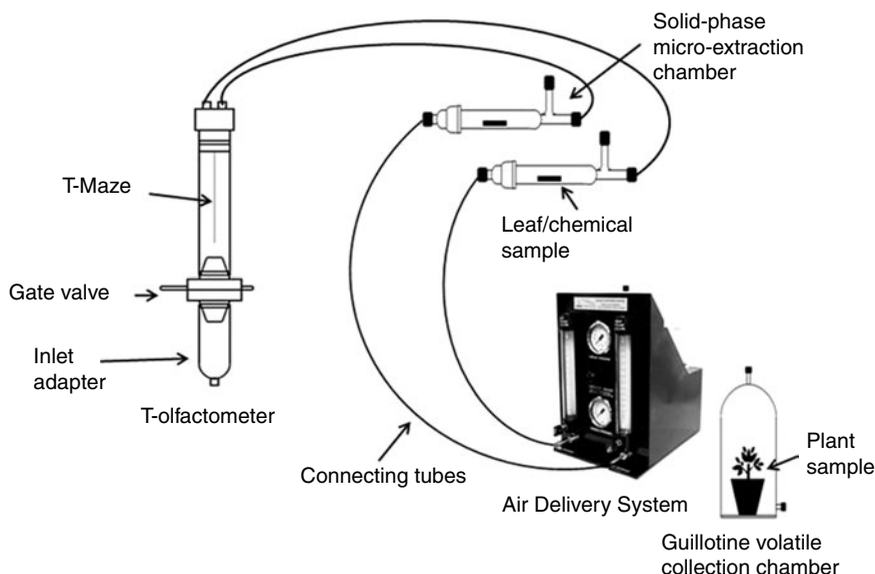


Fig. 1. Schematic diagram of the T-maze olfactometer used for behavioural bioassays. The guillotine volatile collection chambers were replaced with the solid-phase micro-extraction chambers for leaf samples.

purity) and allyl disulfide (ADS) (~80.0% purity) were obtained from Frutarom (UK) Ltd, UK and Penta Chemical Company, USA, respectively.

Behavioural bioassays with plant material

A custom designed two-port divided T-olfactometer (fig. 1) from Analytical Research Systems (ARS) Inc., Gainesville, FL was used to evaluate behavioural response of *D. citri* to *Allium* spp. plants. The olfactometer consisted of a 30 cm long glass tube with 3.5 cm internal diameter that is bifurcated into two equal halves with a Teflon strip forming a T-maze. Each half served as an arm of the olfactometer, enabling the *D. citri* to make a choice between two potential odour fields. The olfactometer arms were connected to odour sources placed in guillotine volatile collection chambers or solid-phase micro-extraction chambers (ARS, Gainesville, FL) through Teflon[®] glass tube connectors (fig. 1). The plant samples were placed directly in 51 guillotine volatile collection chambers (GVCC) (fig. 1). For plant samples, a 35 cm tall × 15 cm wide dome shaped GVCC was placed directly over the pots. The GVCC was supported with a Teflon guillotine support base (GSB) (ARS, Gainesville, FL) (fig. 1). The GVCC was attached with an inlet valve on the side of a chamber and an outlet valve on top of the dome for incoming and outgoing air streams, respectively. The leaf samples were placed in solid-phase micro-extraction chambers (SPMEC) (ARS, Gainesville, FL). The SPMEC consists of a straight glass tube (17.5 cm long × 2.5 cm internal diameter) supported with an inlet and outlet valve for incoming and outgoing air streams, respectively. The leaf samples were wrapped in laboratory tissue (Kimwipes, Kimberly-Clark, Roswell, GA) to minimize contamination of the SPMEC. The purified and humidified air was pushed through the GVCC or SPMEC via two pumps connected to an air delivery system (ARS, Gainesville, FL) (fig. 1). A constant airflow of 0.11 min^{-1} was maintained through both arms of the olfactometer. Purified and humidified air was pushed

through these chambers via two pumps connected to an air delivery system (ARS, Gainesville, FL) (fig. 1). The olfactometer was housed within a temperature controlled room and positioned vertically under a fluorescent 900 lux light bulb positioned within a $1.0 \times 0.6 \times 0.6 \text{ m}$ fibre board box for uniform light diffusion. This position took advantage of the negative geotactic and positive phototactic response of *D. citri* (Mann *et al.*, unpublished results). The olfactometer inlet adapter was covered with black cloth to facilitate insect movement towards odour sources. An odour source was randomly assigned to one of the arms of the olfactometer at the beginning of each bioassay and was reversed after every 30 insects to eliminate positional bias.

Female *D. citri* are more responsive to citrus odours than males (Wenninger *et al.*, 2009); therefore, only female adults were tested in this study. Prior to tests involving putative repellents, *D. citri* female adults were exposed to clean air vs. clean air and citrus vs. citrus in the T-maze olfactometer to verify the absence of positional bias. Thereafter, the following odour source combinations were tested: (i) garlic chive plant vs. clean air; (ii) crushed garlic chive leaves vs. clean air; (iii) wild onion plant vs. clean air; (iv) crushed wild onion leaves vs. clean air; (v) garlic chive essential oil vs. clean air; (vi) citrus leaves vs. clean air; and (vii) clean air vs. clean air (control). The putative repellents were also evaluated in combination with citrus odours to verify their repellent effect in presence of citrus odours. The treatment tested for this set of experiments were: (i) crushed garlic chive leaves + citrus leaves vs. citrus leaves; (ii) crushed wild onion leaves + citrus leaves vs. citrus leaves; (iii) garlic chive essential oil + citrus leaves vs. citrus leaves; (iv) clean air vs. clean air (negative control); and (v) citrus leaves vs. citrus leaves (positive control). The plant samples were composed of 10-week-old whole plants, while leaf samples were composed of ~2.0 g of fresh leaves obtained from *Allium* or citrus plants. A minimum of 120 *D. citri* female adults were examined per treatment combination (four replications of 30 *D. citri* per replication). *D. citri* females were released

individually into the inlet adapter at the base of the olfactometer. Adults were given 300s to exhibit a behavioural response by entering either olfactometer arm. The number of adults entering the treatment arm, control arm or remaining in the inlet adapter (release port) or below T-maze division was recorded. A treatment or control arm choice was recorded when an insect moved into either olfactometer arm by crossing the division in the T-maze olfactometer. A release arm choice was recorded when an insect remained in the release port or below the T-maze division. All experiments were conducted at $26 \pm 1^\circ\text{C}$ and $60 \pm 2\%$ RH. The olfactometer and connecting tubes were thoroughly cleaned with 2% soap solution and baked at 93.3°C between each treatment run. The response of *T. radiata* to *Allium* spp. volatiles was evaluated following the procedures described for *D. citri*. However, for this insect only 60 females were tested per treatment combination given that fewer parasitoids were available.

Chemical analysis

D. citri response to citrus volatiles was inhibited by volatiles from crushed garlic chive essential oil (see results); therefore, the head-space volatiles from these treatments were identified using a static solid phase micro extraction (SPME) technique described in Rouseff *et al.* (2008). Briefly, a sample was weighed into a 40 ml septum-sealed glass vial. The sample was allowed to equilibrate at ambient laboratory conditions for ~30 min. Accumulated volatiles were collected from the glass vials for 1 min post exposure to equilibrate the samples to ambient laboratory conditions (20–25°C). The volatiles were collected using a 75 µm carboxen-polydimethylsiloxane (PDMS) Stable Flex® SPME fiber (Supelco, Bellefonte, PA). At least three replicates of each volatile sample were analyzed. The collected volatiles were analyzed and identified using a Perkin/Elmer® Clarus 500 quadrupole mass spectrometer (GC-MS). The GC-MS was equipped with Turbo Mass software (Perkin/Elmer, Shelton, CT) and a 60 m × 0.25 mm, i.d. × 0.50 µm Restek (RTX-5) capillary column. Helium was used as the carrier gas at a constant flow of 2 ml min⁻¹. The source was kept at 200°C and the transfer line and injector were maintained at 260°C. The oven was programmed from 40 to 260°C at 7°C min⁻¹. We matched mass spectra with NIST 2005 version 2.0 standard spectra (NIST, Gaithersburg, MD); compounds with spectral fit values ≥800 and appropriate LRI values were considered positive identifications. Authentic standards were used to confirm identifications when available.

Behavioural bioassays with synthetic chemicals

The five sulfur volatiles that were found in greatest abundance in garlic chive essential oil and crushed leaves were tested for repellency to *D. citri*. The bioassay techniques used were similar to those described above. The chemicals were evaluated at 0.25, 0.5 and 1.0% concentrations individually and in combination with citrus leaves. The chemical samples were dissolved in 1 ml ethylene glycol (EG) and pipetted onto a 5 cm Richmond cotton wick (Petty John Packaging, Inc., Concord, NC). Given high volatility of sulfur compounds (Dugravot *et al.*, 2004), the synthetic chemicals were dissolved in EG to slow the release rate during bioassays (Arthur, 1996; Dugravot *et al.*, 2002). The

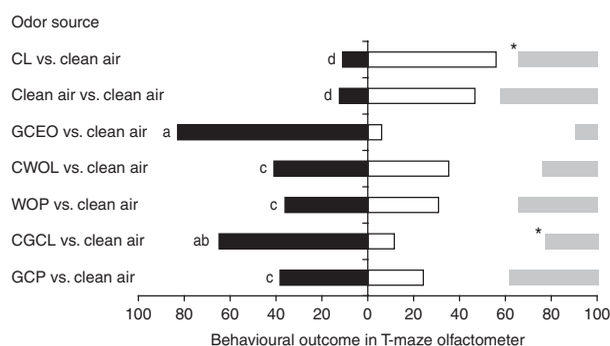


Fig. 2. Responses of *Diaphorina citri* when presented with volatiles emanating from *Allium* spp. plants. Garlic chive plant, GCP; crushed garlic chive leaves, CGCL; wild onion plant, WOP; crushed wild onion leaves, CWOL; garlic chive essential oil, GCEO; citrus leaves, CL. Black bars labelled with the same letters are not significantly different (Tukey's HSD TEST, $P < 0.05$). Grey bars labelled with * are significantly different from white bars (χ^2 test, $P < 0.05$) (■, release arm (%); □, treatment arm (%); ▤, control arm (%)).

treated cotton wick was wrapped in laboratory tissue and placed in solid-phase micro-extraction chambers. The control treatment contained cotton wicks impregnated with 1 ml EG only. For evaluations of chemicals in the presence of citrus odours, ~2.0 g of fresh citrus leaves were placed in both the chemical treatment and control arm of the olfactometer. The compound showing the highest activity against *D. citri* (dimethyl trisulfide) (DMTS) (see results) was also evaluated as a blend with DMDS at a 0.25% concentration. DMTS (0.25%) vs. clean air was used as the control treatment.

Data analysis

For assays in which putative repellent treatments were presented in the T-maze olfactometer with or without citrus and vs. clean air, the number of *D. citri* remaining at the release point and not entering the olfactometer was compared between treatments using one way analysis of variance (ANOVA) followed by Tukey's HSD test ($P < 0.05$). For psyllids leaving the release arm, the number of psyllids choosing the control arm vs. the treatment arm was compared with Chi square (χ^2) analysis at $P < 0.05$. The data from all four replicates were combined for the χ^2 analysis.

Results

Behavioural bioassays with plant material

When clean air was pumped into both arms of the T-maze, the majority of *D. citri* (89.2%) left the release arm, indicating a strong negative geotactic and positive phototactic response (fig. 2). Statistically equivalent numbers of *D. citri* entered each arm when clean air was presented vs. clean air, indicating no positional bias. Significantly ($F = 13.98$; $df = 6, 21$; $P < 0.0001$) more *D. citri* did not move from the release point in treatments in which garlic chive odours were co-presented with clean air compared with when citrus vs. clean air or clean air was presented alone (fig. 2). More *D. citri* remained at the release point when crushed garlic chive leaves or garlic chive essential oil was presented vs.

clean air than when a whole garlic chive plant vs. clean air, crushed wild onion leaves vs. clean air or a whole wild onion plant vs. clean air were presented (fig. 2). The percentage of *D. citri* not moving from the release point ranged between 65 and 82% when crushed garlic chive leaves or garlic chive essential oil was presented vs. clean air. When citrus volatiles were released vs. clean air, significantly ($\chi^2=8.98$; $df=1$; $P<0.0001$) more *D. citri* entered the arm receiving citrus odours than clean air (fig. 2). When garlic chive leaf volatiles were presented vs. clean air, significantly ($\chi^2=5.49$; $df=1$; $P=0.02$) fewer *D. citri* entered the arm with leaf volatiles than the arm receiving the clean air (fig. 2).

Significantly ($F=6.19$; $df=3,12$; $P<0.0087$) more *D. citri* did not move from the release point in treatments in which

crushed garlic chive leaves or garlic chive essential oil were co-presented with citrus leaves compared with when citrus leaves vs. citrus leaves were presented (fig. 3). However, the number of *D. citri* choosing each arm of the olfactometer were comparable when crushed garlic chive leaves + citrus leaves or garlic chive essential oil + citrus leaves were presented vs. citrus leaves (fig. 3). Volatiles from crushed garlic chive leaves and garlic chive essential oil did not affect the behaviour of *T. radiata* ($F=4.69$; $df=2,5$; $P=0.0713$) (fig. 4).

Chemical analysis

Allyl methyl trisulfide (AMTS), diallyl trisulfide (DATS), DMTS, ADS, AMDS, DMDS, diallyl sulfide (DAS) and AMS were identified from garlic chive essential oil (fig. 5). Five of these sulfur volatiles (AMS, ADS, AMDS, DMDS and DMTS) were also found in the head space of crushed garlic chive leaf samples (data not shown).

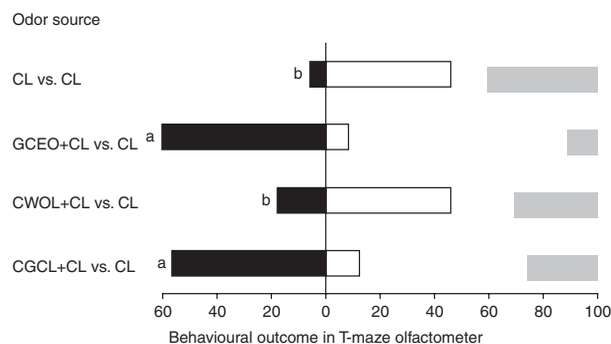


Fig. 3. Responses of *Diaphorina citri* when presented with volatiles emanating from *Allium* spp. plants with or without citrus odours. Crushed garlic chive leaves, CGCL; crushed wild onion leaves, CWOL; garlic chive essential oil, GCEO; citrus leaves, CL. Black bars labelled with the same letters are not significantly different (Tukey's HSD TEST, $P<0.05$) (■, release arm (%); □, treatment arm (%); ▤, control arm (%)).

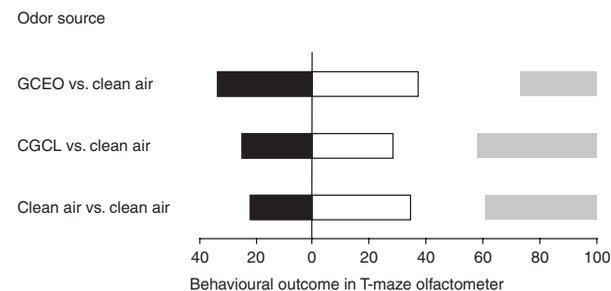


Fig. 4. Responses of *Tamarixia radiata* when presented with volatiles emanating from *Allium* spp. plants. Crushed garlic chive leaves, CGCL; crushed wild onion leaves, CWOL. (■, release arm (%); □, treatment arm (%); ▤, control arm (%)).

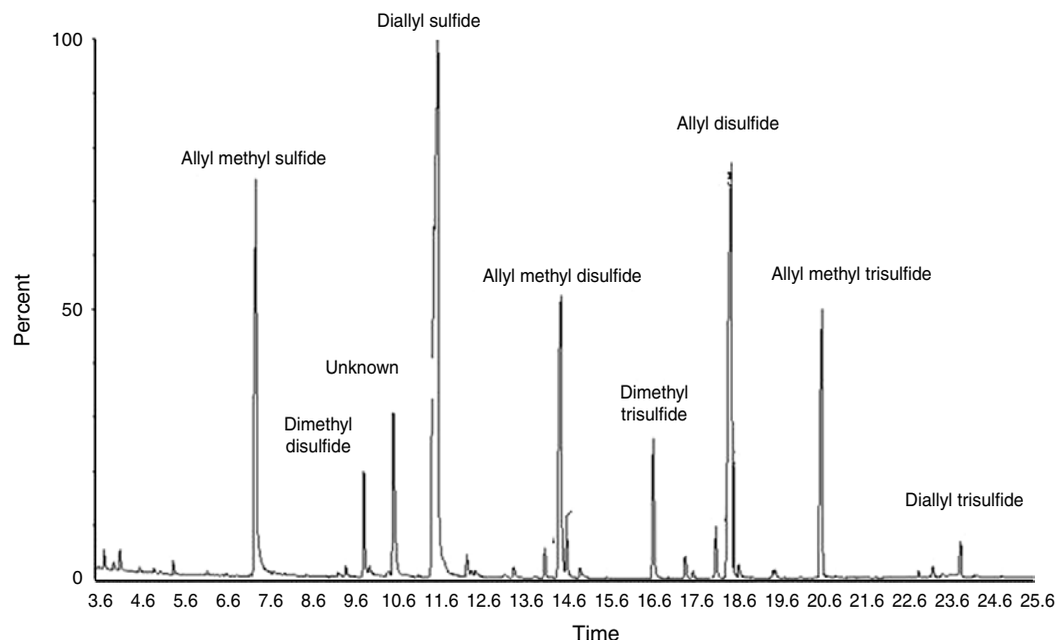


Fig. 5. Example chromatogram showing main sulfur volatiles identified from garlic chive essential oil.

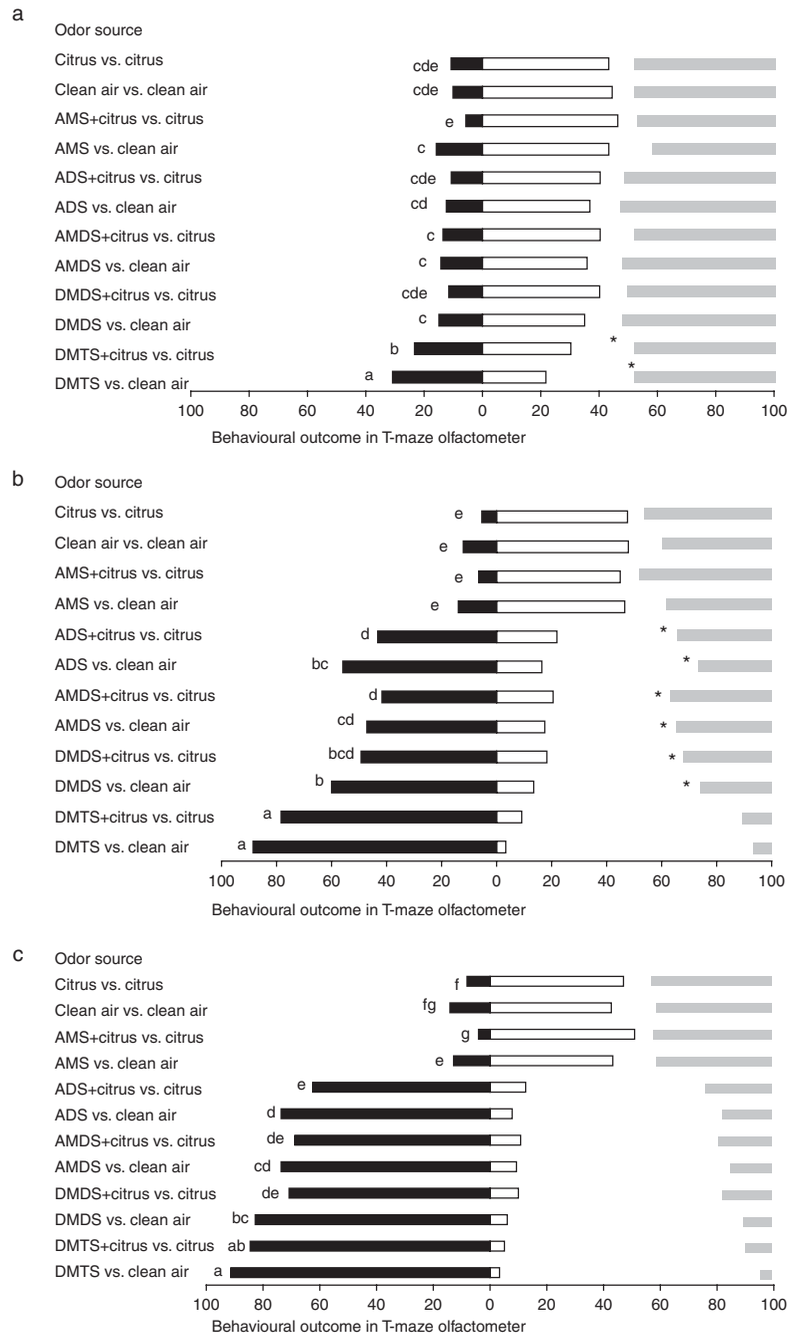


Fig. 6. Responses of *Diaphorina citri* when presented with sulfur volatiles at (a) 0.25%, (b) 0.5% or (c) 1.0% concentration with or without citrus odours. Allyl methyl sulfide, AMS; allyl disulfide, ADS; allyl methyl disulfide, AMDS; dimethyl disulfide, DMDS; dimethyl trisulfide, DMTS. Black bars labelled with the same letters are not significantly different (Tukey's HSD TEST, $P < 0.05$) (■, release arm (%); □, treatment arm (%); ▒, control arm (%)).

Behavioural bioassays with synthetic chemicals

Significantly ($F = 9.67$; $df = 11,36$; $P < 0.0001$) more *D. citri* did not move from the release point in treatments in which DMTS (at 0.25% concentration) was co-presented with clean air or citrus odours than when clean air or citrus were presented simultaneously in both arms of the olfactometer

(fig. 6a). No other synthetic sulfur chemical yielded significant differences from the control treatment (clean air vs. clean air) at the 0.25% concentration with respect to the number of *D. citri* leaving the release point (fig. 6a). Significantly ($\chi^2 = 11.58$; $df = 1$; $P = 0.0007$) more psyllids entered the control arm than the treatment arm when 0.25% DMTS was compared with clean air (fig. 6a).

Significantly ($F = 48.67$; $df = 11,36$; $P < 0.0001$) more *D. citri* did not move from the release point in treatments in which DMTS, DMDS, AMDS or ADS (all at 0.50%) was co-presented with clean air or citrus odours compared with when clean air alone or citrus odour was simultaneously presented in both arms of the olfactometer (fig. 6b). The percentages of psyllids not moving from the release point for DMTS vs. citrus and DMTS vs. clean air ranged between 79 to 89%, respectively, while the percentages of psyllids not moving from point of release for disulfides (DMDS, AMDS and ADS) ranged between 41 and 60% (fig. 6b). Significantly more *D. citri* chose the arm with clean air compared with DMDS ($\chi^2 = 4.4$; $df = 1$; $P = 0.0350$) or AMDS ($\chi^2 = 6.24$; $df = 1$; $P = 0.0124$). Additionally, significantly more *D. citri* chose the arm with citrus odours compared with DMDS ($\chi^2 = 4.03$; $df = 1$; $P = 0.0447$) or AMDS ($\chi^2 = 4.79$; $df = 1$; $P = 0.0285$) (fig. 6b). The percentages of psyllids moving from the release point to DMTS vs. clean air and DMTS vs. citrus were 11 and 21%, respectively.

Significantly ($F = 114.71$; $df = 11,36$; $P < 0.0001$) more *D. citri* did not move from the release point in treatments in which DMTS, DMDS, AMDS or ADS (all at 1.0%) was co-presented with clean air or citrus odours than when clean air alone or citrus was simultaneously presented in both arms of the olfactometer (fig. 6c). The percentages of *D. citri* not moving from the release point for DMTS vs. citrus and DMTS vs. clean air ranged between 84 and 92%. The percentages of psyllids not moving from release point in treatments with disulfides (DMDS, AMDS and ADS) ranged between 62 and 82% (fig. 6c). Percentages of *D. citri* not moving from the release point were statistically equivalent for trisulfides and disulfides (fig. 6c). There were no differences between the number of *D. citri* entering the olfactometer arm containing AMS vs. clean air ($\chi^2 = 0.01$; $df = 1$; $P = 0.9081$). Significantly ($\chi^2 = 8.92$; $df = 1$; $P = 0.0028$) more psyllids remained in the release arm when the 0.25% blend of DMTS+DMDS vs. clean air was presented than when DMTS alone (0.25%) vs. clean air was presented (fig. 7).

Discussion

As expected, more *D. citri* were attracted to citrus volatiles when citrus leaves were compared with clean air. Furthermore, up to 80% of tested *D. citri* entered the olfactometer in response to light and due to negative geotaxis when clean air was presented in both arms. The current results provide evidence that volatiles from crushed garlic chive leaves inhibited the response of *D. citri* to its normally attractive host plant volatiles. These volatiles also appeared to have inhibited the psyllid's normal geotactic and phototactic responses. Furthermore, our results suggest that the sulfur volatiles released by wounded *A. tuberosum* leaves affected the behaviour of *D. citri*. Volatiles released by plants in response to wounding repel herbivores and deter their feeding (Dicke & van Loon, 2000; Dugravot *et al.*, 2003). Volatiles from *Allium* spp. have been previously reported to repel several insect species including hemipterans (Salas, 2001; Assis *et al.*, 2007).

Allium spp. are known to produce sulfur compounds (thiosulfonates) when leaf tissues are wounded. The disulfide and trisulfide breakdown products of the thiosulfonates are toxic to insect herbivores (Auger *et al.*, 1989). We speculated that the effect of garlic chive plants or essential oil on *D. citri* may be due to the release of sulfur volatiles given their

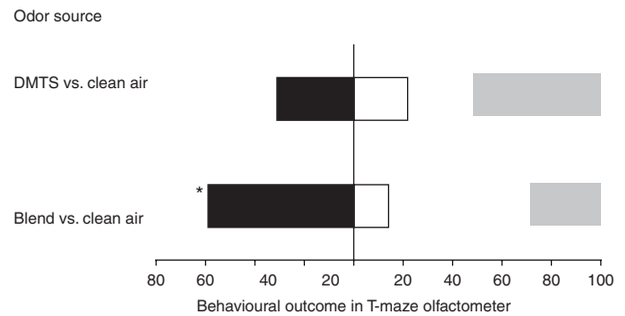


Fig. 7. Responses of *Diaphorina citri* when presented with a blend of 0.25% DMTS+DMDS. Dimethyl trisulfide: DMTS. Blend: DMTS (0.25%) + dimethyl disulfide (0.25%). Black bars labelled with * are significantly different (χ^2 test, $P < 0.05$) (■, release arm (%); □, treatment arm (%); ▒, control arm (%)).

known effect on insect behaviour (Auger *et al.*, 1989; Huang *et al.*, 2000; Block, 2005; Prowse *et al.*, 2006; Amarawardana *et al.*, 2007; Rouseff *et al.*, 2008). GC-MS analysis of garlic chive essential oil and crushed garlic chive leaves revealed five common sulfur-containing compounds (AMS, ADS, AMDS, DMDS and DMTS). Behavioural assays indicated that trisulfides had a greater effect on *D. citri* behaviour than disulfides and that monosulfides had no effect. Crushed wild onion leaves, containing higher proportions of monosulfides than disulfides and trisulfides, as well as wild onion plants did not affect the response of *D. citri*. Disulfides and trisulfides are known plant defensive chemicals and are toxic to several insect species (Auger *et al.*, 1989; Huang *et al.*, 2000; Dugravot *et al.*, 2005; Rouseff *et al.*, 2008). Furthermore, garlic chive essential oil was more repellent to psyllids than garlic chive or wild onion plants, whether they were mechanically wounded or intact. It is possible that the amount of repellent chemical(s) emanating from the essential oil was greater than that from plants. Future descriptive investigations of the effect of airborne concentrations of these odours on psyllid behaviour will be conducted to resolve the role of concentration versus that of specific chemicals on repulsion of *D. citri*.

In addition to effects on behaviour, we also observed that *D. citri* exposed to disulfides and trisulfides died at the 1.0% exposure concentration. DMDS, AMDS and diallyl trisulfide are known to kill several insect species, including *Sitophilus zeamais* Motschulsky (Coleoptera: Curculionidae), *Tribolium castaneum* (Herbst) (Coleoptera: Tenebrionidae) and *Periplaneta americana* (L.) (Dictyoptera: Blattidae) (Norris *et al.*, 1977; Huang *et al.*, 2000; Dugravot, 2003). Sulfur compounds inhibit chemoreception by destroying sulfur electrochemical receptors (Norris *et al.*, 1977) or interfere with the cytochrome oxidase mitochondrial complex, causing loss of neural activity (Dugravot, 2003). Our results indicate that trisulfides were more potent than disulfides. To our knowledge, there are no studies on the mode of action of trisulfide toxicity to insects. Our results on trisulfides corroborate the findings of Huang *et al.* (2000) who reported that trisulfides from *Allium* spp. are more potent insect inhibitors than disulfides.

Both disulfides and trisulfides repelled *D. citri* vs. clean air at the lower concentrations and inhibited response of psyllids to citrus. However, a blend of dimethyl trisulfide and dimethyl disulfide was more inhibitory than the individual

constituents, suggesting an additive effect. Our findings are congruent with that of Podskalska *et al.* (2009) who demonstrated synergistic action of disulfides and trisulfides on *Nicrophorus vespillo* L. (Coleoptera: Silphidae).

In response to herbivore damage, several plant species emit volatiles that attract natural enemies of herbivores (Arimura *et al.*, 2000). Conversely, induced volatiles may have detrimental effects on natural enemies. Therefore, we also investigated whether garlic chive volatiles affect the behaviour of the *D. citri* parasitoid, *T. radiata*. There was no effect of these volatiles on the behaviour of the parasitoid, suggesting their potential use for management of *D. citri* should not affect this natural enemy. The current results indicate that disulfides and trisulfides from crushed garlic chive leaves are potential repellents for development and integration into *D. citri* management programmes. Our current efforts are focusing on formulating these sulfur compounds into controlled release devices for deployment in the field.

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