Manuka Oil and Phoebe Oil are Attractive Baits for Xyleborus glabratus (Coleoptera: Scolytinae), the Vector of Laurel Wilt

JAMES L. HANULA1 AND BRIAN SULLIVAN2


ABSTRACT Redbay ambrosia beetle, Xyleborus glabratus Eichhoff, is a native of Southeast Asia recently established in coastal forests of Georgia, SC and Florida. It vectors a wilt fungus, Raffaelea sp., lethal to redbay trees, Persea borbonia L. Spreng, and certain other Lauraceae. No practical monitoring system exists for this beetle so we conducted studies to identify host attractants and develop lures. Volatiles were collected from redbay wood and bark by steam distillation, direct solvent extraction, and dynamic headspace sampling with a Poropak Q cartridge. Steam, methanol, and pentane extracts were tested as baits in trapping trials but were not attractive to X. glabratus. Major constituents in Poropak aeration identified by gas chromatography-mass spectrometry included α-pinene, β-pinene, δ-3-carene, eucalyptol, p-cymene, α-copaene, terpinene-4-ol, linalool, calamenene, and nonanoic acid. We assayed several of these compounds (including eucalyptol, p-cymene, terpinene-4-ol, linalool, nonanoic acid, and caraphyllene oxide) both individually and in combination, but none were attractive at tested doses. Two other redbay odor components, α-copaene and calamenene, were unavailable in sufficient quantities commercially so we substituted manuka oil, the essential oil extracted from Leptospermum scoparium Forst. and Forst., which contains high proportions of both compounds. Manuka oil was equally attractive as redbay wood to X. glabratus, but increasing release rates >10-fold did not enhance its activity. Phoebe oil, an extract of Brazilian walnut (Phoebe porosa Mez.), which contains significant quantities of α-copaene and calamenene, was also attractive. Fractions of manuka oil were not more attractive than the whole oil. Manuka and phoebe oil are readily available and are good alternatives to redbay wood as a trap bait for monitoring X. glabratus distribution and population trends.

KEY WORDS exotic species, α-copaene, calamenene

Redbay (Persea borbonia L. Spreng.) are evergreen trees in the family Lauraceae indigenous to Atlantic and Gulf coastal plain forests from Virginia to Texas. Although most often occurring as midstory trees or understory shrubs, mature redbay trees can grow to 18–21 m with diameters of 60–90 cm (Brendemuehl 1990). Coastal forests of South Carolina and Georgia contain an average of 200–400 redbay trees (2.5 cm or larger) per hectare (Hanula et al. 2008).

In 2004, redbay trees in the Savannah, GA, area started dying from what was initially thought to be salt intrusion after a storm. Subsequent investigation showed the presence of an exotic ambrosia beetle and a fungal pathogen. Fraedrich et al. (2008) showed that this beetle, Xyleborus glabratus Eichhoff, carries a Raffaelea sp. fungus in its mandibular mycangiium that causes a disease syndrome named "laurel wilt." The beetles are capable of inoculating apparently healthy trees with this fungus and are its only known vector. Additionally, Fraedrich et al. (2008) reported that the beetle and its associated wilt fungus killed all redbay trees over 2.5 cm diameter within 2 yr of first detection in a stand. Since the initial reports of tree mortality in 2004, the beetle has spread north to Charleston County, SC, and south to Brunswick, GA. A second introduction in the Jacksonville, FL, area and subsequent spread has resulted in a continuous distribution of the redbay ambrosia beetle from south of Charleston, SC, to Putnam County, FL (USDA Forest Service 2008). An additional, isolated population exists in Broward and Indian River Counties, FL. It is unclear whether the rapid spread of this beetle has been caused by natural dispersal alone or has been aided by human movement of infested wood.

Although heavy tree mortality was first reported in 2004, beetles were initially detected in ethanol-baited traps at Port Wentworth, GA, 2 yr earlier (Rabaglia et al. 2006). Unlike most ambrosia beetles, redbay ambrosia beetle is weakly or not attracted to ethanol (D. Miller, personal communication), suggesting that populations in the vicinity of the traps may have been high as early as 2002. Anecdotal reports suggest that tree mortality was occurring before 1999 (C. Bates, personal communication).

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Given its already widespread establishment, eradication of *X. glabratus* is not feasible. Furthermore, conventional suppression efforts based on sanitation cutting are unlikely to succeed because of the speed of range expansion by this species, the rapid tree mortality in infested areas, the ability of a single beetle to inoculate and kill a tree, and the high density of suitable hosts. Now that the virulence of this pest is known, detection of *X. glabratus* at points of entry should be a priority. However, the poor performance of standard ethanol-baited ambrosia beetle traps (D. R. Miller, personal communication) suggests that better attractants are needed for detecting and delimiting populations.

Wounded or cut redbay are attractive to *X. glabratus*, but presence of the fungal symbiont or beetles themselves in the wood did not contribute to attraction (Hanula et al. 2008). Currently, cut sections from healthy or diseased redbay trees are the best baits available for trapping this insect (Hanula et al. 2008). We report here a series of experiments to identify and evaluate attractants for the redbay ambrosia beetle.

### Materials and Methods

Field studies were conducted during September and October 2006 and from June to October 2007 at Hunting Island State Park (Beaufort County, SC) located on a small barrier island in the southeastern corner of the state. The forest, located between the dunes and coastal marsh, consisted of a dominant overstory of loblolly pine (*Pinus taeda* L.); a midstory of redbay, live oak (*Quercus virginiana* Nutt.); cabbage palm trees [*Sabal palmetto* (L.) Lodd. ex J.A. and J.H. Schultes]; a shrub layer of waxmyrtle (*Morella cerifera* L. Small) and redbay; and an understory dominated by cabbage palm seedlings. In August 2007, the island had an average of 44 (SE = 6.3) live redbay trees/ha 2.5 cm diameter or larger and 216 (SE = 23.3) dead trees/ha in the same size range that were recently killed, i.e., they still had some dead leaves attached (Hanula et al. 2008).

**Collection of Redbay Volatiles.** A Cleveger apparatus was used to extract lighter-than-water essential oils from redbay tissue. Freshly cut redbay wood with bark was chipped in an 8-horsepower chipper/vac (Troy-Bilt, Jefferson, WI). Sections of wood too large to fit in the chipper were split before chipping. We placed 500 g of fresh chips in a 2-liter Erlenmeyer flask with 1 liter of distilled water and attached the flask to the Cleveger apparatus after the water began to boil. Only a few microliters of oil were collected during extensive boiling of the wood chips (~6 h), so we also collected the distilled water fraction from the Cleveger apparatus. This fraction, which had a strong odor and was available in sufficient quantities, was assayed for its attractiveness to *X. glabratus* instead of the oil. We also extracted 500 g of fresh wood chips with 500 ml of either methanol or pentane. The wood chips were steeped in the solvent for 1 h at room temperature, filtered, and concentrated to 50 ml in a rotary evaporator.

**Behavioral Bioassays.** Responses of *X. glabratus* to candidate attractive substances were assayed in trapping trials. Each trial was replicated four times. Trial 1 used flight-intercept traps that consisted of a single panel (20 by 30 cm) of clear Plexiglas centered above a white, 2-liter capacity plastic bucket partially filled with low-toxicity antifreeze (propylene glycol). These traps were hung ~2 m above the ground on ropes tied between two nonhost trees with bait vials suspended near the middle of the panel. Because it was unknown whether bole-like visual stimuli were necessary for beetles to orient to an odor source, we provided a dark silhouette by hanging a sleeve of black cloth (12 cm wide and 40 cm long) against one side of the Plexiglas barrier, opposite the baits. For trials 2–6, we used sticky traps rather than panel traps because of their similar effectiveness and cost but greater convenience. Sticky traps were constructed from solitary, white, wing-style trap bottoms (23 by 28 cm; Scentry Biologicals, Billings, MT) secured flat with binder clips against a Plexiglas panel (20 by 30 cm). A single trap consisted of two such panels suspended sticky-side-out at 2 m height from opposite sides of a wooden stake. A large binder clip secured both panels to the stake and prevented movement by the wind. Bait vials were suspended near the center of one of the panels. For all trials, traps were ~10 m apart within complete blocks, and blocks were separated by >50 m. Traps were at least 5 m from the nearest host tree. Two blocks were located on both the north and south ends of the island.

**Trapping Trial 1.** Attractiveness of Cleveger, methanol, and pentane extracts of redbay tissue were compared with a methanol blank which was used as a control because alcohols are attractive to some ambrosia beetles. Baits consisted of a piece of cotton dental wick (1 cm diameter; 4 cm long) inside a 25-ml capacity glass scintillation vial filled with 5 ml of the test substance. A single 3-mm-diameter hole was drilled through the vial cap to allow volatiles to escape. Beetles responding to the baits were collected semi-weekly from 21 September to 15 October 2006.

**Trapping Trial 2.** We compared attractiveness of a methanol-only control to seven compounds (Table 1) identified in volatile emissions of redbay tissue (see below), presented either singly or in combination. We also tested manuka oil (Coast Biologicals, Bombay, South Auckland, New Zealand), an essential oil extracted from wild manuka (*Leptospermum scoparium* Forst. and Forst.) plants, because it contained relatively large quantities of two major volatiles (Porter and Wilkins 1999), copaene and calamene, that we had identified in redbay but were not readily available commercially. Liquid test compounds and the manuka oil were absorbed (100 μl) onto one half of a cotton dental wick (1 cm diameter; 2 cm long) placed inside an uncapped 25-ml capacity scintillation vial. Caryophyllene oxide (solid at room temperature) was dissolved in methanol (50 mg/ml) and 100 μl was applied (i.e., 5 mg caryophyllene oxide) to the wick. Catch was collected weekly from 17 to 31 July 2007.
Table 1. Compounds identified in aeration of chipped bark and sapwood of redbay

<table>
<thead>
<tr>
<th>No.</th>
<th>Retention time (min)</th>
<th>Compound name</th>
<th>CAS no.</th>
<th>Compound class</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>6.65</td>
<td>α-Pinene</td>
<td>80-56-8</td>
<td>Monoterpene hydrocarbon</td>
</tr>
<tr>
<td>2</td>
<td>7.84</td>
<td>β-Pinene</td>
<td>127-91-3</td>
<td>Monoterpene hydrocarbon</td>
</tr>
<tr>
<td>3</td>
<td>8.41</td>
<td>δ-3-Carene</td>
<td>13466-78-9</td>
<td>Monoterpene hydrocarbon</td>
</tr>
<tr>
<td>4</td>
<td>9.40</td>
<td>Eucalyptol</td>
<td>470-82-6</td>
<td>Monoterpene ether</td>
</tr>
<tr>
<td>5</td>
<td>10.40</td>
<td>p-Cymene</td>
<td>99-87-6</td>
<td>Monoterpene hydrocarbon</td>
</tr>
<tr>
<td>6</td>
<td>13.67</td>
<td>α-Cubebene</td>
<td>17699-14-8</td>
<td>Sesquiterpene hydrocarbon</td>
</tr>
<tr>
<td>7</td>
<td>14.36</td>
<td>α-Copaene</td>
<td>3856-25-5</td>
<td>Sesquiterpene hydrocarbon</td>
</tr>
<tr>
<td>8</td>
<td>14.89</td>
<td>Limonol</td>
<td>78-70-6</td>
<td>Monoterpene alcohol</td>
</tr>
<tr>
<td>9</td>
<td>16.11</td>
<td>Terpinen-4-ol</td>
<td>562-74-3</td>
<td>Monoterpene alcohol</td>
</tr>
<tr>
<td>10</td>
<td>17.61</td>
<td>α-Terpinol</td>
<td>98-55-5</td>
<td>Monoterpene alcohol</td>
</tr>
<tr>
<td>11</td>
<td>18.23</td>
<td>α-Muurolene&lt;sup&gt;b&lt;/sup&gt;</td>
<td>31083-22-9</td>
<td>Sesquiterpene hydrocarbon</td>
</tr>
<tr>
<td>12</td>
<td>18.31</td>
<td>β-Selinene&lt;sup&gt;b&lt;/sup&gt;</td>
<td>17066-87-0</td>
<td>Sesquiterpene hydrocarbon</td>
</tr>
<tr>
<td>13</td>
<td>18.75</td>
<td>δ-Cadinene&lt;sup&gt;b&lt;/sup&gt;</td>
<td>483-76-1</td>
<td>Sesquiterpene hydrocarbon</td>
</tr>
<tr>
<td>14</td>
<td>20.02</td>
<td>Calamene&lt;sup&gt;c&lt;/sup&gt;</td>
<td>483-77-2</td>
<td>Sesquiterpene hydrocarbon</td>
</tr>
<tr>
<td>15</td>
<td>22.45</td>
<td>Caryophyllene oxide</td>
<td>1139-30-6</td>
<td>Sesquiterpene ketone</td>
</tr>
<tr>
<td>16</td>
<td>24.44</td>
<td>Nonanoic acid</td>
<td>112-05-0</td>
<td>n-Hydrocarbon acid</td>
</tr>
</tbody>
</table>

<sup>a</sup> Peak no. of Fig. 1.
<sup>b</sup> Identification by mass spectral match only.
<sup>c</sup> Identification by mass spectral match and retention time match with compound in previously characterized plant essential oil (see text).

Trapping Trial 3. The most promising candidate lures from trial 2 were assayed at higher release rates. Cleverenger extract of redbay (1 ml), manuka oil (300 µL), eucalyptol (300 µL), caryophyllene oxide (300 µL as 50 mg/ml methanol), and methanol (300 µL) were each applied to half of a cotton dental wick pinned directly to the center of one sticky panel of each trap. Traps were run from 31 July to 7 August 2007.

Trapping Trial 4. We compared attractiveness of manuka oil at three different release rates to freshly cut sections (12 cm diameter; 40 cm long) from a healthy redbay tree and unbaited control traps. The redbay sections were hung vertically at 2 m height from ropes tied between two nonhost trees and had two sticky panels suspended on opposite sides. Manuka oil releasers consisted of a 4-ml glass vial with a 5-mm-diameter hole in the cap through which a cotton wick (5 mm diameter, cut from a cotton deck mop; Libman, Arcola, IL) extended either 0.5, 2, or 4 cm. The wick was threaded through a cone cut from the tip of a transfer pipette that prevented the wick from slipping down into the vial. Each vial contained 6–7 g of manuka oil. Release rates were determined gravimetrically while the releasers were outdoors on a shaded porch in Athens, GA from 20 to 22 August. The experiment was conducted 14–21 August 2007.

Trapping Trial 5. Release rates of wick baits used in trial four were highly variable and not proportional to wick length. The problem was corrected by changing the dimensions of the cone supporting the wick. Releasers were tested outdoors as before from 24 to 30 August. The same treatments as trial 4 were retested using the modified releasers on 21 August to 9 October 2007. Additionally, we tested a commercially developed releaser for manuka oil (Synergy Semichemicals, Burnaby, Canada; labeled release rate, 50 mg/d). The experiment was conducted 27 August to 9 October, 2007.

Trapping Trial 6: Fractions of Manuka Oil and Phoebe Oil. We assayed two distilled manuka oil fractions supplied by the manufacturer (Coast Biologicals), a fraction enriched in the relatively more volatile constituents of manuka oil (fraction 7), and the reciprocal fraction enriched in the relatively less volatile constituents (low-odor fraction), with phoebe oil, (Aripé Citrus Agro Industrial, Montenegro, Brazil), an extract of Brazilian walnut, *Phoebe porosa* Mez. (Lauraceae), containing relatively high amounts of α-copaene (Weyerstahl et al. 1994). We compared beetle responses to traps baited with one of these three substances released from vials with 2-cm-long wicks, redbay bolts, or empty vials from 21 August to 9 October 2007.

Chemical Analyses of Redbay Volatiles and Essential Oil Baits. Volatiles were collected from redbay wood and bark onto a PTFE-encased Poropak Q cartridge (0.1 g, 50–80 mesh; Millipore, Bedford, MA). The cartridge was attached to the outlet of a 50-ml capacity glass cold-finger tube containing redbay chips (8 g; as described previously), and the inlet was attached to a charcoal filter. Air (20 ml/min) was drawn through the tube and cartridge for 3 h at room temperature. The cartridge was extracted with 1.2 ml redistilled pentane. The water fraction from the Cleverenger extract of redbay (10 ml; used as bait in field trial 1) was extracted sequentially three times with 1 ml redistilled pentane and the extracts combined. Manuka oil, its fractions, and phoebe oil were diluted in hexane (1 µL/ml) spiked with 35 µg/ml heptyl acetate as an internal standard. Extracts and diluted essential oils were analyzed on an Agilent 6890–5973 coupled gas chromatograph–mass spectral detector (GC-MS) fitted with an HP-INNOWax (60 m by 0.25 mm by 0.25 µm film; Agilent Technologies, Santa Clara, CA) column. The temperature program was 40°C for 1 min, 16°C/min to 80°C, and 7°C/min to 230°C and held constant for 10 min. Carrier gas (he-
Fig. 1. Total ion chromatogram (TIC) traces from GC-MS analyses of (a) Porapak Q–collected volatiles from pieces of chopped redbay bark and wood, (b) water condensate from distillation of redbay bark and wood on a Clevenger still, (c) unaltered manuka oil, (d) a “low-odor” manuka oil fraction, (e) the reciprocal of the low-odor fraction (fraction 7), and (f) phoebe oil. The numerous large peaks eluting after 22 min in the phoebe oil analysis were predominantly oxygenated sesquiterpenes.

Lium) flow was fixed at 1.0 ml/min, and the injector and detector ports were 200 and 240°C, respectively. Peaks were identified by mass spectral matches to published spectra and retention time matches to commercially obtained standards. When synthetic standards were not available commercially, retention times of peaks were compared with those of constituents of essential oils with previously characterized compositions.

Statistical Analyses. All field trials were replicated four times. Each trapping trial was analyzed as a two-way analysis of variance (ANOVA) with treatment and replicate (blocks) as the independent variables and beetle catch as the dependent variable using Proc
GLM (SAS Institute 2000). Means were separated using the Ryan-Einot-Gabriel-Welsch (REGWQ) multiple comparison test (Day and Quinn 1989). Data were transformed using a log transformation when the Shapiro-Wilk test for normality (Proc Univariate; SAS Institute 2000) indicated the data were not normally distributed.

Results

Extraction and Testing of Redbay Extracts. Major constituents in Poropak aeration of redbay wood and bark (Fig. 1a; Table 1) included α-pinene, β-pinene, δ-3-carene, eucalyptol, p-cymene, α-copaene, terpinene-4-ol, linalool, calamene, and nonanoic acid. Loblolly pine resin contains large amounts of both α- and β-pinene. Because we found no evidence X. glabratus were attracted by freshly cut loblolly pine (Pinus taeda L.) wood (Hanula et al. 2008), we did not investigate these particular monoterpenes further. In trapping trial 1, Clevenger, methanol, and pentane extracts of redbay wood and bark were not more attractive to X. glabratus than methanol controls (Fig. 2). However, the steam extract was attractive to Xylodesandrus crassiusculus (Motschulsky), showing that the release devices were functioning and that the water fraction of Clevenger extract of redbay was attractive to at least one species of xyleborine ambrosia beetle. GC-MS analysis of the Clevenger extract (Fig. 1b) indicated that it contained only oxygenated volatiles identified in the aeration of redbay wood and bark, including eucalyptol, linalool, terpinene-4-ol, and carophyllene oxide.

In trial 2, too few insects were trapped for statistical comparisons, although manuka oil and all chemicals combined caught the most beetles (Table 2). In trial 3, manuka oil was significantly more attractive to X. glabratus than the other compounds or the water frac-

<table>
<thead>
<tr>
<th>Attractant</th>
<th>Trapping trial 2&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Trapping trial 3&lt;sup&gt;b&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Quantity</td>
<td>N</td>
</tr>
<tr>
<td>Caryophyllene oxide (0.5 g/10 ml methanol)</td>
<td>100 µl</td>
<td>4</td>
</tr>
<tr>
<td>Cineole</td>
<td>100 µl</td>
<td>4</td>
</tr>
<tr>
<td>p-Cymene</td>
<td>100 µl</td>
<td>4</td>
</tr>
<tr>
<td>Linalool</td>
<td>100 µl</td>
<td>4</td>
</tr>
<tr>
<td>Nonanoic acid</td>
<td>100 µl</td>
<td>4</td>
</tr>
<tr>
<td>α-Terpipene</td>
<td>100 µl</td>
<td>4</td>
</tr>
<tr>
<td>(−)-Terpinen-4-ol</td>
<td>100 µl</td>
<td>4</td>
</tr>
<tr>
<td>Manuka oil</td>
<td>100 µl</td>
<td>4</td>
</tr>
<tr>
<td>Control—methanol</td>
<td>100 µl</td>
<td>4</td>
</tr>
<tr>
<td>All combined</td>
<td>100 µl of each</td>
<td>4</td>
</tr>
<tr>
<td>Steam extract of redbay</td>
<td>Not tested</td>
<td>—</td>
</tr>
</tbody>
</table>

<sup>a</sup> Liquid test compounds and the manuka oil were absorbed onto a cotton dental wick placed inside an uncapped 25-ml capacity scintillation vial.
<sup>b</sup> Liquid test compounds and the manuka oil were absorbed onto a cotton dental wick and pinned directly to the trap surface.
<sup>c</sup> Means followed by the same letter were not significantly different (P < 0.05; Ryan-Einot-Gabriel-Welch multiple comparison test).
Table 3. Attraction of X. glabratus (RAB) to either manuka oil eluted from vials with wicks of varying lengths, a commercially produced lure, or freshly cut redbay wood

<table>
<thead>
<tr>
<th>Manuka oil releaser</th>
<th>Trapping trial 4</th>
<th>Trapping trial 5</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Release rate</td>
<td>RAB/d</td>
</tr>
<tr>
<td></td>
<td>mg/d (SE)*</td>
<td>(SE)</td>
</tr>
<tr>
<td>Control</td>
<td>4 0.1 (0.06)a</td>
<td></td>
</tr>
<tr>
<td>0.5-cm wick</td>
<td>4 0.1 (0.06)a</td>
<td></td>
</tr>
<tr>
<td>2-cm wick</td>
<td>4 0.1 (0.06)b</td>
<td></td>
</tr>
<tr>
<td>4-cm wick</td>
<td>4 1.3 (0.21)b</td>
<td></td>
</tr>
<tr>
<td>Commercial</td>
<td>—</td>
<td></td>
</tr>
<tr>
<td>Redbay bolt</td>
<td>4 1.3 (0.53)b</td>
<td></td>
</tr>
</tbody>
</table>

*Release rates were determined outdoors with similar temperatures as occurred during trapping trials.

Comparison of Manuka Oil to Redbay. In trapping trial 4, manuka oil releasers were not eluting properly, and all wick lengths had similar release rates ranging from 25 to 33 mg/d (Table 3). The three wick lengths caught significantly more X. glabratus than un baited controls, and all three attracted beetles in numbers similar to redbay wood.

In trial 5, we corrected the problem with the manuka oil releasers. The 0.5-cm wicks eluted significantly less manuka oil (3.3 mg/d) than 2- or 4-cm-long wicks (15-17 mg/d). All lures with manuka oil caught significantly more X. glabratus than un baited controls (Table 3) and, except for the 4-cm-long wicks, the various release devices of manuka oil (including the commercial lure) attracted beetles in numbers similar to those captured in traps baited with fresh redbay wood.

Fractions of Manuka Oil and Phoebe Oil. All attractants tested caught significantly more beetles than un baited controls (Fig. 3). Phoebe oil, manuka oil, and fraction 7 of manuka oil did not attract beetles in significantly different numbers than redbay wood. Low odor manuka oil (the reciprocal of fraction 7) attracted significantly fewer beetles than redbay bolts or phoebe oil but not significantly fewer than whole manuka oil or fraction 7. The low odor manuka oil contained smaller amounts of higher volatility monon and sesquiterpenes than fraction 7, manuka oil, or phoebe oil (Fig. 1).

Discussion

All seven Xyleborus species for which attractants have been identified respond to ethyl alcohol (El-Sayed 2007 and references therein). However, D. Miller (personal communication) was unsuccessful in capturing X. glabratus using ethanol lures, so other compounds released from redbay wood presumably serve as host attractants for this species. Redbay ambrosia beetles are attracted to odors from both redbay and avocado, Persea americana Mill. (Hanula et al. 2008), although redbay wood is more readily available and is currently used as the operational lure.

Aerations of redbay wood contained α-pinene, β-pinene, δ-3-carene, eucalyptol, and p-cymene, but these were absent from low-odor manuka oil. Because this oil fraction was similar in attractiveness as whole manuka oil to X. glabratus (Fig. 3), these compounds were presumably unessential components of the X. glabratus attractant. Manuka oil was found to contain relatively large quantities of three sesquiterpene hydrocarbons (calamene, α-cubebene, and α-co-pa e na; Fig. 1c; Table 1), which were the only compounds that had in common with redbay extracts other than α-pinene and myrcene. In contrast, phoebe oil (Fig. 1f; Table 1), which was similarly attractive to the beetles as manuka oil or redbay wood, contained calamene and α-cubebene in much smaller quantities relative to α-co-pa e na. These results suggest α-co-pa e na may be the primary attractive component of manuka oil, phoebe oil, and redbay, although other compounds may contribute to attraction as well.

![Fig. 3. Mean (±SEM) daily catch of X. glabratus in traps baited with manuka oil or its fractions, phoebe oil, or a freshly cut redbay bolt. Columns with the same letter were not significantly different (REGWQ multiple comparison test; n = 4; P < 0.05; SAS Institute 1985).](image-url)
Crook et al. (2008) pointed out that a variety of other insects, including a number of bark beetles, respond to these sesquiterpenes. However, if α-copaene, α-cubebene, and/or calamenene are the essential attractive components of redbay wood, the currently limited commercial availability and high price of these compounds in pure form implies that manuka oil or other essential oils may be the most cost-effective sources of artificial X. glabrat us attractant. One disadvantage of using essential oils in applications requiring a standardized bait (such as population monitoring) is that the chemical composition may vary among suppliers or batches because of changes in the geographic origin, cultivar, and growing conditions of the extracted plants. This variability has been documented in manuka oil (Porter and Wilkins 1999). Nonetheless, it is likely that essential oil–based baits will provide greater consistency in trapping X. glabrat us than baits of fresh redbay wood.

Recently, Crook et al. (2008) reported manuka oil and phoebe oil were attractive to emerald ash borer, Agrilus planipennis Fairmaire (Coleoptera: Buprestidae), and that six constituents of these oils, including α-copaene and α-cubebene, stimulated olfactory sensilla of this insect. In addition, they found these compounds were produced by standing ash trees after application of stressors (girdling) capable of stimulating attraction of emerald ash borers. However, logs cut from similar, healthy trees were not as attractive as girdled trees (Poland et al. 2004).

Artificially wounded trees and bolts cut from either healthy or diseased trees are attractive to X. glabrat us (Hanula et al. 2008). Our volatiles collections were made from wood and bark of diseased trees; however, Hanula et al. (2008) found no evidence that trees either infected with the laurel wilt fungus or heavily infested with X. glabrat us were more attractive to X. glabrat us than healthy trees.

Manuka and phoebe oil should provide a practical and cost-effective bait alternative to redbay wood in traps used for monitoring X. glabrat us distribution and population trends. Additionally, the availability of practical artificial baits for X. glabrat us will permit study of attractant–based control strategies for X. glabrat us and laurel wilt. X. glabrat us populations declined to very low levels in areas where it had killed most of the mature redbay trees (Hanula et al. 2008), hence trap-out or attract-and-kill techniques might be useful for further reducing such suppressed populations giving redbay seedlings, saplings, and stump sprouts an opportunity to develop into mature trees. Such techniques combined with effective sanitation of infested trees could be used to allow redbay populations to recover. In addition, our results in combination with those of Crook et al. (2008) suggest that manuka oil may be an effective attractant for a diversity of hardwood pests and should be considered as bait for early detection of new pest introductions.

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