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Pezet, J., Elkinton, J., Gomez, S., McKenzie, E. A., Lavine, M., & Preisser, E. (2013). Hemlock woolly adelgid and elongate hemlock scale induce changes in foliar and twig volatiles of eastern hemlock. Journal of Chemical Ecology, 39(8), 1090-100. Available at:<http://dx.doi.org/10.1007/s10886-013-0300-5>

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## Hemlock Woolly Adelgid and Elongate Hemlock Scale Induce Changes in Foliar and Twig Volatiles of Eastern Hemlock

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**Abstract**— Eastern hemlock (*Tsuga canadensis)* is in rapid decline because of infestation by the invasive hemlock woolly adelgid (*Adelges tsugae;* 'HWA') and, to a lesser extent, the invasive elongate hemlock scale (*Fiorinia externa;* 'EHS'). For many conifers, induced oleoresin-based defenses play a central role in their response to herbivorous insects; however, it is unknown whether eastern hemlock mobilizes these inducible defenses. We conducted a study to determine if feeding by HWA or EHS induced changes in the volatile resin compounds of eastern hemlock. Young trees were experimentally infested for three years with HWA, EHS, or neither insect. Twig and needle resin volatiles were identified and quantified by gas chromatography/mass spectrometry. We observed a suite of changes in eastern hemlock's volatile profile markedly different from the largely terpenoid-based defense response of similar conifers. Overall, both insects produced a similar effect: most twig volatiles decreased slightly, while most needle volatiles increased slightly. Only HWA feeding led to elevated levels of methyl salicylate, a signal for systemic acquired resistance in many plants, and benzyl alcohol, a strong antimicrobial and aphid deterrent. Green leaf volatiles, often induced in wounded plants, were increased by both insects, but more strongly by EHS. The array of phytochemical changes we observed may reflect manipulation of the tree's biochemistry by HWA, or simply the absence of functional defenses against piercing-sucking insects due to the lack of evolutionary contact with these species. Our findings verify that HWA and EHS both induce changes in eastern hemlock's resin chemistry, and represent the first important step toward understanding the effects of inducible chemical defenses on hemlock susceptibility to these exotic pests.

**Key Words**—*Adelges tsugae*; *Fiorinia externa*; *Tsuga canadensis*; plant-insect interactions; conifer volatiles: induction.

#### INTRODUCTION

Conifers in the family Pinaceae are among the largest and longest-living organisms on earth. Their striking longevity means that individual trees face an imposing array of biotic and abiotic challenges. They respond to these challenges via complex constitutive and inducible defenses that enable them to survive under highly diverse and taxing conditions and dominate vast areas of the earth's temperate and alpine forests (Trapp and Croteau 2001, Dudareva et al. 2006).

Conifers commonly use oleoresin-based chemical defenses to combat herbivorous insects and pathogens (Zulak and Bohlmann 2010). Oleoresin, or simply 'resin,' is a complex and species-specific mixture of phytochemicals that is usually dominated by volatile monoterpenoids and non-volatile diterpenoid acids but also contains smaller amounts of volatile organic chemicals such as sesquiterpenoids, benzenoids (including phenolics), and fatty acid derivatives. These compounds are produced in resin-cells of buds, needles and woody tissue, and in some conifers (such as *Pinus* species) they accumulate in intercellular ducts or canals either constitutively or in response to trauma (Keeling and Bohlmann 2006). Many conifers can respond to insect and microbial challenges via inducible increases in the biosynthesis and accumulation of resin (Hudgins et al. 2004). These defenses variously act to physically engulf and expel insects from the tree by the force of resin flow, seal off infected regions from surrounding tissue, 80 deter herbivory or oviposition, chemically interfere with insect developmental pathways, ATP production and nervous system functioning, and disrupt microbial cell membranes causing cell leakage and death (Langenheim 1994, Eyles et al. 2010). Herbivore attack can also induce the release of volatile resin semiochemicals that attract predators of the colonizing plant-feeder (Mumm et al. 2003, Koepke 2010). 84 Over the last century, factors such as non-native pest introductions, forestry practices, and climate change have sharply increased the amount of conifer mortality due to pests or pathogen (Trapp and Croteau 2001, Cudmore et al. 2010). The increasing frequency and severity of such outbreaks have spurred intensive molecular and biochemical research into the factors underlying host susceptibility and pest/pathogen defense in spruce (*Picea*; Bohlmann 2008), fir (*Abies*; Hain et al. 1991, Lewinsohn et al. 1993a), and pine (*Pinus*; Sampedro et al. 2011) species. Defense induction in conifers by mechanical wounding (Lewinsohn et al. 1993a), experimental insect attack (Miller et al. 2005, Sampedro et al. 2011) or 'simulated' herbivory by application of chemical elicitors such as methyl jasmonate (Martin et al. 2002,

2003, Sampedro et al. 2010) leads to dramatic increases in bark and stem-wood terpenoid accumulation and volatile release from needles. An increasing number of the active genes and biosynthetic enzymes underlying defensive chemical outputs in these conifer systems have been identified, establishing strong evidence that resin-based—and primarily terpenoid-based—chemical defenses are central to the trees' evolved responses to insect or pathogen colonization (Franceschi et al. 2005, Keeling and Bohlmann 2006). In eastern North America, the invasive twig-feeding hemlock woolly adelgid (*Adelges tsugae*; 'HWA') threatens to extirpate the native eastern hemlock (*Tsuga canadensis* Carr.; McClure and Cheah 99 1999). The first documented population of the adelgid in eastern North America was detected in the early 1950s, and appears to be of Japanese origin (Havill et al. 2006). The insect has now spread to the southern extent of eastern hemlock's range in northern Georgia, and is moving northward into Vermont, New Hampshire, and Maine (Preisser et al. 2008, Forest Health Protection Program 2011). The insect can take a 103 year or two to reach high densities, but its effect on hemlocks is needle desiccation, branch mortality, and marked suppression of new spring growth, often leading to tree death in four years or less (McClure 1991a). As the only native shade-tolerant conifer in the eastern United States, eastern hemlock acts as a foundation species (sensu Ellison et al. 2005) that creates cool and moist microclimates in the midst of deciduous forests. The nearly complete removal of mature and seedling eastern hemlocks following HWA infestation (Preisser et al. 2011) substantially increases soil and stream temperatures, alters soil chemistry and nutrient cycling patterns, and favors fast-growing, early-successional trees—a series of changes that dramatically transforms the forest landscape (Orwig et al. 2008, Gandhi and Herms 2010). The elongate hemlock scale (*Fiorinia externa*; 'EHS') is another exotic pest of eastern hemlock; an armored scale 112 introduced to the Northeastern United States in the early 20<sup>th</sup> century, this insect is also now present in much of the tree's range and continues to spread northward (Preisser et al. 2008). Reports seemingly based on observational, rather than experimental, evidence suggest that although EHS is usually not lethal, high densities can cause significant needle loss and contribute to the mortality of already stressed trees (McClure 1980, Abell and Van Driesche 2012).

Despite the existence of several studies documenting a correlation between terpenoid levels and herbivory in eastern hemlock, there has been no direct investigation into whether either of these exotic 119 pests elicits resin defenses in eastern hemlock. One study reported a positive correlation between volatile



or composition of volatile resin compounds. We hypothesized that an agent as rapidly lethal as HWA would elicit a defensive resinosis typical of many conifers of Pinaceae: pronounced increases in toxic or deterrent phytochemicals, especially terpenoids. We also predicted that the much milder effects of EHS on the host tree's physiology (Miller-Pierce et al. 2010, Radville et al. 2011, Gonda-king et al. 2012, Gomez et al. 2012) would be accompanied by an induced resin response distinct from that of HWA.

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#### 154 METHODS AND MATERIALS

*Study System.* Eastern hemlock buds begin opening in May in the Northeastern United States, and the young new growth shoots, at first green and pliant, complete their elongation at approximately the end of summer. By that time the foliage has hardened and taken on the form and appearance of the fully mature, previous year growth.

The HWA completes two clonal generations per year in eastern North America, as in its range of origin in Japan (McClure and Cheah 1999). In the Northeastern United States, first instar nymphal crawlers of the progredien generation settle in April (before bud-break) on already mature previous year's growth 162 just below the needle abscission layer and feed through a stylet bundle on xylem ray parenchyma cells in the twig (Young et al. 1995). The sexuparae, a winged, sexually reproducing generation of HWA, hatch concurrently with the clonal female progrediens and, in Japan, subsequently disperse to a spruce (Picea) primary host to complete reproduction. Sexuparae in North America are unable to complete their life-cycle due to the absence of a suitable spruce host andthus, only asexual reproduction occurs. The sessile progredien adults complete egg laying in June, at which point the crawlers of the sisten generation emerge, settle preferentially on the new, young current year's growth, and promptly enter aestivation. In early fall, by the time the new growth has matured, the sisten nymphs resume feeding, completing development and oviposition in April (McClure and Cheah 1999).

The EHS completes two full generations per year in its natural range in Japan, but in the Northeastern United States it appears to lack a distinct and regular cycle of life stage development, and completes between one and two generations annually (Abell and Driesche 2012). First instar nymphs begin to hatch in early June and settle preferentially on the undersides of young hemlock needles. EHS is also a sessile stylet feeder, inserting a thread-like stylet bundle and sucking fluid from needle mesophyll cells

(McClure 1980). Since generation times in the Northeast are irregular, life stages appear to overlap and often two or more instars may be found developing concurrently on the same foliage (Abell and Driesche 2012).

*Experimental Design.* In April 2007, eastern hemlock saplings (0.7-1.0m) were removed from Cadwell Experimental Forest (Pelham, MA, USA) and planted in an open field setting (East Farm, Kingston, RI, USA) in a rectangular grid. The source forest was free of both HWA and EHS at the time of collection, and careful inspection of the sapling trees revealed no prior infestation by either insect. Artificial infestation with HWA, EHS, or neither insect was randomly applied to the saplings. Because both insects are wind-dispersed during their first-instar crawler phase, each tree (including all uninfested controls) was enclosed in a mesh cage annually from early spring to late fall to prevent cross-contamination. Each of the 187 1 m x 1 m x 2 m (length by width by height) cages consisted of a plastic PVC pipe frame covered by 188 mosquito netting (97 holes/cm<sup>2</sup> mesh size). Weed-inhibiting fabric  $(1 \text{ m}^2)$  was placed around the base of each tree. By 2010, a combination of insect cross-contamination and tree death from transplantation-related stress reduced the level of replication to nine trees in the HWA treatment, seven trees in the EHS treatment, and eight trees in the control treatment. *Insect Inoculations.* Insect inoculations were conducted following standard procedures (see Butin et al. 2007). Briefly, trees were inoculated with insects each spring from 2007 to 2010 to mimic natural infestation cycles. Immediately prior to crawler emergence (May for HWA, June for EHS), naturally-infested branches with comparable insect densities were collected from sites in southern New England and attached to trees in the appropriate treatment group; control trees received uninfested branches. Individual branches were placed in aquapics to slow needle desiccation and decrease insect mortality. *Plant Material.* Plant tissue samples were collected from each tree in late June 2010 (fully mature, previous year foliage segments) after the first-instar crawlers of both insects had settled and commenced feeding, and again in mid-October 2010 (young, current year growth twigs) after settled HWA had ceased

aestivation and resumed feeding. An average of 10 cm of twig with foliage was clipped; in the case of the

insect treatments, infested foliage samples were selected. Each sample was placed in a polypropylene 205 cryovial, flash-frozen in liquid nitrogen, transported to the laboratory on dry ice and stored at  $-80^{\circ}$ C until extraction and analysis.

*Extraction of Resin Volatiles.* Extraction of resin volatiles was modified from a protocol developed by

Lewinsohn et al. (1993b). All reagents and reference standards were obtained from Sigma-Aldrich (St.

Louis, MO, USA). Solvents were HPLC or GC grade purity.

Needles were separated from twigs and ground to a homogenous powder using a mortar and pestle under liquid nitrogen. Approximately 100-200 mg (dry weight) of needle tissue was combined with methyl 213 *tert*-butyl ether (MTBE; 1.3-1.5 mL) containing a known concentration of isobutylbenzene (40 µg mL<sup>-1</sup>) as an internal standard in a pre-weighed 2 mL vial (glass with PTFE-coated screw cap, Sigma-Aldrich, St. Louis, MO, USA). Needle samples were extracted overnight (20 h) with constant shaking at room 216 temperature. Each extract was transferred to a fresh glass vial and washed with aqueous  $(NH_4)_2CO_3 (0.3)$ mL, 1 M) to neutralize acidic impurities. The organic layer was then filtered through a Pasteur pipette 218 column packed with silica gel (0.3 g, Sigma-Aldrich,  $60\text{\AA}$ ) overlaid with MgSO<sub>4</sub> (0.2 g). Oxygenated 219 volatile compounds were subsequently eluted by washing the filter with diethyl ether (1 mL), and 220 combined eluates were collected in a GC vial (PTFE-coated screw cap, Agilent Technologies, Santa Clara, 221 CA, USA) and stored at -20<sup>o</sup>C until analysis. Twig samples of approximately 10-50 mg (dry weight) were cooled with liquid nitrogen in a mortar and pestle, ground to a coarse powder, and combined with MTBE (1.0 mL) containing 224 isobutylbenzene (2  $\mu$ g mL<sup>-1</sup>) in a 2 mL glass vial. Twigs were extracted overnight (19 h) with constant 225 shaking at room temperature. Aqueous  $(NH_4)_2CO_3$  (0.2 mL; 1 M) was added to each extract, followed by 226 thorough mixing. The organic layer was then transferred directly to a Pasteur pipette filter packed with 227 silica gel (0.2 g, 60Å) overlaid with MgSO<sub>4</sub> (0.13 g). The filter was washed with diethyl ether (0.5 mL), and combined eluates were collected and stored as described above. After extraction, each sample was dried for at least 48 hours at 55-60°C and weighed for the determination of tissue dry weight. 

*Analysis of Resin Volatiles.* Needle volatile extracts were analyzed on a Hewlett-Packard (HP) 6890 GC 232 equipped with a flame ionization detector (FID). For all analyses, the injection volume was  $1 \mu L$ , injector temperature 220°C. Volatile compounds were separated on an Agilent DB-5, 0.25 mm i.d. x 30 m, 0.25 µm 234 coating thickness, fused silica capillary column. H<sub>2</sub> carrier gas flow was a constant 1.0 mL min<sup>-1</sup> and the 235 split ratio was 20:1. The FID was heated to 250°C, with H<sub>2</sub> flow at 40 mL min<sup>-1</sup>, air flow 350 mL min<sup>-1</sup>, 236 and constant make-up flow  $(N_2)$  at 45 mL min<sup>-1</sup>. The GC oven was programmed with an initial temperature 237 of 60°C (no hold), an increase at 3°C min<sup>-1</sup> to 156°C, then 50°C min<sup>-1</sup> to 300°C (hold 3 min). GC-FID generated peaks were integrated using HP ChemStation software (Agilent technologies). Datafiles for five of the October needle samples were corrupted, reducing the level of replication to seven trees in the HWA treatment, six trees in the EHS treatment, and six trees in the control treatment. For all compound identifications, as well as all twig volatile quantification, analyses were 242 performed on a Shimadzu GC-2010 system equipped with a QP2010-Plus mass spectrometer (EI mode, 70 eV), running GCMSolution software (Shimadzu Corporation, Kyoto, Japan). Separations were performed 244 on the same column as described above for GC-FID. The injection volume was 1 µL and injector 245 temperature 220 $^{\circ}$ C. Helium carrier gas flow was in constant linear velocity mode at 36.5 cm sec<sup>-1</sup>, with 246 column flow set at 1.0 mL min<sup>-1</sup> and a split ratio of 5:1. The GC oven was programmed with an initial 247 temperature of 60°C (no hold), an increase at 3°C min<sup>-1</sup> to 175°C, then 30°C min<sup>-1</sup> to 300°C (hold 5 min). The interface and ion source temperatures were both set at 300°C, and the MS scan range was *m/z* 40-400. Identification of each volatile compound was, wherever possible, based on comparison of the experimental retention time and mass spectrum with those of an authentic standard (indicated in Table 1); when a pure standard was unavailable, tentative identification was based on comparison with retention index and mass spectral information reported in the literature (Adams 2001) and with mass spectra in the NIST05 and NIST05s mass spectral libraries (Stein 2005). Concentrations of all compounds were determined by normalizing integrated peak areas against that of the internal standard isobutylbenzene in 255 each chromatogram. Each tissue volatile concentration value was standardized to 'ug  $g^{-1}$  dry weight' by dividing by the sample dried weight. Since both the HWA and the EHS are quite small and adhere tightly to their twig or needle feeding

sites, complete removal of insects and their ovisacs from infested samples prior to analysis was not

practical. To test whether detected volatiles could potentially be of insect, rather than hemlock, origin, we obtained several samples of HWA-infested foliage of comparable size and insect density to our experimental samples, collected the insects, eggs, and the wax of ovisacs into vials, and extracted and analyzed the insect material using the plant-volatile protocol described herein. 

*Statistical Analysis.* Resin volatile concentrations were log transformed prior to statistical analysis to reduce heterogeneity of variance. Two-way mixed-model ANOVAs (Proc Mixed, SAS 9.3; SAS Institute

266 2011) were used to test twigs and needles separately for treatment-level differences in the concentration of

individual volatiles, total monoterpenoids, total sesquiterpenoids, total green leaf volatiles (needles only)

and total combined benzenoids (including phenolics; twigs only) with month (June vs. October) and

treatment (HWA, EHS and control) as fixed factors and tree as a random factor. Mixed-model analyses

were appropriate because we sampled from the same trees in both months. We also used ANOVA planned

contrasts to separately test HWA and EHS treatment means against the control mean, using treatment as the

fixed factor (R 2.14.0; R Development Core Team 2012).

Familywise error rate for the mixed model analyses was evaluated using a false discovery rate (FDR) estimation method ('fdrtool' software package; R 2.14.0; Strimmer 2008). FDR techniques are now used widely with multiple simultaneous hypothesis testing to estimate the proportion of tests with incorrectly rejected null hypotheses among tests with statistically significant findings. This is in contrast to traditional familywise error rate correction methods (e.g. the sequential Bonferroni) that estimate the probability of a false rejection among all tests conducted and, arguably, unnecessarily sacrifice statistical power.

As an additional measure of the overall strength of evidence for our mixed-model hypothesis test findings, we used the following binomial equation (sensu Moran 2003) to calculate the overall probability 282 of obtaining *K* tests with *P*-values smaller than our specified  $\alpha$ -level:

 $P_B = [N!/(N-K)!K!] \times \alpha^K (1-\alpha)^{N-K}$ 

284 where *N* = number of tests. This procedure allowed us to estimate the probability that so many statistically significant treatment effects could arise by chance (i.e. could be 'false positives').



- significance. HWA feeding significantly decreased total monoterpenoids, while EHS feeding decreased both total monoterpenoids and total volatiles with marginal significance. HWA feeding marginally increased total benzenoids, while EHS feeding marginally decreased these compounds (Online Resource 2A). In needles, (Online Resource 2B) EHS feeding increased *cis*-3-hexenal and total GLVs
- significantly, and increased *trans*-2-hexenal and the benzenoid p-cymene with marginal significance.
- There were no significant effects of HWA feeding on needle volatile levels.
- Results of planned contrast ANOVA comparisons of average control versus treatment volatile
- concentrations were quite similar to those we obtained using the mixed model analyses. Significance
- values from these simpler analyses are indicated in Table 1.
- The binomial probability that the 60 twig volatile mixed model tests we ran would generate *P*-
- 326 values smaller than the ones we observed was  $P_B=0.00014$  if calculated at the  $\alpha$ =0.05 level, or 4.7 x 10<sup>-9</sup> if
- 327 calculated at the  $\alpha$ =0.10 level. For the 54 needle volatile tests, the overall probability of no 'real' effect
- 328 was greater:  $P_B$ =0.18, and 0.12, respectively.
- Estimated false discovery rate for twig volatile hypothesis tests is reported as *q*-value alongside each test's nominal *P*-value (Online Resource 2A). The *q*-value is the minimum FDR level that would be needed to reject that hypothesis. Selection of an appropriate FDR level, in turn, depends on the proportion of false rejections considered tolerable. We did not report FDR for needle volatile hypothesis tests (Online Resource 2B). Since the method estimates the proportion of false rejections among only tests with significant findings—and there was just one out of 54 needle volatile hypothesis tests that was statistically significant—in that case an estimate of FDR was superfluous.
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#### DISCUSSION

- We found evidence of an induced response in eastern hemlock during infestation by both HWA and EHS ,
- encompassing a number of feeding-elicited changes in the tree's resin volatile profile. However, the
- modest induction (mostly decreases) of resin metabolites in twig tissue and the non-significant trend of
- modest increases in needle tissue produced by both insects, was conspicuously different from the profuse
- resinosis observed in insect-infested pines, spruces, and firs (Trapp and Croteau 2001). In light of the

considerable evidence that HWA induces more extensive changes in eastern hemlock physiology than does EHS (Miller-Pierce et al. 2010, Radville et al. 2011, Gonda-King et al. 2012, Gomez 2012), the observation that HWA and EHS produced similar overall changes in the tree's volatiles was intriguing and ran counter 346 to our predictions.

In contrast to the modest changes in terpenoid levels, a number of the non-terpenoids were sharply increased by HWA feeding, in what may reflect a hemlock defense response (Table 1). Benzyl alcohol was induced in HWA-infested trees; this compound is a common plant volatile (Dudareva et al. 2008) previously detected in the stem-wood of mountain hemlock (*T. mertensiana*; Shepherd et al. 2008) and in volatiles released from mite-infested spruce foliage (Kannaste 2008; Fig. 3). In screening studies, benzyl alcohol deterred feeding by the greenbug aphid *Schizaphis graminum*, reducing fecundity and causing substantial mortality (Formusoh et al. 1997). MeSA, which was also induced by HWA (Fig. 4), has been found in the volatile mix released after aphid feeding and identified as a deterrent to aphid settling and fecundity in a number of plant-insect systems (Hardie et al. 1994, Quiroz et al. 1998).

The sharp increase of these two compounds in HWA-infested trees (Table 1) is notable in light of the growing body of evidence that some plants respond to piercing-sucking hemiptera by activating biosynthetic pathways similar or identical to those used in pathogen defense (Kaloshian and Walling 2005). Benzyl alcohol is a strong antimicrobial agent against diverse microorganisms (Shenep et al. 2011), while MeSA, the volatile methyl ester of salicylic acid (SA), activates a SA-dependent biosynthetic cascade in numerous plants that leads to systemic acquired resistance (SAR) against pathogen infection (Durrant and Dong 2004). For aphids, close relatives of adelgids, feeding has been shown in many studies to activate the SA-dependent biosynthetic pathways normally associated with pathogen defense (Moran and Thompson 2001, Martinez de Ilarduya et al. 2003, Zhu-Salzman et al. 2004) or to induce pathogen-resistance outright in their host plant (Russo et al. 1997). The elevated levels of these two compounds in HWA-induced hemlock tissue is a sign that a SA-driven insect defense syndrome may be active in HWA-infested trees. It is also possible that increased production of these volatiles reflects the tree's detection of a microbial associate of HWA rather than of the insect itself. An endosymbiont was recently found throughout the body of the HWA and appears essential to the insect's survival (Shields and Hirth 2005). It

is possible that the hemlocks may be responding to this bacterium, if it is introduced into the vascular tissue of eastern hemlock during HWA feeding, by mobilizing a pathogen defense response.

Our results may help elucidate why HWA causes more extensive damage to eastern hemlock than EHS. Radville et al. (2011) detected evidence of a local hypersensitive response (elevated hydrogen peroxide levels) in both EHS- and HWA-infested trees, and showed that this hypersensitive response occurs systemically in response to HWA-infestation. The hypersensitive response usually precedes the development of SAR (Durrant and Dong 2004, Kaloshian and Walling 2005). Research on tobacco has revealed that in pathogen-infected plant tissue SA is enzymatically converted to the volatile MeSA, which acts as a mobile agent that is taken up by receptors on distant, uninfected tissue. There, the MeSA is demethylated and transformed back to SA, which in turn activates an induced resistance response to the invading organism (i.e. SAR; Shulaev et al. 1997, Park et al. 2007). Our discovery that MeSA levels were elevated in only the HWA-infested trees suggests this compound could be a mobile signal that propagates the 'pathogen-like' effects of the adelgid on uninfested foliage, extending the insect's effects and intensifying the overall damage to the tree. The observation that HWA elicited such a response, but EHS did not, may reflect the species-specific nature of the hemlock defense elicitors carried in the insects' salivary secretions, as has been observed in at least one other hemipteran-plant interaction (Ven et al. 2000).

The HWA-driven increases we observed in levels of benzyl alcohol and MeSA may also help explain previously noted changes in the primary chemistry of the hemlock saplings of the present study (Gomez et al. 2012). Although much of the biosynthetic pathway for the benzenoids has yet to be determined, radio-labeling experiments show they are derived from L-phenylalanine (Dudareva et al. 2006). As with benzyl alcohol and MeSA, a marked increase in L-phenylalanine and many other free amino acids occurred in trees infested with HWA, but not EHS (Gomez et al. 2012). Thus the increased amino acid levels in HWA-infested trees may constitute an adaptive mobilization of precursors of defense-related volatile compounds.

Alternatively, adelgid manipulation of host-plant biochemistry could explain a number of the insect-induced changes in resin chemistry we have shown. HWA, like many adelgids, forms extensive galls on the buds of its primary spruce host in its original range in Asia (Havill and Foottit 2007). Galling insects are known to be adept at manipulating host plant physiology to create a more nutritious and less defended environment (Tooker and Moraes 2009). We have demonstrated a substantial decrease in monoterpenoids, often compounds of direct defense against herbivory (Eyles et al. 2010, Schiestl 2010), in the tissue where the adelgid feed. Our results also show a less pronounced elicitation of GLVs (typical wounding response volatiles; Fig. 2; Shiojiri et al. 2006) in response to the feeding of HWA, relative to EHS, despite the adelgid's much greater impacts on tree physiology (Miller Pierce et al. 2010, Radville et al. 2011, Gonda-King et al. 2012). These observations, considered together with the noted increase in free amino acids only in HWA-infested trees (Gomez at al. 2012), may constitute evidence that the host-manipulating capacity conserved in adelgid biology may be an underlying mechanism in this system. Lagalante et al. (2006) suggested that the lack of a co-evolutionary history between eastern hemlock and sessile piercing-sucking insects resulted in the absence of biosynthetic pathways with which eastern hemlock can defend against insects like HWA and EHS. This hypothesis is consistent with the finding of little or no output of anatomical of chemical resin defenses. However, we did observe a resin chemical response to HWA, and to the co-occurring EHS, though perhaps of a subtler nature than that often seen in other conifers. It is possible that the resistance traits HWA elicits in eastern hemlock are simply not well matched to the actual challenge of this introduced insect and do not confer resistance. A comparison of the induced response of susceptible eastern hemlocks to those of HWA-resistant *Tsuga* species and strains of eastern hemlock believed resistant to HWA, as well as conifers with putative resistance to EHS (McClure and Fergione 1977), will test these hypotheses. Nonetheless, our findings establish that HWA and EHS both induce changes in the resin chemistry of eastern hemlock, and constitute the first critical step toward understanding the role inducible chemical defenses play in determining hemlock susceptibility to these exotic hemipteran pests. ACKNOWLEDGMENTS This work was supported by USDA Forest Service cooperative agreements #103-CA-11244225- 130 and

#09-CA-11420004-360 to JSE, and NSF grant DEB-0715504 and NIFA grant #2011-67013-30142 to ELP.

We also gratefully acknowledge the conceptual and technical help of R. D'Alonzo, S. Eyles, J. Gershenzon,





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### 591 FIGURES AND TABLES

592 **Table 1** Resin volatile concentration relative change (treatment average/control average ratio) for eastern

593 hemlock saplings treated with 3-year artificial infestation with hemlock woolly adelgid (HWA) or elongate

594 hemlock scale (EHS)



<sup>a</sup> Foliage sampled in June was mature, previous year growth infested with EHS or progredien-generation HWA; foliage sampled in October was young, current year growth infested with EHS or sisten-generation HWA.

**b** Compounds are ordered first by structural class, then by ascending order of elution from a non-polar DB-5 GC column. A summed total for each class of phytochemical is included.

 $c$  Values >1 and <1 indicate an increase (dark gray shading) and decrease (light gray shading), respectively, from the control trees. Statistically significant differences from uninfested trees (planned contrast,  $P<0.05$ ) are marked in bold text with asterisks. Marginally significant differences from uninfested trees (planned contrast,

<sup>d</sup> Tentative identification based on comparison of GC retention time and mass spectrum with those of an authentic standard

595

- **Fig. 1** (A) GC-FID total ion chromatogram showing volatiles tentatively identified in HWA-infested
- eastern hemlock needles: **1**, *cis*-3-hexenal; **2**, *n*-hexanal; **3**, *trans*-2-hexenal; **4**, tricyclene; **5**, α-pinene; **6**,
- camphene; **7**, sabinene; **8**, β-pinene; **9**, myrcene; **10**, α-phellandrene; **11**, isobutylbenzene (internal
- standard); **12**, *p*-cymene, **13**, D-limonene; **14**, γ-terpinene, **15**, terpinolene; **16**, camphor; **17**, borneol; **18**, 4-
- carvomenthenol; **19**, *p*-menth-1-en-9-ol; **20**, α-terpineol; **21**, *trans*-piperitol; **22**, piperitone; **23**, bornyl
- acetate; **24**, β-caryophyllene; **25**, α-humulene; **26**, germacrene-D; **27**, α-amorphene; **28**, δ-cadinene. (B)
- 603 GC-MS total ion chromatogram showing volatiles in HWA-infested twigs: 1, tricyclene; 2,  $\alpha$ -pinene; 3,
- camphene; **4**, β-pinene; **5**, myrcene; **6**, isobutylbenzene (internal standard); **7**, *p*-cymene; **8**, D-limonene; **9**,
- benzyl alcohol; **10**, unknown; **11**, unknown; **12**, α-campholenal; **13**, L-*trans*-pinocarveol; **14**, *cis*-verbenol;
- **15**, *trans*-verbenol; **16**, pinocarvone; **17**, borneol; **18**, *p*-cymen-8-ol; **19**, methyl salicylate; **20**, myrtenol; **21**,
- verbenone; **22**, *cis*-carveol; **23**, bornyl acetate; **24**, unknown; **25**, β-caryophyllene; **26**, 3,4-
- dimethoxyphenol; **27**, α-humulene; **28**, germacrene-D; **29**, α-amorphene; **30**, raspberry ketone; **31**,
- caryophyllene dioxide.
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- **Fig. 2** Green leaf volatile ('GLV') content (average ± SE) in needle tissue of control and insect-infested
- eastern hemlocks. 'HWA' or 'EHS' represents 3-year artificial infestation with hemlock woolly adelgid (*A.*
- *f* tsugae) or elongate hemlock scale (*F. externa*). Data represents the average concentration ( $\mu$ g⋅g dry wt<sup>-1</sup>) of
- total GLVs in mature previous year growth (sampled 28 June) and young current year growth (sampled 19
- October), calculated from 6 to 9 trees per treatment group. *P*-values are shown when the difference
- between the treatment and control trees was significant (*P*<0.05), or marginally significant (0.05<*P*<0.10;
- planned contrast).
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 **Fig. 3** Benzyl alcohol content (average  $\pm$  SE) in twig tissue of control and insect-infested eastern hemlock trees. 'HWA' or 'EHS' represents 3-year artificial infestation with hemlock woolly adelgid (*A. tsugae*) or 621 elongate hemlock scale (*F. externa*). Data represents the average concentration ( $\mu$ g⋅g dry wt<sup>-1</sup>) of benzyl alcohol in mature previous year growth (sampled 28 June) and young current year growth (sampled 19 October), calculated from 7 to 9 trees per treatment group. *P*-values are shown when the difference



- *tsugae*) or elongate hemlock scale (*F. externa*). Data represents the average concentration (μg⋅g dry wt<sup>-1</sup>) of
- methyl salicylate in mature previous year growth (sampled 28 June) and young current year growth
- (sampled 19 October) calculated from 7 to 9 trees per treatment group. *P*-values are shown when the
- difference between the treatment and control trees was significant (*P*<0.05), or marginally significant
- (0.05<*P*<0.10; planned contrast).
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**Fig. 1** 











**Online Resource 1** Structures of representative volatile compounds of eastern hemlock (*Tsuga canadensis* Carr.): (A) monoterpenoids, (B) sesquiterpenoids, (C) green leaf volatiles (GLVs) and (D) benzenoids detected in twig or needle tissue of young trees.

	Previous year growth (June sampling)			Current year growth (October sampling)					HWA versus control analyses	EHS versus control analyses		
							Sample	Insect		Sample	Insect	
	Control HWA		EHS	Control HWA		EHS	Date	Effect	q-value	Date	Effect	q-value
	$(N=8)$	$(N=9)$	$(N=7)$	$(N=8)$	$(N=9)$	$(N=7)$	F(P)	F(P)	$(FDR)^d$	F(P)	F(P)	(FDR)
Tricyclene	15.55	7.70	12.49	26.32	16.45	25.04	$41.56^{b}$	7.46		37.46	0.50	
α-Pinene	(2.44) 29.48	(1.29) 22.96	(3.50) 27.45	(3.55) 345.86	(2.72) 372.80	(4.10) 245.02	(0.00) 163.97	(0.01) 1.10	0.05	(0.00) 218.00	(0.49) 1.02	0.24
Camphene	(4.27) 15.43	(5.81)	(4.48) 11.75	(90.76)	(127.32) 22.71	(51.46) 30.47	(0.00) 70.93	(0.31) 11.09	0.18	(0.00) 73.19	(0.33) 1.19	0.19
	(2.06)	7.53 (1.24)	(2.87)	33.34 (3.65)	(3.85)	(4.50)	(0.00)	(0.00)	0.03	(0.00)	(0.29)	0.18
β-Pinene	2.74	2.09	2.94	35.65	43.59	26.84	145.38	0.50		205.05	0.29	
Myrcene	(0.36) 3.31	(0.53) 2.78	(0.40) 4.30	(10.07) 31.51	(15.50) 44.46	(5.92) 51.46	(0.00) 61.94	(0.49) 0.32	0.24	(0.00) 91.03	(0.60) 0.72	0.28
	(0.90)	(0.39)	(1.11)	(4.13)	(19.41)	(17.22)	(0.00)	(0.58)	0.28	(0.00)	(0.41)	0.22
Limonene	0.30 (0.18)	0.22 (0.11)	0.29 (0.20)	26.99	18.59 (4.38)	39.72	155.13 (0.00)	2.83 (0.11)		829.59 (0.00)	1.78 (0.20)	
α-Campholenal	5.37	3.15	4.01	(2.13) 14.59	9.99	(5.49) 8.97	64.95	5.63	0.11	53.25	4.38	0.15
	(0.89)	(0.73)	(0.48)	(2.53)	(1.52)	(0.88)	(0.00)	(0.03)	0.07	(0.00)	(0.06)	0.09
L-trans-Pinocarveol	3.31 (0.51)	2.44 (0.61)	2.90 (0.56)	11.08 (1.89)	7.87 (1.57)	6.47 (1.01)	63.33 (0.00)	4.04 (0.06)	0.09	34.49 (0.00)	4.48 (0.05)	0.09
cis-Verbenol	0.27	0.25	0.31	1.12	0.82	0.51	28.27	1.37		8.36	3.00	
	(0.09)	(0.08)	(0.15)	(0.15)	(0.17)	(0.25)	(0.00)	(0.26)	0.17	(0.01)	(0.11)	0.11
trans-Verbenol	4.76 (0.65)	3.65 (0.98)	4.45 (0.73)	21.63 (3.74)	15.58 (2.91)	12.45 (2.28)	70.76 (0.00)	3.46 (0.08)	0.10	58.47 (0.00)	3.56 (0.08)	0.10
Pinocarvone	3.62	2.47	3.40	9.66	7.43	5.45	69.94	3.42		26.51	4.22	
	(0.55)	(0.50)	(0.42)	(1.59)	(1.22)	(0.83)	(0.00)	(0.08)	0.10	(0.00)	(0.06)	0.09
Borneol	6.62 (1.37)	4.82 (1.28)	5.78 (1.11)	7.77 (1.00)	5.63 (0.86)	9.63 (2.41)	4.31 (0.05)	2.29 (0.15)	0.12	4.90 (0.04)	0.01 (0.93)	0.38
Myrtenol	5.09	3.12	3.17	22.18	15.59	12.46	137.21	4.61		50.32	8.48	
	(0.56)	(0.79)	(0.92)	(4.16)	(3.01)	(1.99)	(0.00)	(0.05)	0.09	(0.00)	(0.01)	0.05
Verbenone	9.74 (1.75)	5.85 (1.26)	6.87 (1.36)	15.40 (2.60)	11.49 (1.84)	10.06 (1.64)	14.32 (0.00)	3.50 (0.08)	0.10	7.82 (0.01)	3.02 (0.10)	0.11
trans-Carveol	0.72	0.52	0.63	4.77	3.45	2.98	84.00	1.14		76.51	1.10	
Bornyl Acetate	(0.29)	(0.21)	(0.22)	(0.84)	(0.81)	(0.39)	(0.00)	(0.30)	0.18	(0.00)	(0.31)	0.18
	32.87 (4.62)	14.86 (2.98)	19.63 (5.52)	78.83 (7.51)	56.32 (13.02)	64.61 (10.93)	86.15 (0.00)	9.76 (0.01)	0.04	74.67 (0.00)	3.69 (0.08)	0.10
β-Caryophyllene	0.50	0.52	0.00	2.88	6.85	3.61	12.20	0.28		11.83	0.65	
	(0.28)	(0.26)	(0.00)	(0.98)	(3.30)	(1.79)	(0.00)	(0.61)	0.28	(0.00)	(0.43)	0.23
α-Humulene	0.45 (0.24)	0.85 (0.34)	1.68 (0.57)	7.32 (2.57)	20.56 (10.00)	10.71 (3.80)	29.39 (0.00)	0.57 (0.46)	0.23	45.48 (0.00)	2.20 (0.16)	0.13
Germacrene-D	0.14	0.14	0.18	1.22	1.43	0.67	43.44	0.07		16.44	1.12	
	(0.09)	(0.10)	(0.14)	(0.28)	(0.37)	(0.32)	(0.00)	(0.79)	0.34	(0.00)	(0.31)	0.18
α-Amorphene	0.43 (0.39)	1.64 (1.14)	0.45 (0.45)	4.71 (2.90)	15.25 (7.88)	5.42 (4.01)	10.38 (0.01)	1.92 (0.18)	0.14	8.71 (0.01)	0.05 (0.83)	0.35
Caryophyllene Oxide	4.02	3.82	3.41	16.89	14.87	8.48	57.84	0.20		33.64	2.41	
	(0.79)	(0.86)	(0.96)	(3.93)	(3.81)	(1.29)	(0.00)	(0.66) 0.02	0.30	(0.00)	(0.14)	0.12
p-Cymene	4.91 (1.29)	3.69 (1.17)	3.55 (0.77)	26.64 (9.25)	33.32 (9.91)	19.54 (7.38)	87.06 (0.00)	(0.90)	0.37	55.30 (0.00)	0.61 (0.45)	0.23
Benzyl Alcohol	0.79	24.81	0.42	1.22	11.83	11.83	1.57	12.56		5.95	0.99	
p-Cymen-8-ol	(0.33) 3.72	(10.70) 3.46	(0.42) 1.86	(0.24) 8.81	(3.94) 8.11	(3.94) 6.28	(0.23) 28.28	(0.00) 0.00	0.03	(0.03) 15.41	(0.34) 2.34	0.19
	(1.01)	(0.82)	(0.95)	(1.99)	(1.56)	(1.86)	(0.00)	(0.99)	0.39	(0.00)	(0.15)	0.12
Methyl Salicylate	0.02	2.91	0.00	0.08	0.96	0.09	3.69	8.03		2.34	0.10	
3,4-Dimethoxyphenol	(0.02) 2.83	(1.72) 2.48	(0.00) 2.80	(0.04) 5.15	(0.45) 5.84	(0.09) 4.45	(0.07) 16.37	(0.01) 0.00	0.05	(0.15) 4.20	(0.75) 0.39	0.33
Raspberry Ketone	(0.41)	(0.44)	(0.52)	(0.76)	(0.83)	(0.92)	(0.00)	(0.96)	0.39	(0.06)	(0.54)	0.26
	4.59	3.34	3.33	16.46	10.79	10.53	67.80	4.27		21.20	3.02	
Unknown A	(0.62) 0.82	(0.59) 0.71	(0.70) 0.68	(2.28) 3.63	(1.70) 2.86	(3.30) 1.83	(0.00) 35.78	(0.06) 1.05	0.09	(0.00) 44.52	(0.10) 1.92	0.11
Unknown B	(0.35)	(0.22)	(0.50)	(0.52)	(0.45)	(0.62)	(0.00)	(0.32)	0.19	(0.00)	(0.19)	0.14
	3.03	1.95	2.43	22.35	15.03	13.24	488.65	6.06		316.95	2.56	
Unknown C	(0.44) 16.20	(0.44) 4.78	(0.67) 9.99	(4.30) 14.40	(3.92) 4.22	(1.72) 6.97	(0.00) 2.06	(0.03) 15.01	0.07	(0.00) 6.35	(0.13) 2.40	0.12
	(2.77)	(0.70)	(1.79)	(4.10)	(0.67)	(1.30)	(0.17)	(0.00)	0.02	(0.02)	(0.14)	0.12
<b>Total Monoterpenoids</b>	139.16	84.40	110.36	686.71	652.75	552.14	103.68	5.60		118.05	3.14	
<b>Total Sesquiterpenoids</b>	(12.82) 5.53	(16.83) 6.98	(18.05) 5.71	(117.13) 33.04	(190.09) 58.96	(88.28) 28.88	(0.00) 35.13	(0.02) 0.44		(0.00) 41.86	(0.09) 0.16	
	(1.31)	(1.87)	(1.65)	(10.17)	(23.51)	(8.88)	(0.00)	(0.52)		(0.00)	(0.69)	
Total Benzenoids/Phenolics	16.87	40.69	11.94	58.36	70.84	41.94	33.17	3.07		46.06	3.35	
<b>Total Twig Resin Volatiles</b>	(2.07) 181.60	(12.45) 139.50	(2.26) 141.10	(12.67) 818.49	(12.09) 804.66	(11.62) 645.00	(0.00) 82.54	(0.10) 2.36		(0.00) 119.96	(0.09) 3.92	
	(15.13)	(22.67)	(20.22)	(141.03)	$(228.07)$ $(103.38)$		(0.00)	(0.14)		(0.00)	(0.06)	

**Online Resource 2A** Twig volatile concentrations (average  $\pm$  SE) with mixed model ANOVA results

"Familywise error rate was estimated using a false discovery rate (FDR) method. FDR estimates the proportion of incorrectly rejected null hypotheses among tests with statistically significant findings. The q-value for each

 $<sup>b</sup>$  There were no significant sampling date x insect treatment interactions, except as noted.</sup>

 $\textdegree$  The HWA versus control model includes a significant sampling date x treatment interaction term; F=1.83, P=0.018.

	Previous year growth (June sampling)				Current year growth							
				(October sampling)			HWA versus control analyses			EHS versus control analyses		
							Sample Insect			Sample Insect		
	Control HWA		EHS	Control HWA		<b>EHS</b>	Date	Effect	q-value	Date	Effect	q-value
	$(N=8)$	$(N=9)$	$(N=7)$	$(N=8)$	$(N=9)$	$(N=7)$	F(P)	F(P)	(FDR)	F(P)	F(P)	(FDR)
cis-3-Hexenal	1.21	1.26	1.66	1.59	2.07	4.04	4.85	$\overline{1.13}$	$*^a$	6.45	5.18	$\ast$
n-Hexanal	(0.64) 0.54	(0.31) 0.58	(0.48) 0.76	(0.31) 13.43	(0.31) 17.13	(1.13) 20.87	(0.05) 174.51	(0.31) 0.81		(0.03) 118.68	(0.04) 1.62	
trans-2-Hexenal	(0.15) 3.27	(0.25) 4.26	(0.31) 3.58	(2.80) 4.81	(1.19) 8.76	(4.32) 12.25	(0.00) 6.23	(0.39) 2.78		(0.00) 14.47	(0.23) 3.29	
	(1.31)	(0.92)	(0.58)	(0.79)	(2.03)	(3.03)	(0.03)	(0.12)	×	(0.00)	(0.10)	$\ast$
Tricyclene	26.39 (2.29)	28.26 (2.35)	26.88 (4.66)	50.67 (3.01)	56.86 (4.61)	55-33 (8.66)	153.14 (0.00)	0.64 (0.44)		73.17 (0.00)	0.09 (0.77)	
α-Pinene	91.56	98.81	96.22	186.27	213.45	202.57	177.07	0.63		63.48	0.00	
Camphene	(8.08) 60.41	(9.21) 62.60	(14.25) 63.14	(13.69) 115.01	(20.86) 123.94	(26.49) 129.77	(0.00) 148.38	(0.44) 0.19		(0.00) 65.98	(0.97) 0.00	
	(5.13)	(5.52)	(9.42)	(6.74)	(10.52)	(16.85)	(0.00)	(0.67)		(0.00)	(0.95)	
Sabinene	3.60	4.08	3.54	7.06	8.40	6.40	114.12	1.50		38.00	0.37	
	(0.35)	(0.38)	(0.57)	(0.82)	(0.77)	(0.86)	(0.00)	(0.24)	÷	(0.00)	(0.56)	
β-Pinene	12.92 (1.39)	14.11 (1.42)	12.91 (2.00)	29.00 (2.80)	33.21 (2.50)	29.00 (3.78)	215.16 (0.00)	0.91 (0.36)		70.42 (0.00)	0.05 (0.82)	
Myrcene	15.26	16.05	15.57	29.20	31.65	31.40	139.98	0.33		63.15	0.01	
	(1.23)	(1.52)	(2.27)	(1.69)	(2.48)	(3.68)	(0.00)	(0.57)		(0.00)	(0.91)	×
α-Phellandrene	6.01	7.11	6.74	17.87	23.87	19.53	145.20	1.03		75.42	0.23	
	(1.08)	(1.31)	(1.19)	(2.45)	(2.07)	(3.06)	(0.00)	(0.33)		(0.00)	(0.64)	
Limonene	25.18	29.89	29.08	58.90	66.21	63.23	31.97	1.55		23.70	0.39	
y-Terpinene	(4.00) 0.48	(2.76) 0.42	(4.76) 0.39	(3.99) 0.99	(5.61) 1.13	(7.40) 0.91	(0.00)	(0.24) 0.02		(0.00)	(0.55) 0.72	
	(0.08)	(0.11)	(0.15)	(0.05)	(0.12)	(0.20)	50.44 (0.00)	(0.89)		29.47 (0.00)	(0.41)	
Terpinolene	2.03	2.10	2.33	4.75	6.01	6.07	130.28	0.70		77.55	0.81	
	(0.18)	(0.26)	(0.37)	(0.30)	(0.84)	(0.82)	(0.00)	(0.42)		(0.00)	(0.39)	
Camphor	0.60	0.62	1.06	1.84	1.35	2.58	22.70	0.30		14.11	0.79	
	(0.15)	(0.32)	(0.29)	(0.44)	(0.69)	(1.07)	(0.00)	(0.59)	÷	(0.00)	(0.39)	$\ast$
Borneol	1.52	0.08	0.16	0.09	0.21	0.15	0.28	0.62		0.71	0.33	
	(1.43)	(0.08)	(0.10)	(0.09)	(0.14)	(0.10)	(0.61)	(0.45)		(0.42)	(0.58)	
4-Carvomenthenol	0.87 (0.11)	0.72 (0.11)	0.71 (0.15)	1.50 (0.17)	1.39 (0.18)	1.37 (0.31)	31.05 (0.00)	0.64 (0.44)		17.99 (0.00)	0.56	×
p-Menth-1-en-ol	1.72	1.44	1.27	2.73	3.13	2.94	65.17	0.36		61.76	(0.47) 1.20	
	(0.22)	(0.31)	(0.35)	(0.20)	(0.70)	(0.54)	(0.00)	(0.56)	$\ast$	(0.00)	(0.30)	
a-terpineol	0.79	0.93	0.90	1.21	0.59	0.94	0.31	0.60		0.10	0.03	
	(0.06)	(0.08)	(0.04)	(0.48)	(0.11)	(0.39)	(0.59)	(0.45)		(0.76)	(0.86)	
Piperitol	0.88	0.62	0.52	3.10	3.86	3.67	92.22	0.02		56.01	0.50	
	(0.08)	(0.14)	(0.19)	(0.62)	(0.42)	(0.70)	(0.00)	(0.90)	*	(0.00)	(0.49)	
Piperitone	24.20	20.02	21.16	37.11	39.23	47.58	37.05	0.01		29.88	0.01	
<b>Bornyl Acetate</b>	(4.60)	(2.30) 126.13	(4.72)	(4.67) 212.80	(4.83)	(6.37)	(0.00)	(0.91) 0.07		(0.00)	(0.94) 0.01	
	125.05 (10.85)	(11.00)	130.54 (18.86)	(16.12)	229.55 (27.16)	235.76 (29.51)	92.52 (0.00)	(0.79)	$\ast$	48.47 (0.00)	(0.91)	$\star$
β-Caryophyllene	11.85	11.99	12.67	22.22	27.53	25.15	108.43	1.11		66.54	0.01	
	(0.91)	(0.72)	(2.12)	(2.09)	(2.66)	(3.69)	(0.00)	(0.31)	$\star$	(0.00)	(0.92)	$\star$
α-Humulene	13.51	14.16	15.12	26.54	32.89	30.09	121.09	1.42		70.01	0.05	
	(1.20)	(0.91)	(2.55)	(2.54)	(3.21)	(4.43)	(0.00)	(0.26)		(0.00)	(0.82)	
Germacrene D	4.61	5.79	3.28	7.64	13.45	5.24	47.97	1.32		31.87	1.80	
	(0.91)	(1.19)	(1.12)	(2.38)	(2.99)	(2.06)	(0.00)	(0.27)		(0.00)	(0.21)	
α-Amorphene	2.29 (0.22)	2.09 (0.22)	2.30 (0.34)	4.10 (0.53)	4.73 (0.69)	4.46 (0.64)	61.99 (0.00)	0.01 (0.91)		66.56 (0.00)	0.05 (0.83)	
δ-Cadinene	2.34	2.28	2.40	4.16	5.14	4.72	58.84	0.26		60.82	0.06	
	(0.19)	(0.24)	(0.41)	(0.58)	(0.70)	(0.76)	(0.00)	(0.62)		(0.00)	(0.81)	$\star$
p-Cymene	5.76	1.92	1.46	3.66	2.71	2.31	0.87	1.83		0.84	3.66	
	(3.66)	(0.28)	(0.27)	(0.79)	(0.26)	(0.35)	(0.37)	(0.20)	$\star$	(0.38)	(0.08)	$\star$
Total Green Leaf Volatiles	5.02	6.10	5.99	19.83	27.96	37.16	62.45	3.04		85.22	6.02	
	(1.89)	(1.23)	(0.81)	(2.84)	(3.11)	(4.49)	(0.00)	(0.11)	$\star$	(0.00)	(0.03)	$\star$
Total Monoterpenoids	399.47	413.99	413.11	760.09	844.04	839.20	22.89	1.12		18.00	0.40	
	(28.46)	(34.62)	(59.97)	(46.43)	(78.55)	(102.74)	(0.00)	(0.31)	×	(0.00)	(0.54)	
<b>Total Sesquiterpenoids</b>	34.60	36.31	35.76	64.66 (7.03)	83.75	69.65	32.83 (0.00)	1.79 (0.21)	*	20.64 (0.00)	0.16 (0.69)	$\ast$
Total Needle Volatiles	(2.66) 444.84	(1.87) 458.33	(5.67) 456.32	848.23	(9.16) 958.46	(9.99) 948.32	183.62	0.44		75.63	0.00	
	(27.90)	(35.81)	(65.18)	(51.45)	(84.71)	(112.67)	(0.00)	(0.52)	÷	(0.00)	(0.99)	$\star$

**Online Resource 2B** Needle volatile concentrations (average  $\pm$  SE) with mixed model ANOVA results

 $\overline{P}$  FDR results were not reported for needle volatile tests due to the scarcity of statistically significant findings. See text for a more detailed explanation.