Variation for Epicuticular Waxes on Onion Foliage and Impacts on Numbers of Onion Thrips

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ABSTRACT. Natural variation exists in onion (Allium cepa L.) for amounts of epicuticular waxes on foliage, and plants with lower amounts of these waxes suffer less feeding damage from onion thrips (Thrips tabaci Lind.). Wild-type onion possesses copious amounts of epicuticular waxes and is often referred to as “waxy.” The recessively inherited “glossy” phenotype has significantly less wax relative to waxy types and shows resistance to onion thrips but is vulnerable to spray damage, foliar pathogens, and excessive transpiration. Phenotypes visually intermediate between waxy and glossy also exist in onion, which we refer to as “semiglossy.” Epicuticular waxes on the leaves of glossy, semiglossy, and waxy onions were evaluated for appearance using scanning electron microscopy (SEM) and amounts and types were analyzed using gas chromatography/mass spectrometry. Wax crystals were clearly visible on the surface of waxy foliage with decreasing amounts on semiglossy and none on glossy leaves. The ketone hentriacontanone-16 was the most prevalent wax on leaves of waxy onion and was significantly (P < 0.01) less on semiglossy relative to waxy plants and on glossy relative to waxy and semiglossy plants. Numbers of adult and immature onion thrips were significantly reduced (P < 0.05) on glossy and/or semiglossy accessions relative to waxy in field and greenhouse cage experiments. These results indicate that semiglossy plants possess intermediate amounts of epicuticular waxes that may protect leaves from diseases or environmental stresses while still conferring resistance to onion thrips. Therefore, the semiglossy phenotype should be useful in integrated programs managing this important onion pest.
is referred to as “waxy.” Jones et al. (1934) observed that the onion cultivar White Persian had a lighter green foliage color, which reduced onion thrips populations and the amount of feeding damage as compared with blue-gray foliage. Jones et al. (1944) subsequently referred to the foliage type from ‘White Persian’ as “glossy” and demonstrated that it is conditioned by the recessive gl locus. Subsequent reports have documented that glossy onions show resistance to onion thrips (Alimousavi et al., 2007; Molenaar, 1984; Mote and Sonone, 1977; Pawar et al., 1975). Molenaar (1984) observed that glossy onions have less epicuticular wax than waxy leaves. Although imparting greater resistance to onion thrips, glossy foliage is not commercially viable because plants show susceptibility to leaf pathogens, excessive transpiration, and spray injury (Baker, 1982; Mohan and Molenaar 2005; Molenaar, 1984). There exists natural variation in onion for amounts of epicuticular waxes intermediate between glossy and waxy types. Jones et al. (1935) described ‘Sweet Spanish’ populations as having an intermediate “shade” of foliage relative to glossy and waxy onions, which we refer to as “semiglossy.” These semiglossy types are lighter green in color, are associated with reduced onion thrips populations, and suffer less damage relative to waxy onions (Diaz-Montano et al., 2010; Maughan and MacLeod, 1936). Because semiglossy onions have intermediate amounts of wax, this phenotype may be useful in an integrated approach to manage onion thrips by reducing sprays and the incidence of thrips-transmitted diseases while maintaining the protection of some wax accumulation on leaves. In this study, we evaluated amounts and types of epicuticular waxes on the leaves of glossy, semiglossy, and waxy onions as well as assessed numbers of adult and larval onion thrips in replicated field and greenhouse cage experiments.

Materials and Methods

Plant materials. Doubled-haploid (DH) onions 2107 and 2150 (Alan et al., 2004) have waxy foliage. U.S. Department of Agriculture (USDA) PI 546303 is a glossy inbred line (B9885C) derived from ‘White Persian’ (Molenaar, 1984). USDA inbred B3531C (cage 905-3) selected from ‘Colorado #6’ and an S1 family from PI 264320 (USDA breeding plot 22142) of ‘Sweet Spanish’ background (SG-A and SG-B, respectively) are semiglossy types with light green foliage and less visible wax as compared with waxy onions.

Scanning electron microscopy of epicuticular wax morphology. Seeds of accessions DH2107, B9885C, 22142, and B3531C were planted in Jan. 2012 in a greenhouse on the University of Wisconsin-Madison (UW) campus. Initial day-length was set at 12 h and was increased on the first of each subsequent month after the spring equinox to the natural daylength in Madison, WI. Temperatures were 25 °C days and 20 °C nights. In Apr. 2012, leaf samples were collected from each accession for analysis (Laudate, 2003) on an FEI Quanta scanning electron microscope. Because the electron beam began to melt waxes within 30 s, various settings were evaluated and the best were 30 kV, pressure = 3 torr, spot size = 5, brightness = 47, and contrast = 97. Micrographs were visually inspected for relative amounts and morphologies of wax crystals on the leaf surface of the waxy, semiglossy, and glossy phenotypes.

Gas chromatography/mass spectrometry analyses of epicuticular waxes. In Feb. 2012, five plants of each of the four accessions were transplanted into 20-cm pots for a total of 20 plants. In Apr. 2012, three samples were collected from a single leaf from each of the 20 plants for a total of 60 samples. In May 2012, two samples were collected from an older leaf and two from a younger leaf from each of the 20 plants for a total of 80 samples. In June 2012, the same procedure was followed as for the May sampling to yield another 80 samples.

In May 2013, seeds of the four accessions were planted in 10-cm pots at the UW greenhouses and in four randomly assigned field plots at the Dean Kincaid farm near Palmyra, WI. In June 2013, five greenhouse-grown plants of each accession were transplanted into 20-cm pots for a total of 20 plants. Greenhouse-grown plants of the four accessions were also transplanted in 2013 into four randomly assigned field plots at the UW Horticulture Research Station near Arlington, WI. At each of the three environments, leaf samples from the four accessions were collected once per month for 3 months for analysis of epicuticular waxes by gas chromatography/mass spectrometry (GC/MS). In June 2013 at the UW greenhouses, two samples were collected from different leaves from each of the 20 plants for a total of 40 samples. In July and Aug. 2013, the same procedure was followed as for the June sampling to yield another 40 samples each month. In June 2013 at each of the Arlington and Palmyra field locations, five randomly selected plants were chosen from each of four plots containing the four accessions for a total of 20 plants. Two samples were collected from different leaves from each of the 20 plants for a total of 40 samples. In July and Aug. 2013, the same procedure was followed as for the June sampling to yield another 40 samples each month from each field location. In total, 12 samplings occurred across four (two greenhouse and two field) environments and yielded a total of 580 samples.

Immediately after sampling, the fresh weight in grams of each leaf piece was measured and an aliquot of docosane (Sigma-Aldrich, St. Louis, MO), dissolved in high-performance liquid chromatography (HPLC) reagent grade chloroform (Fisher Scientific, Hampton, NH), was applied to the leaf sample at the rate of 1 μg per 0.3 g of tissue and allowed to air-dry. Leaf pieces were placed in 16 × 100-mm glass vials (Fisher Scientific) and covered with aluminum foil for immediate transport to the laboratory. Epicuticular waxes were extracted by adding enough HPLC reagent-grade chloroform to each vial to submerge the leaf sample for 60 s. Leaf tissue was removed and discarded. Chloroform in vials was allowed to evaporate in a fume hood without agitation. After evaporation, 500 μL of anhydrous grade chloroform, 600 μL of acetoni-trile, and 210 μL of the derivatization agent N,O-Bis(trimethylsilyl)trifluororaceta-mide (BSTFA) were added to each vial (all from Sigma-Aldrich). Vials were capped with phenolic polytetrafluoroethylene (PTFE)-lined caps (Kimble Chase, Rockwood, TN) and derivatized in an oven at 80 °C for 30 min. Each sample was filtered through a Supelco Iso-disc PTFE-13-2 filter (Sigma-Aldrich) into a Supelco 2-mL PTFE/Silcone (Sigma-Aldrich) capped glass vial using a 1-mL disposable syringe (Fisher Scientific). The samples were analyzed on a GC/MS (GCMS-QP2010 SE; Shimadzu, Kyoto, Japan) equipped with an SHRXI-5MS column and an AOC-5000 autoinjector. The machine used a split/splitless glass inlet liner, Thermolite septum (Restek, Bellefonte, PA), and a 10-μL Hamilton syringe. Each sample cycle began with split injection (ratio = 1/20) of 1 μL of sample at 250 °C and used a programmed column temperature consisting of 10 min at 150 °C, 15 min increasing temperature.
at a rate of 10 °C·min⁻¹, and 10 min at 300 °C. Mass scan range was between 35 and 600 m/z. Chemical compounds in each sample were identified by post-run software (Shimadzu) comparing mass spectra with a library (NIST11) with a similarity threshold of 80%. Quantities were estimated by the area under peaks on the total ion chromatogram. All peak areas were normalized to the docosane internal standard. Relative retention times were calculated by dividing compound retention times by those of the internal standard docosane.

**Statistical analyses of GCMS results.** A mixed-model analysis of variance (ANOVA) using the repeated-measures option was performed in SAS (Version 9.3; SAS Institute, Cary, NC). Individual plants were regarded as experimental units and the three sampling time points were considered as repeated measures. GC/MS data for each compound was considered as a unique, dependent variable. Accessions and samplings were considered as fixed and environments, plants, and subsamples as random variables. Departure from normality of model residuals was corrected by a square-root transformation. Significant differences for quantities of compounds detected by GC/MS were determined using least-squared means and probability values of all pairwise comparisons among accessions.

**Confirmation of compounds detected by GCMS.** After determining the most prevalent compounds associated with the foliar phenotypes, hentriacontanone-16, triacontanol-1, and octacosanol-1 were obtained from TCI Chemicals (Portland, OR). These compounds and docosane were prepared for GC/MS analysis according to the method of standard addition (Harris, 2003) to confirm the identities of these epicuticular waxes and to calculate absolute quantities. Four leaf samples from waxy DH2150 were separately submerged into 2 mL of HPLC reagent-grade chloroform as previously described. Three dilutions of the four compounds were made in HPLC reagent-grade chloroform at 1, 5, and 10 µg·mL⁻¹ in a final volume of 500 µL. A fourth vial was filled with 500 µL of chloroform for each set of dilutions. Five hundred microliters of the epicuticular wax solution from DH2150 were added to each of the four vials for each compound; then 500 µL of acetonitrile and 200 µL of BSTFA were added to each vial. Vials were capped, derivatized, filtered, and analyzed on the GC/MS as previously described. The retention times and mass spectra of peaks for the three compounds from TCI Chemicals were compared with the epicuticular waxes from DH2150. A linear regression of peak areas against concentrations for each set of dilutions was performed. The difference between the origin and X-intercept provided the concentration of the compound from the DH2150 leaf sample. The regression slopes for octacosanol-1, triacontanol-1, and hentriacontanone-16 were then used to calculate concentrations represented by peak area least square means for each of the waxes. Concentrations of other fatty alcohols were calculated using the average slope of octacosanol-1 and triacontanol-1. Other alkane compound concentrations were calculated using the docosane regression. Based on sample fresh tissue weights and compound concentrations, micrograms of wax per gram of fresh leaf tissue were calculated for each compound for each of the four accessions.

**Assessment of onion thrips populations on foliage types.** In May 2013, the four accessions were planted in fields at the UW Horticulture Research Farm near Arlington, WI, and at Dean Kincaid farm near Palmyra, WI. Each field had four replications of the four accessions in a 4 × 4 Latin-square design with 30 to 50 plants in each 4-m plot. A Latin-square design was selected to increase the likelihood that replicate sets of experimental treatments (accessions) would be equally exposed to mobile, colonizing adult onion thrips that could disperse into experiment plots from several directions. Plots at Arlington were not sprayed with insecticide; Palmyra plots received five insecticide sprays between 7 Aug. and 3 Sept. based on established thresholds for onion thrips management (Nault et al., 2012). Plots were sampled by cutting two randomly selected plants at the soil level and placing them in 3.8-L plastic zip-lock bags. The bags were transported to Madison where 500 mL of 95% ethanol was added. Each bag was agitated for 3 min to wash all life stages (except eggs) off of the plant. Plants were discarded and insects were recovered by filtering the ethanol through 150-µm screens (Greenhouse Megastore, Danville, IL). All adult and larval stages of onion thrips were counted under a stereoscope. Onion thrips were first observed on accessions at Arlington in mid-July and sampling occurred on a weekly basis for 4 weeks thereafter. Onion thrips were first observed at Palmyra at the end of August and population sampling occurred twice over the next 2 weeks.

For choice and no-choice cage experiments in the greenhouse, onion thrips were obtained from G.G. Kennedy (North Carolina State University, Raleigh, NC) and maintained on the laboratory bench at ambient conditions using a 13 × 20-cm plastic container with fresh cabbage leaves replaced every 3 to 4 d. For the choice experiments, 10 cylindrical cages were constructed from 30 × 76-cm polyethylene plastic sheets (Interstate Plastics, Sacramento, CA) bonded on the 30-cm edges with tape (Killer Red Tape; ePlastics, San Diego, CA). One end of each cylinder was covered with thrips proof 150-µm screen using plastic adhesive glue (Weld-on 16; ePlastics). The open end of each cage fit inside the rim of a 25-cm pot. One plant of each accession was placed in each of 10 25-cm pots and cages were placed over each pot. Fifty adult onion thrips were collected from the colony by aspiration into a glass pipet and introduced into each cage by placing the pipet upright in the center of the pot and left undisturbed for 14 d. For the no-choice experiments, 20 cylindrical cages were built from 30 × 61-cm polyethylene plastic sheets bonded on the 30-cm edges with killer red tape and covered with thrips-proof 150-µm screen as described previously. The open end of each cage fit within the rim of a 20-cm pot. Five plants of each accession were planted separately in 20 20-cm pots and cages placed over each pot. Each cage was infested with 30 adult onion thrips as described previously. Plants from choice and no-choice experiments were harvested into gallon plastic zip-lock bags and weighed. Plants were washed with ethanol and insects counted as previously described.

**Experimental design and statistical analysis of numbers of onion thrips.** For 2013 field experiments at Arlington and Palmyra, a mixed-model ANOVA with the repeated-measures option was performed in SAS (SAS Institute, Inc., Cary, NC) to assess the effect of onion accession on numbers of adult and larval onion thrips. Plots were considered as experimental units and the four sampling at Arlington and two at Palmyra were considered as repeated measures. Adult and larval numbers per plant were dependent variables. Accessions were considered fixed and environments, range, row, and subsamples as random. For the experiments at UW greenhouses, a mixed-model ANOVA was performed and cages were considered as replications and individual plants as experimental units. Adult and larval counts were considered as dependent variables.
Accessions were considered fixed and replication as random. Plant weight was used as a covariate to account for a possible attraction of onion thrips to larger plants.

**Results and Discussion**

**EPICUTICULAR WAX MORPHOLOGY.** SEM micrographs (Fig. 1) revealed no obvious wax crystals on glossy leaves, a thin scattering of spiky, plate-like crystals on semiglossy leaves, and large quantities of the spiky, plate-like crystals on waxy leaves. According to descriptions of epicuticular wax crystals by Jeffree (1986), the spiky crystals on onion foliage are likely formed by a ketone. Studies by Gülz et al. (1992, 1993) concluded that characteristic lipid crystals can be observed on leaves if one lipid class comprises at least 40% of the total profile and if at least 80% of that class is one compound. Therefore, SEM indicated that ketone(s) may predominate in the epicuticular waxes on onion foliage.

**GC/MS ANALYSIS OF EPICUTICULAR WAX.** GC/MS is a powerful technique for identifying and quantifying epicuticular waxes (Kolattukudy and Espelie, 1985; Tulloch, 1976; Walton, 1990) and revealed carbohydrate, short hydrocarbon, and long hydrocarbon chain (wax) compounds in chloroform-extracted samples from onion leaves. Pairwise comparisons among wax compounds accounted for the vast majority of the highly significant ($P < 0.01$) differences among glossy, semiglossy, and waxy phenotypes. Twenty-one comparisons among eight wax compounds (Table 1) and two comparisons among two non-wax compounds (butanedioic acid and myo-inositol) were significantly different at $P < 0.01$. The two non-wax compounds likely originated from injured leaf cells and may not contribute significantly to compounds on leaf surfaces. Accession-by-sampling interactions were not significant for wax compounds, revealing that differences for amounts of waxes were similar over time in both field and greenhouse environments.

Hentriacontanone-16 as the most prevalent on waxy leaves (Table 1) and its quantities were significantly different among waxy, semiglossy, and glossy accessions (Table 2). Relative to waxy, quantities of this ketone were intermediate on semiglossy and lowest on glossy leaves. These GC/MS results are consistent with the morphology of wax crystals under SEM. Hentriacontanone-16 forms crystals at concentrations as low as 23% of total waxes (Gülz et al., 1992); crystals were observed on waxy (52%) and semiglossy (35% to 39%) but not glossy (19%), leaves (Table 1). The crystals observed on waxy leaves were also similar in appearance to those of hentriacontanone-16 on the "cross-rippled" leaf surface of *Liriodendron tulipifera* L. (Gülz et al., 1992). SEM and GC/MS analyses provide strong evidence that hentriacontanone-16 is the predominant epicuticular wax on onion foliage and the most reduced in the glossy and semiglossy phenotypes. Rhee et al. (1998) reported that hentriacontanone-16 is the predominant epicuticular wax on leek (*Allium ampeloprasum* L.) foliage.

Other waxes detected by GC/MS included four fatty alcohols (hexacosanol-1, octacosanol-1, heptadecanol-1, and triacontanol-1) and three alkanes (heptacosane, 1-ethenylxyloctadecane, and 2-methyl octacosane). Similar amounts of octacosanol-1 and triacontanol-1 on waxy and glossy leaves suggest that these compounds do not contribute to the visual phenotype. Quantities of other waxes were relatively small, albeit significantly ($P < 0.01$) different between waxy DH2107 and the other accessions. Four of the eight waxes in Table 1 were significantly ($P < 0.01$) different between two semiglossy accessions (B5351C and 22142), revealing that phenotypically similar plants have different amounts of specific waxes. The total amount of wax on the leaves of semiglossy B5351C was not significantly different from glossy B9885C (Table 2); however, B5351C accumulated enough hentriacontanone-16 (39% of total waxes) to appear visually as semiglossy (Table 1).

Comparison of mass spectra and retention times between commercially available hentriacontanone-16, octacosanol-1, and triacontanol-1 and samples of epicuticular waxes from leaves of waxy DH2150 confirmed the identities of these compounds. Standard addition regression lines were used to estimate quantities in micrograms of waxes per gram of leaf tissue across the four accessions (Table 2).

**NUMBERS OF ONION THRIPS.** Numbers of adult and larval onion thrips in the field choice experiments were significantly ($P < 0.05$) different between waxy DH2107 and the other accessions, but there were no significant differences among the glossy and semiglossy onions (Table 3). Larval numbers were higher than adults (Table 3) and presumably are responsible for most of the direct damage to leaf tissues (Coudriet et al., 1979). As expected, infestation levels at Palmyra were much lower than Arlington as a result of insecticide sprays. Numbers of adults were not significantly different among accessions at Palmyra; however, differences for larvae between DH2107 and

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Fig. 1. Scanning electron micrographs of surfaces of glossy B9885C (left), semiglossy 22142 (middle), and waxy DH2107 (right) onion leaves at magnification of 5000×. Elliptical shapes are stomata and 10-μm scales are shown in lower right corners.
both B9885C and B5351C were significant ($P < 0.05$). Overall, the field results indicate that semiglossy plants may support fewer adults and larvae of onion thrips relative to waxy while retaining more epicuticular waxes than the glossy phenotype.

In the caged choice experiments, larval numbers were significantly ($P < 0.05$) different between waxy DH2107 and the two semiglossy accessions, but not between glossy B9885C and waxy DH2107 (Table 3). Numbers of adult onion thrips in the no-choice experiment revealed significant differences between waxy DH2107 and the other accessions and between DH2107 and semiglossy 22142 for larvae. These results do not agree with Molenaar (1984), who observed significant differences between waxy and glossy plants in greenhouse choice experiments. However, Lowe et al. (1985) observed resistance to grain aphid (Sitobion avenae F.) on glossy wheat (Triticum aestivum L.) cultivars in the field but found that differences in both wax bloom and resistance were diminished in a greenhouse environment. Similarly, we observed fewer significant differences for numbers of onion thrips among accessions in our greenhouse experiments.

Fewer onion thrips in the field and cage experiments were consistent with reduced amounts of epicuticular waxes, suggesting that specific waxes or total wax amounts may be important for either host recognition by the insect or preference for feeding or ovipositioning. Although statistical analyses considered accessions as a fixed variable and greenhouse experiments had fewer significant differences among the onion accessions for numbers of onion thrips, our results agree with other reports that the glossy and semiglossy phenotypes of onion significantly contribute to onion thrips resistance (Alimousavi et al., 2007; Diaz-Montano et al., 2010; Jones et al., 1934; Maughan and MacLeod, 1936; Molenaar, 1984; Mote and Sonone, 1977; Pawar et al., 1975). Types of insect resistance in plants have been classified into three general categories: antixenosis (non-preference), antibiosis, and tolerance (Kogan and Ortman, 1978; Painter, 1951). Antibiosis refers to plant characteristics that have an adverse effect as the insect feeds. Tolerance refers to the ability of an infested plant to perform similarly to non-infested plants under comparable pest population densities. Antixenosis refers to the existence of plant characteristics that repel insects or do not induce an important host-recognition behavioral response. Lower amounts of total wax, or less of specific waxes such as hentriacontanone-16, on leaf surfaces could be less attractive to onion thrips (antixenosis) and/or have an adverse effect on the insect as it feeds or reproduces (antibiosis). Lower numbers of onion thrips on semiglossy foliage.

### Table 1. Major epicuticular waxes, formulas, relative retention time (RRT), and percent of total waxes detected using gas chromatography/mass spectrometry on leaves of four onion accessions evaluated across four environments.

<table>
<thead>
<tr>
<th>Compound</th>
<th>Formula</th>
<th>RRT</th>
<th>GL</th>
<th>SG-A</th>
<th>SG-B</th>
<th>WX</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hentriacontanone-16</td>
<td>C₃₁H₆₂O</td>
<td>1.47</td>
<td>19.0</td>
<td>39.0</td>
<td>34.7</td>
<td>52.0</td>
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<td>Heptacosane</td>
<td>C₂₇H₅₆</td>
<td>1.37</td>
<td>1.4</td>
<td>9.5</td>
<td>2.5</td>
<td>1.4</td>
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<td>Heptadecanol-1</td>
<td>C₁₇H₃₄O</td>
<td>1.45</td>
<td>3.3</td>
<td>8.5</td>
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<td>6.2</td>
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<td>Hexacosanol-1</td>
<td>C₂₆H₄₂O</td>
<td>1.31</td>
<td>6.3</td>
<td>3.0</td>
<td>4.6</td>
<td>2.5</td>
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<tr>
<td>Octacosanol-1</td>
<td>C₂₈H₅₂O</td>
<td>1.39</td>
<td>52.0</td>
<td>24.3</td>
<td>35.3</td>
<td>26.4</td>
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<tr>
<td>1-Ethenyloxy octadecane</td>
<td>C₂₀H₄₀O</td>
<td>1.35</td>
<td>3.4</td>
<td>2.3</td>
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<tr>
<td>2-Methyl octacosane</td>
<td>C₂₉H₄₀</td>
<td>1.29</td>
<td>1.6</td>
<td>4.1</td>
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</tr>
<tr>
<td>Triacontanol-1</td>
<td>C₃₀H₆₂O</td>
<td>1.50</td>
<td>13.0</td>
<td>9.4</td>
<td>16.1</td>
<td>9.6</td>
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</table>

*GL = glossy B9885C; SG-A = semiglossy B5351C; SG-B = semiglossy 22142; WX = waxy DH2107.

### Table 2. Amounts in micrograms of eight epicuticular waxes per gram of fresh leaf tissue and least significant differences calculated from least squares means of gas chromatograph peak areas for four onion accessions across four environments.

<table>
<thead>
<tr>
<th>Compound</th>
<th>Amount of waxes (µg g⁻¹)*</th>
<th>Least significant difference</th>
</tr>
</thead>
<tbody>
<tr>
<td>GL</td>
<td>SG-A</td>
<td>SG-B</td>
</tr>
<tr>
<td>Hentriacontanone-16</td>
<td>13.1</td>
<td>20.5</td>
</tr>
<tr>
<td>Heptacosane</td>
<td>0.9</td>
<td>5.0</td>
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<td>Heptadecanol-1</td>
<td>2.3</td>
<td>4.5</td>
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<td>Hexacosanol-1</td>
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<td>1.6</td>
</tr>
<tr>
<td>Octacosanol-1</td>
<td>35.9</td>
<td>12.8</td>
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<tr>
<td>1-Ethenyloxy octadecane</td>
<td>2.4</td>
<td>1.2</td>
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<tr>
<td>2-Methyl octacosane</td>
<td>1.1</td>
<td>2.1</td>
</tr>
<tr>
<td>Triacontanol-1</td>
<td>9.0</td>
<td>4.9</td>
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<tr>
<td>Total wax</td>
<td>69.1</td>
<td>52.6</td>
</tr>
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</table>

*GL = glossy B9885C; SG-A = semiglossy B5351C; SG-B = semiglossy 22142; WX = waxy DH2107.

### Table 3. Least squares means and least significant differences for numbers of adult and larval onion thrips per plant in field choice and greenhouse choice and no-choice experiments.

<table>
<thead>
<tr>
<th>Location</th>
<th>Treatment</th>
<th>Stage</th>
<th>Mean accessions (no.)</th>
<th>Least significant difference (0.05)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arlington</td>
<td>Choice</td>
<td>Adults</td>
<td>3.1</td>
<td>2.7</td>
</tr>
<tr>
<td>Arlington</td>
<td>Choice</td>
<td>Larvae</td>
<td>15.7</td>
<td>12.1</td>
</tr>
<tr>
<td>Palmyra</td>
<td>Choice</td>
<td>Adults</td>
<td>0.7</td>
<td>1.1</td>
</tr>
<tr>
<td>Palmyra</td>
<td>Choice</td>
<td>Larvae</td>
<td>3.4</td>
<td>1.9</td>
</tr>
<tr>
<td>Greenhouse</td>
<td>Choice</td>
<td>Adults</td>
<td>3.9</td>
<td>3.4</td>
</tr>
<tr>
<td>Greenhouse</td>
<td>Choice</td>
<td>Larvae</td>
<td>44.3</td>
<td>25.2</td>
</tr>
<tr>
<td>Greenhouse</td>
<td>No-choice</td>
<td>Adults</td>
<td>3.6</td>
<td>3.0</td>
</tr>
<tr>
<td>Greenhouse</td>
<td>No-choice</td>
<td>Larvae</td>
<td>45.8</td>
<td>55.1</td>
</tr>
</tbody>
</table>

*GL = glossy B9885C; SG-A = semiglossy B5351C; SG-B = semiglossy 22142; WX = waxy DH2107.
together with protection from diseases or environmental stresses provided by intermediate amounts of epicuticular waxes, support the commercial use of this phenotype to potentially reduce the number of insecticide sprays over the production season.

**Literature Cited**


