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1 Performance of *Sirex noctilio*'s biocontrol agent *Deladenus siricidicola*, in 2 known and predicted hosts

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11 Abstract

12 Survival of the free-living mycetophagous form of *Deladenus siricidicola*, the major biological control
13 agent of Sirex woodwasp, *Sirex noctilio*, was tested in known (*Pinus taeda*) and predicted novel (*P.*
14 *elliottii* subsp. *elliottii* x *P. caribaea* var. *hondurensis*) hybrid host taxa. Trials were established in the
15 field to simulate nematode dispersal both naturally by infected wasps and following commercial
16 inoculation, as well as in the laboratory under controlled conditions. Nematodes showed reduced
17 survival in hybrid pine compared with *P. taeda* for all tree-associated treatments, but performed
18 equivalently in petri-dish bioassays containing substrate of each taxon. Growth of *Amylostereum*
19 *areolatum*, the food source of *D. siricidicola* was lower on plates containing ground hybrid substrate
20 than on plates containing ground *P. taeda*. Some physical differences were found between taxa,
21 including differences in bordered pit diameters, tracheid widths, and basic density, but these did not
22 consistently explain reduced performance. More plant secondary compounds (predominantly
23 oleoresins) were present in hybrid taxa than in *P. taeda*, and in standing trees compared with felled
24 trees. Our results suggested that *D. siricidicola* may not be as effective in hybrid pine taxa for the
25 biological control of *S. noctilio* as it is in its current known host taxa, possibly because of reduced
26 growth of its food source, *A. areolatum* in hybrid pine.

27

28 Keywords: *Beddingia*; *Pinus taeda*; nematode; *Amylostereum areolatum*;

29

30 1. Introduction

31 Sirex woodwasp, *Sirex noctilio* F. (Hymenoptera: Siricidae) and its obligate symbiotic fungus,
32 *Amylostereum areolatum* (Chaillet ex Fr.) Boidin (Russulales: Amylostereaceae) are major invasive
33 pests of softwoods worldwide. In concert, the duo kills trees through the combined action of the
34 wasp's phytotoxic mucus and growth of the fungus, both deposited into trees during oviposition
35 (Ryan and Hurley 2012). The two species are established as exotic invaders in Australia, New Zealand,
36 South Africa, China, South America and North America (Adams et al. 2011; Li et al. 2015), and have
37 gradually expanded their Australian geographic range, reaching southern (temperate) Queensland in
38 2009 (Carnegie and Bashford 2012). Climate modelling predicts subtropical regions of Queensland
39 will support *S. noctilio* establishment (Carnegie et al. 2006), suggesting the wasp is likely to continue
40 to move north into valuable subtropical pine estates, where previously unencountered hosts,
41 particularly southern pines (*Pinus elliottii* (Engelm) var. *elliottii* and *P. caribaea* var. *hondurensis*
42 (Sénécl.) W.H.Barrett and Golfari) (Pinales: Pinaceae) and their hybrids, are grown in large-scale
43 plantations (Gavran 2014). No *Pinus* species are known to be resistant to Sirex, although the level of
44 susceptibility varies between species (Ryan and Hurley 2012): it is likely that these hybrids will be
45 suitable hosts but uncertain how host characteristics may interact with biological control (Bedding
46 2009).

47
48 The parasitic nematode, *Deladenus (=Beddingia) siricidicola* Bedding (Tylenchida: Neotylenchidae), is
49 the most effective biological control agent used in *S. noctilio* management worldwide, but shows
50 very variable and inconsistent parasitism rates within and between regions (Hurley et al. 2007).
51 Mechanisms behind the variation in biological control success using nematodes have not been
52 clearly explained (Hurley et al. 2008; Slippers et al. 2012). Extremely low genetic diversity of *D.*
53 *siricidicola* may affect their ability to adapt to novel environments and hosts (Mlonyeni et al. 2011),
54 and aspects of the host trees themselves may also affect nematode survival (Bedding 2009). The
55 efficacy of the standard biological control program developed in winter rainfall regions appears to be
56 reduced in different climatic conditions, as was the case in South America and South Africa (Slippers
57 et al. 2012). Thus, there are concerns that if the woodwasp establishes in the subtropical pine estate
58 in Australia, the current biological control practice may not be effective.

59
60 *Deladenus siricidicola* has a free-living fungal-feeding form that is mass-produced for commercial
61 inoculation in *Sirex* biological control programs (Slippers et al. 2012). In proximity to *Sirex* larvae,
62 female nematodes shift phase to a parasitic form that ultimately sterilises female *Sirex* by entering
63 the ovaries and eggs, and is then transmitted (as the free-living, fungal-feeding form) during
64 oviposition by the infected wasp (Slippers et al. 2012). The free-living form is thus spread naturally
65 when infected *S. noctilio* females lay packets of nematodes in place of viable eggs into trees, and
66 operationally by inoculating laboratory-cultured nematodes into felled trees. The success of the
67 biological control program is influenced by interactions between wasp, nematodes and fungal strains
68 (Morris et al. 2012; Slippers et al. 2015), other insects and fungi (Yousuf et al. 2014), climatic
69 conditions (Hurley et al. 2007) and host tree characteristics (Bedding 2009).

70
71 Loblolly pine, *Pinus taeda* L., native to SE USA, is grown extensively in South America and, along with
72 *P. elliottii* is the most commonly planted commercial pine species in Brazil and Argentina (Klasmer
73 and Botto 2012) and the most susceptible to *S. noctilio* (Fenili et al. 2000; Iede et al. 2012). In this
74 host, *S. noctilio* parasitism by *D. siricidicola* varies between 0 and 90% (Iede et al. 2012; Nahrung et
75 al., 2015). *Pinus taeda* is grown commercially in southern Queensland and northern New South
76 Wales, where it is readily attacked by *S. noctilio* (Carnegie et al. 2005), albeit less successfully than *P.*
77 *radiata* (Nahrung et al. 2015). A commercial hybrid *Pinus* taxon, *Pinus elliottii* subsp. *elliottii* x *P.*
78 *caribbea* var. *hondurensis* (PEE x PCH) was developed in 1958 (Nikles 1996) and is now widely grown
79 in coastal plantation regions of south-eastern Queensland. The performance of *Sirex* wasp, fungus
80 and nematode in this host is unknown, and hence the overall impact of the *Sirex* wasp in this host is
81 impossible to predict. Here, we use field and laboratory studies, in the absence of *S. noctilio*, to
82 compare performance and survival of both the *Amylostereum* fungus and the free-living
83 mycetophagous form of *D. siricidicola* on these different host taxa. We also compare physical and
84 chemical properties of the two hosts with potential to impact on the survival and development of
85 nematodes and/or fungus, including moisture content, wood density, tracheid and bordered pit
86 diameter.

87

88 **2. Materials and Methods**

89 *2.1. Nematode survival and development*

90 *2.1.1. Field trials*

91 Field trials were established in a subtropical plantation (Beerburum, Site 1, PEE x PCH F2 hybrids)
92 and a higher altitude, temperate plantation (Passchendaele, Site 2, *P. taeda*) (Figure 1) in
93 Queensland, and intended to mimic (a) natural nematode spread by ovipositing females (*i.e.* into
94 standing trees, $n = 5/\text{site}$) and (b) commercial inoculation into felled trees ($n = 5/\text{site}$). Because *S.*
95 *noctilio* is not yet present in the subtropics (Site 1), surrogate techniques were used to introduce *A.*
96 *amylostereum* and *D. siricidicola* into all trees. All trees were poisoned using standard trap tree plot
97 techniques designed to stress trees and make them susceptible to *S. noctilio* oviposition (Haugen et
98 al. 1990, Gitau et al. 2013) to provide conditions similar to those that nematodes would encounter
99 either deposited by Sirex females (standing trees), or during artificial inoculation (felled trees). Ten
100 trees at each site were poisoned with Dicamba (3,6-dichloro-2-methoxybenzoic acid) at a rate of 1
101 mL/10 cm circumference in March 2012. In May 2012, five trees at each site were felled, and five
102 were left standing. Each tree was inoculated at four points (approximately 0, 90, 180 and 270
103 degrees around the stem) at each 1 m interval along the trunk for 4 m (*i.e.* 16 inoculation points per
104 tree) with a slurry of nematode rearing media containing both nematodes and the *Amylostereum*
105 fungus supplied by Ecogrow Environment Ltd. Nematodes were mixed to a concentration of
106 approximately 1 million nematodes/500 mL of *Amylostereum* rearing slurry, and inoculated into the
107 tree using the standard hammer punch method (Carnegie & Bashford 2012), such that each
108 inoculation point received approximately 995 nematodes. Samples were taken by removing a 4 cm
109 diameter core around the inoculation point to a depth of 1 cm with a hole saw at each inoculation
110 height (1-4 m) one and three months after inoculation. The outer bark was removed to minimise
111 inclusion of any bark feeding nematodes, and then cores were soaked in 30 mL of tap water for 12
112 hours and the supernatant examined under a dissecting microscope (40 \times) for the presence of
113 nematodes. The proportion of cores per tree that scored positive for nematodes was calculated,
114 and results for each sample time and treatment were compared using chi-square tests.

115

116 2.1.2. Laboratory trials

117 To enable comparison between taxa under identical conditions, three inoculated billets from the top
118 of each felled tree from the field trial above were taken to the laboratory immediately following
119 inoculation, and stored in a controlled environment room (20 °C, 60% RH). One billet from each tree
120 was destructively sampled at one and three months post-inoculation, by cutting into slices 3.5 ± 1.5
121 cm thick, and soaking each in 300 mL of water separately in a plastic bag for 12 hours. The number
122 of nematodes in each sample was estimated by decanting the supernatant into a flat dish and
123 allowing nematodes to settle to the bottom. The number of nematodes in each of four fields of view
124 of a dissecting microscope (40 \times) was counted and the total number of nematodes per sample was
125 estimated by correcting for the proportion of the total surface area that our counts represented. We
126 also examined the distribution of nematodes relative to the inoculation point by combining data
127 from slices to the equivalent of approximately 10 cm intervals, and mapping the proportion of
128 nematodes recovered from each section.

129

130 2.2. *Amylostereum* survival and spread

131 Growth of *Amylostereum* fungus on an artificial substrate incorporating each of the two taxa was
132 compared. Outer sapwood samples were collected from the same trees as the field trials by
133 removing cross-sections between 4 and 10 cm thick about 2 metres from the tree base. The bark
134 was removed from each biscuit and sections of outer sapwood chosen to minimise blue-stain
135 fungus. Samples were ground in a rotary mill (Siemens) with a 1 mm mesh, then sieved to separate

136 and collect all material between 0.5 and 2 mm (coarse), and less than 0.5 mm (fine). Approximately
137 equal amounts of the coarse ($0.503 \pm 9.9 \times 10^{-4}$ g) and fine ($0.502 \pm 1.0 \times 10^{-3}$ g) ground sapwood
138 samples from each tree were mixed together, sterilised by irradiation (25 kGy for 25 hours), mixed
139 with agar (1.2%, Oxoid Technical Agar #3, 15 mL) and poured into separate Petri dishes. Control
140 plates (agar only) were also prepared.

141

142 *Amylostereum areolatum* used in the trials was sourced from a commercial culture maintained by
143 Ecogrow Environment Pty Ltd. The Ecogrow culture was subcultured onto Potato Dextrose Agar
144 (PDA, 3.9%, Difco, 15 mL) plates and grown for use in the trials. The prepared agar plates containing
145 ground sapwood were inoculated with a plug (3.3 mm diameter) of the *A. areolatum* culture placed
146 on the surface of the agar in the centre of each plate. Plates were sealed with Parafilm™ and stored
147 in the dark at 26 °C for the duration of the experiment.

148

149 Three measures of fungal growth were made for each plate. Radial growth was measured from the
150 central inoculation point to the furthest point of hyphal growth visible under a dissecting microscope
151 (40×) (mean of four readings from 0, 90, 180 and 270 °). Overall hyphal density was a subjective
152 measure of density of hyphal growth across the plate (scale of 0 - 3). Hyphal expansion was a
153 subjective measure of the density of hyphae at the leading edge of the radial expansion (scale of 1 -
154 6). Assessments were made at 4, 7, 11 and 14 days post inoculation, except hyphal density and
155 hyphal expansion which began on Day 7. By Day 14 hyphal growth had reached the edge of the petri
156 dish in most cases, preventing accurate measures of hyphal density. Data were analysed using
157 Mann-Whitney U-tests, adjusted for multiple comparisons using a Bonferroni correction.

158

159 Half of the plates used for assessing fungal growth were inoculated with nematodes eight days after
160 initial inoculation with *Amylostereum*. A 10 µL suspension of nematodes from Ecogrow (1 in 24
161 dilution) was inoculated at three positions on each plate (0, 120 and 240 °) giving approximately 60
162 nematodes per plate. Plates were sealed with Parafilm™ and incubated at 23 °C in darkness.

163

164 After twelve days, plates were visually assessed under a dissecting microscope for the presence
165 (low/medium/high) or absence of nematodes and given a score from 0 - 3. Plates were then stored
166 at 4 °C until final assessment based on Morris et al. (2012). Briefly, plates were filled with 15 mL of
167 distilled water and allowed to sit for 10 minutes then swirled and the supernatant was poured off.
168 Plates were then rinsed four times and all washings were combined. The number of nematodes
169 present (viewed under a dissecting microscope at 25×) was assigned to 'high', 'medium' and 'low'
170 categories, and given a score from 1 - 3.

171

172 2.3. Physical properties

173 2.3.1. Moisture content and wood density

174 Moisture content was measured at inoculation using the wood plug removed for sampling and at
175 three months using the core removed for nematode sampling. Samples were transported cooled, to
176 the laboratory in sealed airtight bags, then weighed prior to soaking for nematode extraction (wet
177 mass). After soaking, samples were dried at 103 °C, then reweighed until they reached constant
178 mass (dry mass). Moisture content was then estimated as ((wet mass-dry mass)/(wet mass)). These
179 data were arcsine-square root transformed prior to analysis using a two-sample t-test. Basic wood
180 density was estimated using the equation (dry mass/wet volume) using cross-sectional slices of mid-

181 bole billets stored under controlled conditions. These data were compared using a Mann-Whitney U-
182 test because of non-homogeneity of variance.

183

184 2.3.2. Tracheid and bordered pit diameter

185 Test billets were cut 2 m from the base of the standing trees used in inoculation trials (n = 3 per
186 taxon). A 4 cm thick disc was cut from the lower end of each billet, and a 15 mm diametral strip was
187 cut from each disc through the “pith” from edge-to-edge (bark removed). Particular attention was
188 paid to avoiding any obvious defects in the sample, such as compression wood, resin pockets and
189 branch knots. The study looked at the 2nd and 5th growth rings from the bark, to approximately
190 mimic the hammer inoculation depth and ovipositor penetration range of *Sirex* females. The 2nd
191 and 5th ring segments were extracted from the best radii of the sample and dissected into separate
192 earlywood and latewood material. Each separated sample was tangentially cut into “matchstick”
193 sized pieces and placed in individual 30 mL test tubes ensuring sample integrity throughout. Thus,
194 for each taxon, there were replicated samples from early and late wood for each of ring 2 and ring 5
195 for each of three trees (n = 12 samples per taxon).

196

197 Samples were macerated by adding 10 mL of freshly prepared macerating fluid (1:1 mixture of
198 hydrogen peroxide 100 vols – 35 % W/W and glacial acetic acid) to each 30 mL test tube with vented
199 cap and placing test tubes into 500 mL beakers containing 250 mL gently simmering water. The
200 beakers were placed on top of a heating plate in a fume cabinet and simmered gently for 5 hours
201 (keeping water in the beakers at an appropriate level) by which time the samples had reached a
202 totally colourless state. The test tubes were removed from the beakers and the macerating fluid was
203 carefully poured off, taking care not to loosen fibres in the process. The samples were washed
204 several times with deionised water to remove all trace of the macerating fluid and allowed to cool
205 before storage at 4 °C. Temporary slides were made in order to capture digital images of individual
206 fibres. Samples were shaken vigorously to separate fibres and a drop of the mixture was slide
207 mounted using Safranin stain (1% in 95% alcohol diluted 50/50 with deionised water). Six slides for
208 each of the 12 samples were prepared for digital photography using a “Leica” DMLB compound
209 microscope with digital image capture capability coupled with a NIKON “Digital Sight” using DS-L2
210 Version 322 software. Each slide was viewed at 40× magnification and visually scanned, in a
211 systematic manner to locate suitable pits for image capture. Sufficient slides were scanned and
212 images captured to ensure at least 50 pits were included. The required measurements were
213 obtained using *Image-Pro Plus V6.3* digital imaging software.

214

215 Pit apertures on the top plane of the fibre were measured, initially along the longest diameter and
216 then at 90 ° to the initial diameter. As many of the suitable pit apertures as possible were measured
217 to ensure adequate co-efficient of variation. Three internal fibre diameters were measured, at the
218 point where the pit apertures were measured and averaged to give the “mean fibre width”. Data
219 were analysed using one-way ANOVA, with Fisher’s protected LSD post-hoc tests.

220

221 2.4. Chemical properties

222 Further samples of ground sapwood used in the petri dish bioassays were taken, and approximately
223 equal amounts of the coarse ($1.004 \pm 1.0 \times 10^{-3}$ g) and fine ($1.004 \pm 1.3 \times 10^{-3}$ g) samples were mixed
224 together and extracted in dichloromethane (Ajax, 10 mL) for 48 h in the dark at 4 °C (after Cadahia
225 1997, Eyles 2003, Hayes *et al*, 2014). Samples were stored in the freezer until analysis.

226

227 Samples (1 μ L) were analysed using a gas chromatograph (GC) (Agilent 6890 Series) coupled to a
228 mass spectrometer (MS) (Agilent 5975) and fitted with a silica capillary column (Agilent, model HP5-
229 MS, 30 m \times 250 μ m ID \times 0.25 μ m film thickness). Data were acquired under the following GC
230 conditions - inlet temperature: 250 $^{\circ}$ C, carrier gas: helium at 51 cm/s, split ratio 13:1, transfer-line
231 temperature: 280 $^{\circ}$ C, initial temperature: 40 $^{\circ}$ C, initial time: 2 min, rate: 10 $^{\circ}$ C/min, final
232 temperature: 260 $^{\circ}$ C, final time: 6 min. The MS was held at 280 $^{\circ}$ C in the ion source with a scan rate
233 of 4.45 scans/s.

234

235 Peaks present in blank dichloromethane (control) samples were discarded from analysis in test
236 samples. Tentative identities were assigned to peaks with respect to the National Institute of
237 Standards and Technology (NIST) mass spectral library. Mass spectra of peaks from different samples
238 with the same retention time were compared to ensure that the compounds were indeed the same.

239

240 Mann-Whitney U-tests were used to compare the number of compounds between treatments, and
241 ANOSIM was used to determine differences in overall chemical profiles, with post-hoc t-tests used to
242 determine whether relative peak areas differed significantly for compounds found to separate the
243 groups.

244

245 3. Results

246 3.1. Field trials

247 a) *standing trees* No *D. siricidicola* were recovered from any cores from any standing hybrid trees at
248 either of the sample times (n = 20 cores/sample). Nematodes were recovered from all *P. taeda*
249 trees, from an average of $45 \pm$ s.e. 9.3 % of cores/tree at the one-month sample, and from 60 % of
250 trees and 12.5 ± 7.2 % of cores/tree at three months post-inoculation. Taxa were significantly
251 different at the one-month sample ($\chi^2_1 = 11.6$, $P < 0.001$), but not after three months ($\chi^2_1 = 3.2$, $P =$
252 0.07).

253

254 b) *felled trees* One month after inoculation, nematodes were recovered from all felled *P. taeda*
255 trees, and three-quarters of cores/tree (75 ± 16 %), but only from three (60%) hybrid trees, and less
256 than half of cores sampled (40 ± 17 %) ($\chi^2_1 = 5.01$, $P = 0.03$). After three months, however,
257 nematodes were recovered similarly from both taxa, at 30 ± 0.15 % of cores from *P. taeda*, and $30 \pm$
258 0.12 % cores from hybrid pine ($\chi^2_1 = 0.01$, $P = 0.09$).

259

260 3.2. Laboratory trials

261 3.2.1. *Whole tree sections* After one month, there was a greater than seven-fold difference in the
262 number of nematodes recovered from *P. taeda* billets than hybrid billets stored under controlled
263 conditions since inoculation. On average, each ten-centimetre section of *P. taeda* had $17,341 \pm$
264 4,591 estimated nematodes, significantly more than hybrids, which had only $2,248 \pm 557$ per section
265 (Mann-Whitney U-test, $U = 25$, $P = 0.008$). On examining the distribution of nematodes throughout
266 the billet, there was a significant, negative relationship between the number of nematodes and the
267 distance from the inoculation point for *P. taeda* (Spearman rank correlation, $\rho = -0.65$, $P < 0.001$),
268 but not for hybrids ($\rho = -0.13$, $P = 0.25$), with nematodes appearing normally distributed around the
269 inoculation point for *P. taeda*, but not for hybrids (Figure 2).

270

271 Three months after inoculation, although *P. taeda* supported over five times as many nematodes as
272 hybrids (*P. taeda*: 1770 ± 1504; hybrid 322 ± 228), the average estimated number of nematodes per
273 section was not significantly different between the taxa (Mann-Whitney U-test, U = 19, P = 0.222).

274

275 2.3.3. Petri dish bioassays

276 *Amylostereum areolatum* grew on both ground-wood+agar substrates (Figure 3), but not on agar-
277 only control plates. The radial growth (mm) of *A. areolatum* did not differ between felled and
278 standing trees within each taxon (Mann-Whitney U tests, U = 1.5 – 6.0, P = 0.04 – 1.0; Bonferroni-
279 adjusted P = 0.02), or between *P. taeda* and hybrid pine (Mann-Whitney U test, U = 23 – 30, P = 0.38
280 – 0.88) (Figure 3).

281

282 However, hyphal density (Figure 4: Mann-Whitney U test, U = 9-13.5, P = 0.01 – 0.04) and hyphal
283 expansion (density at the leading edge of fungal growth) (Figure 5: Mann-Whitney U test, U = 4.5 -
284 11, P = 0.003 – 0.03) were always higher in *P. taeda* than the hybrid, and both were always higher
285 than the plain agar control, which scored 0 and 1, respectively, at each assessment.

286

287 Twelve days after plates were inoculated with nematodes, nematode eggs were present on 7/8
288 hybrid-substrate plates, and 4/8 *P. taeda* substrate plates; this difference was not statistically
289 different ($\chi^2_1 = 2.6$, P = 0.11). Likewise, the number (high, medium or low) of nematodes did not
290 differ between plates of different taxa (Mann-Whitney U test, U = 21.0, P = 0.27), with hybrid plates
291 scoring an average 2.13 ± 0.13 and *P. taeda* scoring 1.75 ± 0.16. At the conclusion of the experiment,
292 nematode populations persisting on both taxa were not statistically separable (U = 17.5, P = 0.17),
293 with an average score of 2.13 ± 0.30 for hybrid plates, and 1.38 ± 0.32 per *P. taeda* plate.

294

295

296 3.3. Physical properties

297 3.3.1 Moisture content and wood density

298 Initial moisture content, and moisture content three months after inoculation did not differ between
299 taxa for any treatment (Table 1), however the basic density of hybrid wood (0.037 ± 0.001 g/m³) was
300 significantly greater than that of *P. taeda* (0.034 ± 0.001 g/m³) (Mann-Whitney U test, U = 1403, P =
301 0.004).

302

303 3.3.2 Tracheid and bordered pit diameter

304 Overall, pit diameter was larger in hybrids than in *P. taeda* (t-test, $t_{22} = 4.7$, P < 0.001). Considering
305 samples separately, however, the difference was significant only in ring 5 (ANOVA, $F_{7,23} = 3.4$, P =
306 0.02, Fisher's protected LSD post-hoc test) (Figure 6).

307

308 Tracheid width also differed between taxa, this time being larger in *P. taeda* than in hybrids but
309 only in ring 2 earlywood (ANOVA, $F_{7,23} = 3.9$, P = 0.01, Fisher's protected LSD post-hoc test)
310 (Figure 7).

311

312

313 3.4. Chemical analysis

314 For each taxon considered separately, there were significantly more compounds in standing than
315 felled trees (*P. taeda*: $U = 0.0$, $P = 0.036$; hybrid: $U = 0.0$, $P = 0.036$), and significantly more in
316 standing trees when both taxa were combined (Mann-Whitney $U = 0.0$, $P < 0.001$). However, there
317 was no difference between the taxa with respect to number of compounds for either standing ($U =$
318 1.5 , $P = 0.400$) or felled trees ($U = 5.5$, $P = 0.175$). Standing hybrids contained three unique
319 compounds, and standing *P. taeda* had one unique compound (Figure 8). Overall, seven compounds
320 were detected in hybrids that were not found in *P. taeda* (Figure 6) and two in *P. taeda* but not in
321 hybrid.

322

323 Considering all trees, there was no difference between the taxa in their total chemical profile
324 (ANOSIM: Global $R = 0.08$, $P = 0.135$). However standing trees differed from felled trees (Global $R =$
325 0.268 , $P = 0.047$) and so were considered separately. The taxa differed for standing trees only,
326 although the groups overlapped (Global $R = 0.481$, $P = 0.10$), with significantly more palustic acid,
327 tolurosol and an unidentified compound (RT 20.233, MW 286) detected in hybrids than in *P. taeda*
328 (Table 2). For felled trees, the two taxa were barely separable (Global $R = 0.208$, $P = 0.143$), with
329 relative amounts of callistic acid, dehydroabietic acid (DHA) and methyl 4-hydroxy-2-methoxy-3,5,6-
330 trimethyl benzoate contributing most strongly to group separation. For standing trees, seven
331 compounds contributed to the top 50% of dissimilarity between taxa – of which 71% had retention
332 times greater than 20 minutes (i.e. non-volatile compounds). For felled trees, 8 compounds
333 contribute to the top 50% of dissimilarity – of which 63% had retention times greater than 20
334 minutes (Table 3).

335

336 Of the total 88 compounds detected during the analysis less than one third had retention times
337 greater than 20 minutes (ie non-volatile compounds), although they were 60-70% of the compounds
338 that differed between taxa. Oleoresins such as these present both a physical and chemical defence
339 system against stem-boring insects and are composed mostly of monoterpenes and diterpene resin
340 acids.

341

342 4. Discussion

343 The results of this study suggest poorer survival of *Deladenus siricidicola* nematodes in the novel
344 PEE x PCH hybrid host compared with the known host *P. taeda*. Nematodes were never recovered
345 from standing hybrid pine, and nematode survival was lower in hybrid felled trees and billets stored
346 in common conditions compared with *P. taeda*. We identified physical and chemical differences
347 between the two taxa and here consider the potential impact of these in influencing the observed
348 patterns of survival and performance of both nematodes and fungus.

349

350 Some physical sapwood properties differed between the taxa studied, and may partly explain the
351 distribution of nematodes observed in the hybrid billets in the laboratory compared with *P. taeda*.
352 Tracheid widths of both pine species were similar to the width of adult nematodes ($\approx 30 \mu\text{m}$ –
353 Bedding 1968), so are unlikely to have restricted the spread of nematodes through the timber. The
354 generally larger size of bordered pits in the hybrid may explain the more even spread of the
355 nematodes through these billets compared with that of *P. taeda*, facilitating movement of

356 nematodes between tracheids via bordered pits. We found higher wood density associated with
357 lower nematode success, with hybrids having significantly higher density than *P. taeda*. Likewise,
358 *Bursaphelenchus xylophilus* nematodes have been associated with *P. pinaster* of lower wood density
359 (Rodrigues et al. 2010). Fast drying has also been postulated as a possible explanation for low *D.*
360 *siricidicola* performance in areas with warm dry winters (Bedding 2009) but both treatment taxa
361 here did not differ in moisture content at inoculation or after three months. Consistent with Hurley
362 et al's (2008) results, we found lower nematode survival in inoculated standing trees compared with
363 felled ones.

364

365 Although we identified some differences in chemical profiles of the two taxa, we did not detect any
366 differences in performance of nematodes inoculated onto plates of each, suggesting that these
367 differences in plant chemistry did not directly impact their survival. Propagation of the pine wilt
368 nematode, *B. xylophilus*, is influenced by monoterpene concentrations in *P. massoniana* (Niu et al.
369 2012) but the taxa investigated in our study did not differ in their concentrations of these.

370

371 The most likely explanation for differential nematode performance between the taxa tested here is
372 the lower growth of free-living *D. siricidicola*'s food source, *Amylostereum areolatum*, on hybrid
373 substrate. *Amylostereum areolatum* grew more densely on plates containing *P. taeda* than the
374 hybrid substrate, and this may also explain the difference in nematode performance in billets of
375 each under controlled conditions. There was a significant relationship between the number of
376 nematodes recovered in *P. taeda* and the distance from the inoculation point, suggesting that the
377 fungus established and grew from the inoculation point, enabling nematodes to feed and reproduce.
378 In contrast, both the lower numbers and broader distribution of nematodes throughout inoculated
379 hybrid billets suggests that nematodes migrated in search of food, and were unable to reproduce at
380 as high a rate as in *P. taeda*. We were unable to isolate *A. areolatum* from the billets to confirm its
381 distribution because of contamination, but our plate bioassay clearly showed reduced growth of
382 *Amylostereum* on hybrid substrate. In contrast, *D. siricidicola* performed equivalently on plates
383 containing substrate of each taxon, presumably because despite its lower growth on hybrid
384 substrate, the amount of fungus was not limiting to nematode survival. *Deladenus siricidicola*
385 performs better on slower-growing fungus (Morris et al. 2012), and in circumstances where the
386 population of nematodes is very small in the presence of a large *A. areolatum* biomass, the fungus
387 can invade and kill nematode eggs (Morris et al. 2014). However, we found no evidence of
388 diminished nematode survival on plates with differential growth rates of *A. areolatum* in our study.

389

390 The chemical differences we found between taxa may have influenced the growth of *A. areolatum*.
391 For example, the terpenoid torulosol was among the major compounds that differentiated hybrids
392 and *P. taeda* in both standing and felled trees: torulosol and its derivatives have been identified in
393 extracts that show antifungal activity (Meneses et al. 2009). The majority of compounds we detected
394 in our samples were non-volatile oleoresins, composed mostly of monoterpenes and diterpene resin
395 acids, which can present both a physical and chemical defence system against invading pathogens
396 and insects. Growth of a mycangial basidiomycete fungus was inhibited by several terpenoid
397 compounds tested by Bridges (1987). Similarly, in a study of timber decay resistance, heartwood
398 extracts from *P. elliotii* and *P. caribaea* protected *Pinus radiata* sapwood from decay by individual
399 basidiomycete fungi in the laboratory (Kennedy et al. 1995). In this case, protection against
400 basidiomycete decay did not occur in contact with unsterile soil, where it was proposed that extracts
401 were detoxified by organisms absent from living trees.

402

403 If the lower survival of *D. siricidicola* in hybrid taxa is explained by the lower growth of *A. areolatum*
404 in that taxon as occurred in the laboratory, it is possible that the large variation in parasitism rates

405 reported globally (Hurley et al. 2007; Hurley et al. 2012; Slippers et al. 2012) could be the result of
406 differential fungal growth on different genotypes, phenotypes or chemotypes of trees. Breeding
407 pines for resistance to *Sirex* has not been a focus of management because of the success of existing
408 management strategies (Slippers et al. 2015), but it is possible that germplasm could be selected
409 that would promote nematode survival. Further work in identifying these factors is warranted.

410

411 Because *S. noctilio* is not yet present in the regions of Australia where hybrid pines are grown, we
412 tested only the survival of the mycetophagous form of the nematode. Further examination of the
413 ability of hybrid pine taxa to support *S. noctilio* development, and parasitism by *D. siricidicola* is
414 required before the wasp reaches these regions.

415

416

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420 Queensland) for their laboratory assistance, analyses, site access and preparation. We also thank
421 Robin Bedding for useful discussions, confirming the identification of recovered nematodes, and
422 valued input. Ecogrow supplied the nematode and fungal cultures, and this work was partially
423 funded by the National *Sirex* Coordination Committee.

424

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520 **Figure and Table captions**

521 **Figure 1:** Map of Queensland, Australia, showing the location of the two field sites. Site 1
522 (Beerburrum) comprises hybrid pine grown in the subtropics; Site 2 (Passchendaele) comprises *Pinus*
523 *taeda* grown in higher altitude, temperate conditions.

524

525 **Figure 2:** Mean + s.e. proportion of nematodes recovered from each 10 cm section of *Pinus taeda*
526 (grey) and hybrid pine (black) billets kept under controlled conditions, for one month after
527 inoculation into the centre of the billet.

528

529 **Figure 3:** Mean \pm s.e. radial growth (mm) of *Amylostereum areolatum* on agar plates containing
530 either felled (open symbols) or standing (solid symbols) ground *Pinus taeda* (squares) or hybrid pine
531 (circles).

532 **Figure 4:** Hyphal density of *Amylostereum areolatum* measured seven and eleven days after
533 inoculation onto agar plates containing ground *P. taeda* (grey) and hybrid (black). Hyphal density
534 differed between taxa at both assessments (see text).

535

536 **Figure 5:** Hyphal expansion (density at the leading edge of fungal growth) of *Amylostereum*
537 *areolatum* measured seven, eleven and fourteen days after inoculation onto agar plates containing
538 ground *P. taeda* (grey) and hybrid (black). Hyphal expansion differed between taxa at each
539 assessment (see text).

540

541 **Figure 6:** Average + s.e. pit diameter (μm) from growth ring 2 (R2) and 5 (R5) early and late wood of
542 *Pinus taeda* (grey) and F2 hybrids (black). Different letters above bars designate means that differ
543 significantly.

544 **Figure 7:** Average + s.e. tracheid width (μm) from growth ring 2 (R2) and 5 (R5) early and late wood
545 of *Pinus taeda* (grey) and F2 hybrids (black).

546

547 **Figure 8:** Number of compounds detected using GC-MS in felled and standing *Pinus taeda* and hybrid
548 pine; intersection of rectangles shows the number of compounds in common between groups.

549

550 **Table 1:** Mean \pm s.e. moisture content of *Pinus taeda* and hybrid pine treatments at inoculation (i),
551 and three months after inoculation (3) with *Deladenus siricidicola* and *Amylostereum areolatum*.

552

553 **Table 2:** Retention time, compound ID, and peak area of the seven compounds that best separated
554 the two taxa as standing trees.

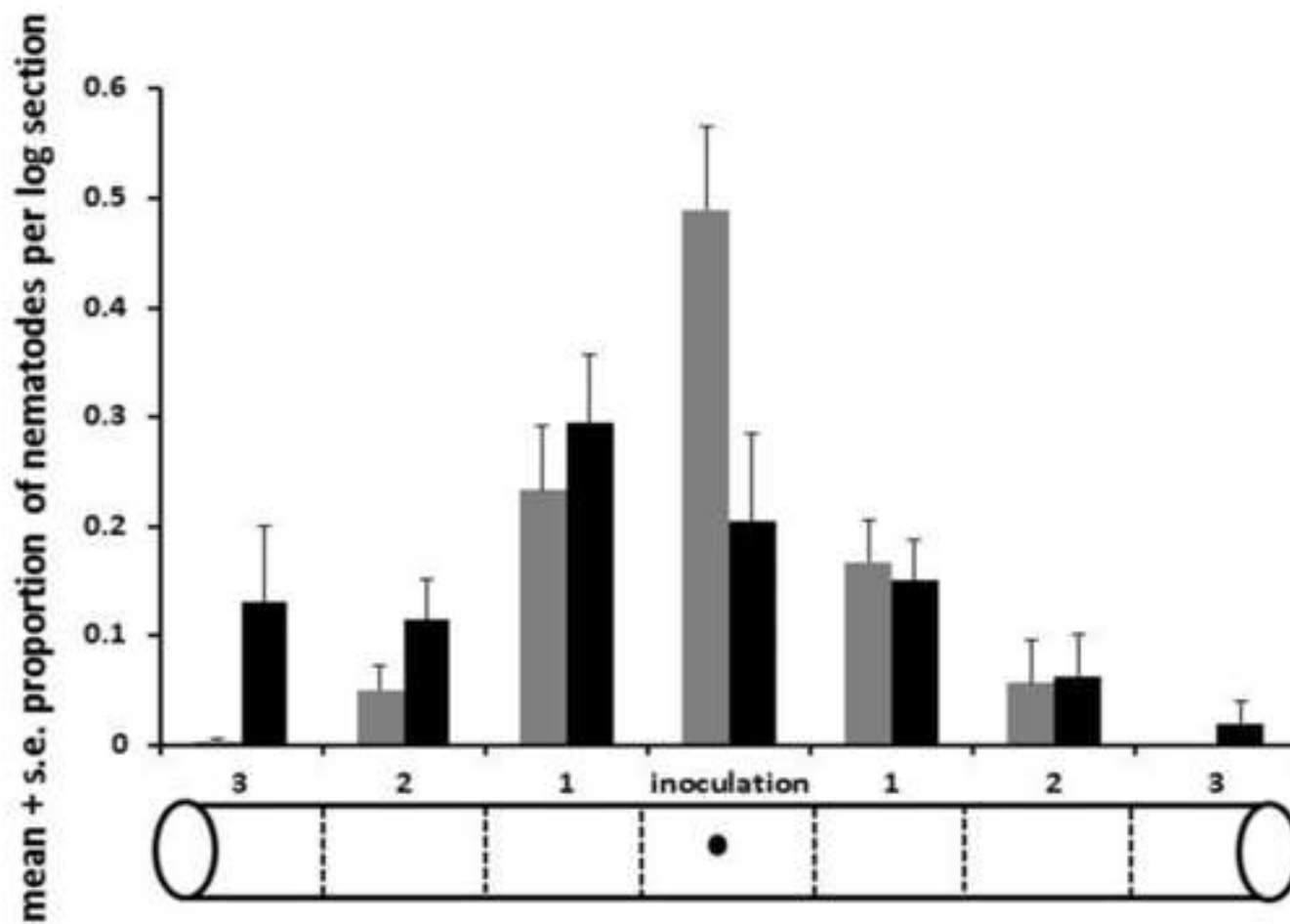
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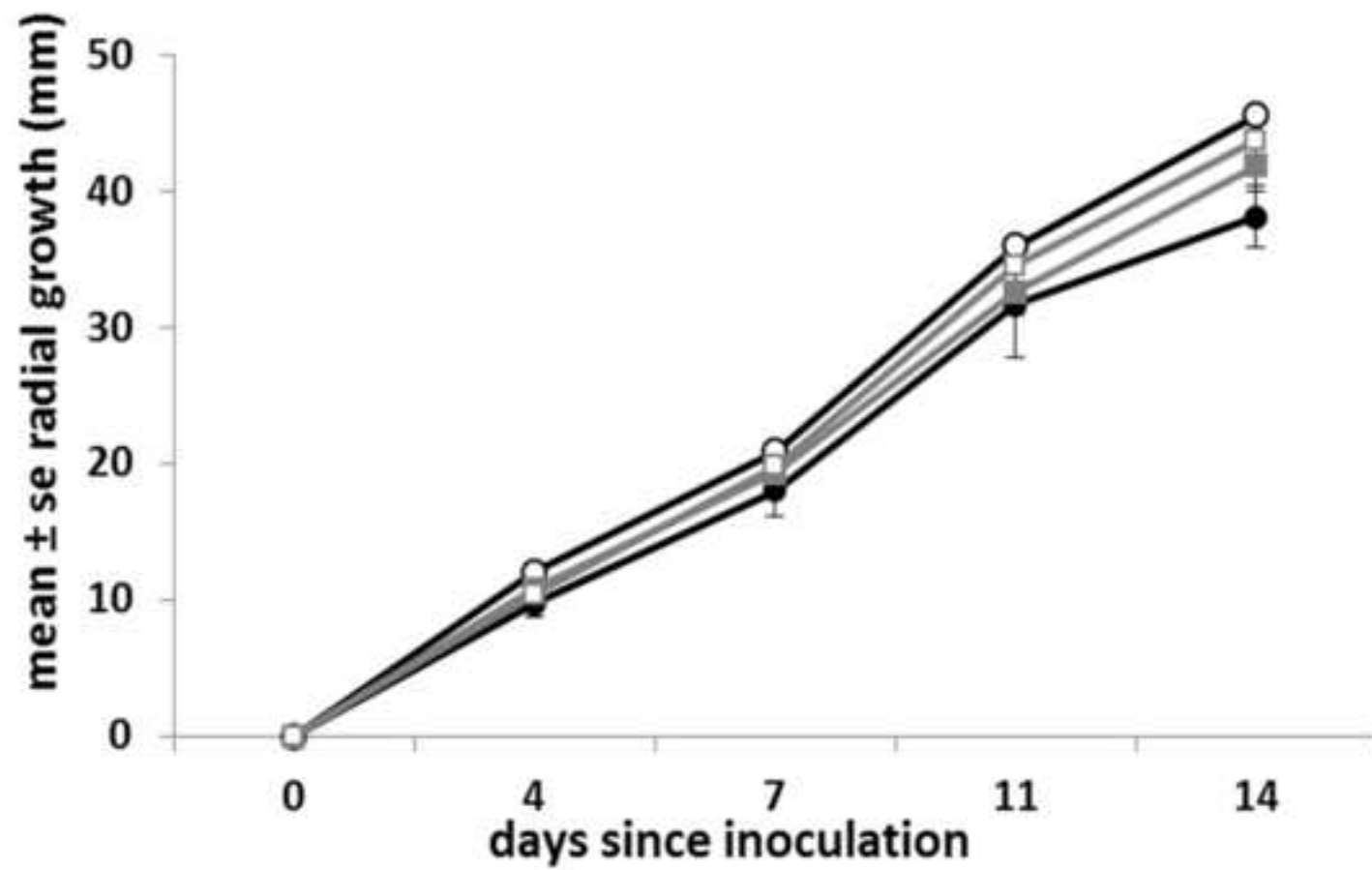
556 **Table 3:** Retention time, compound ID, and peak area of the seven compounds that best separated
557 the two taxa as felled trees.

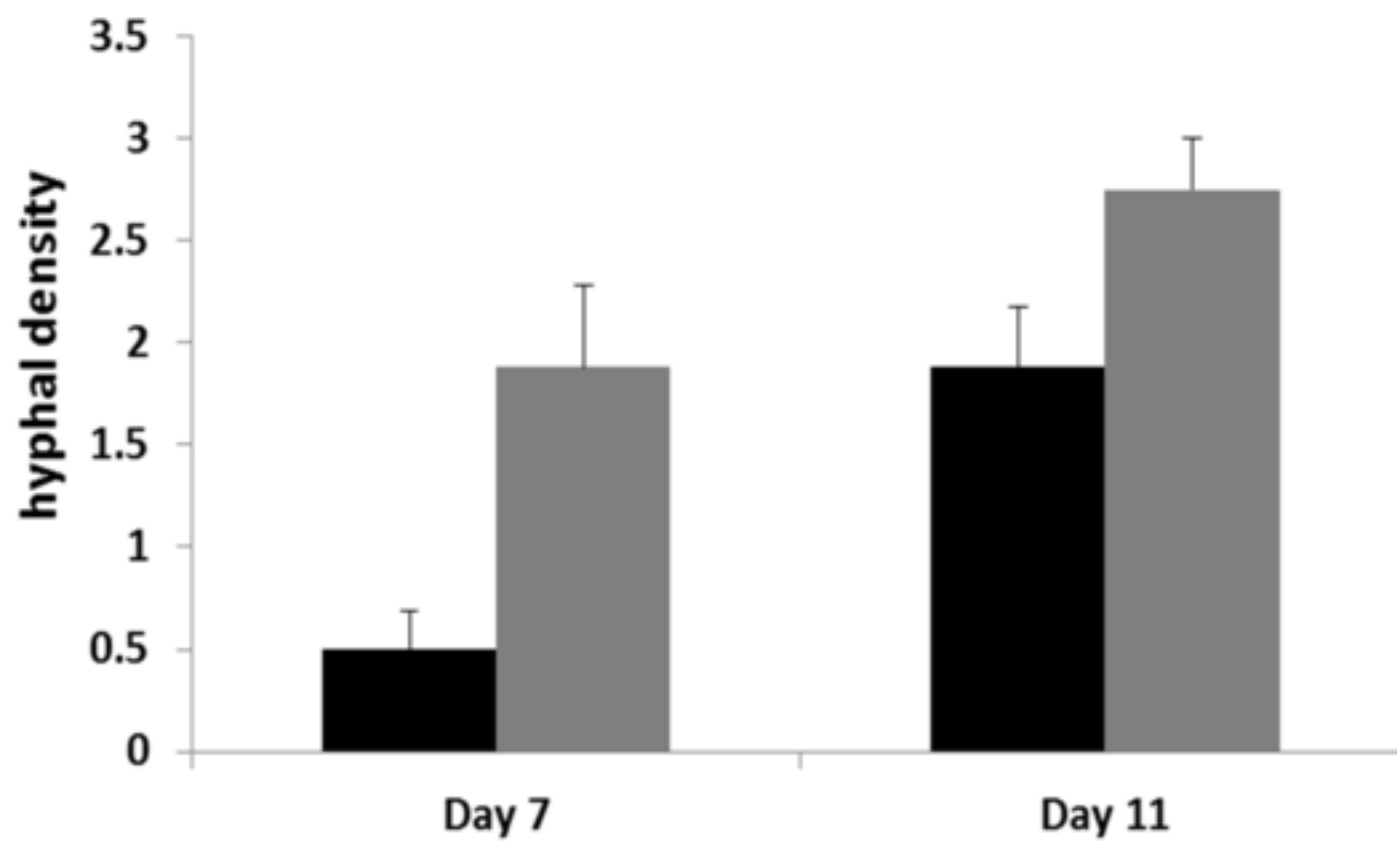
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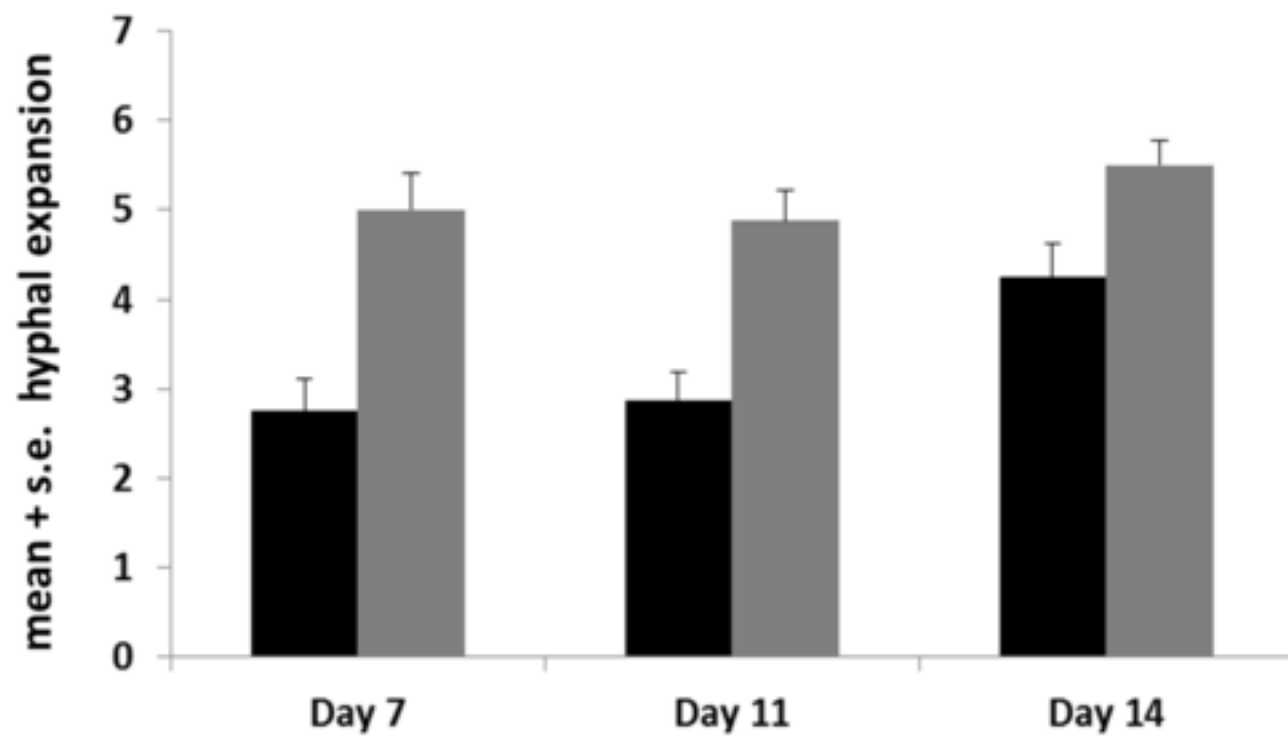


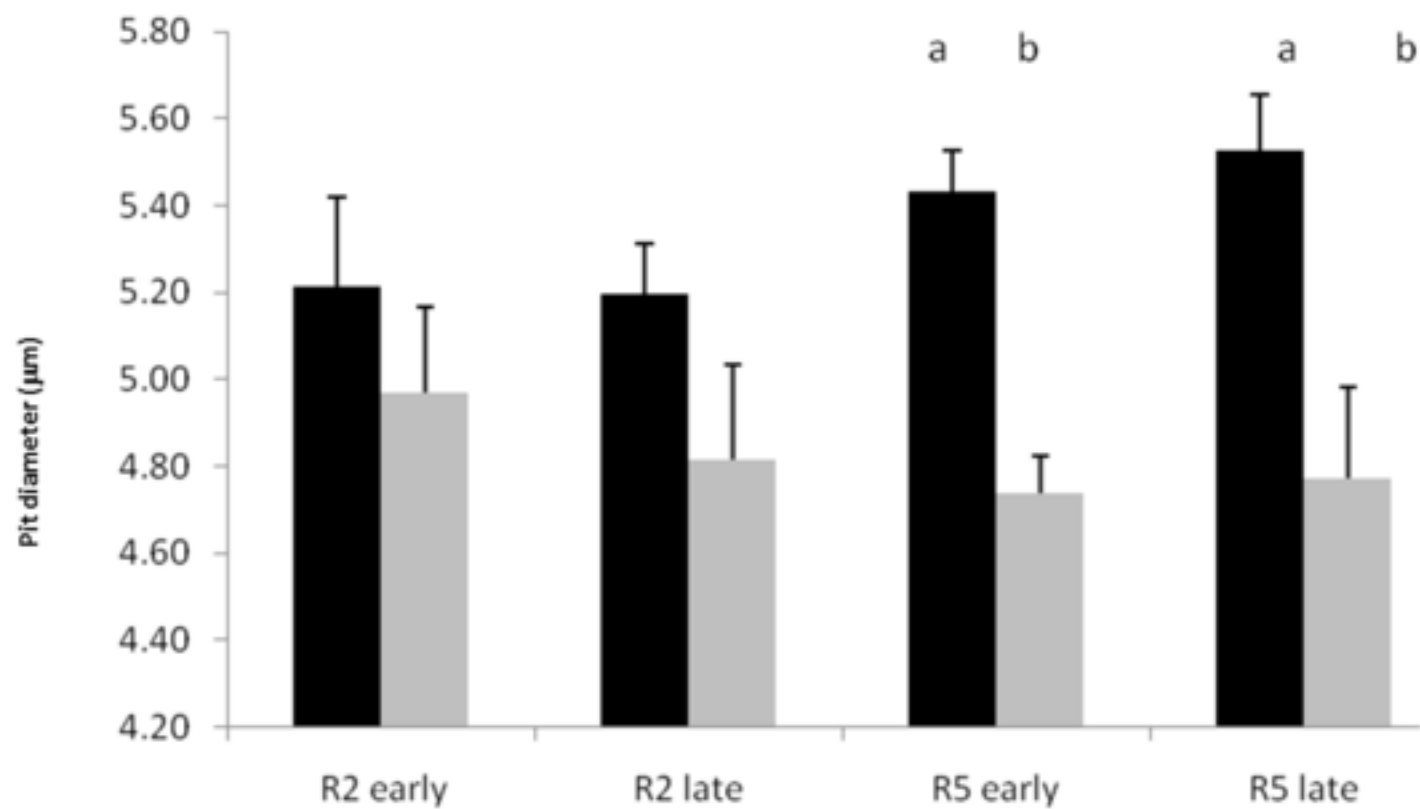
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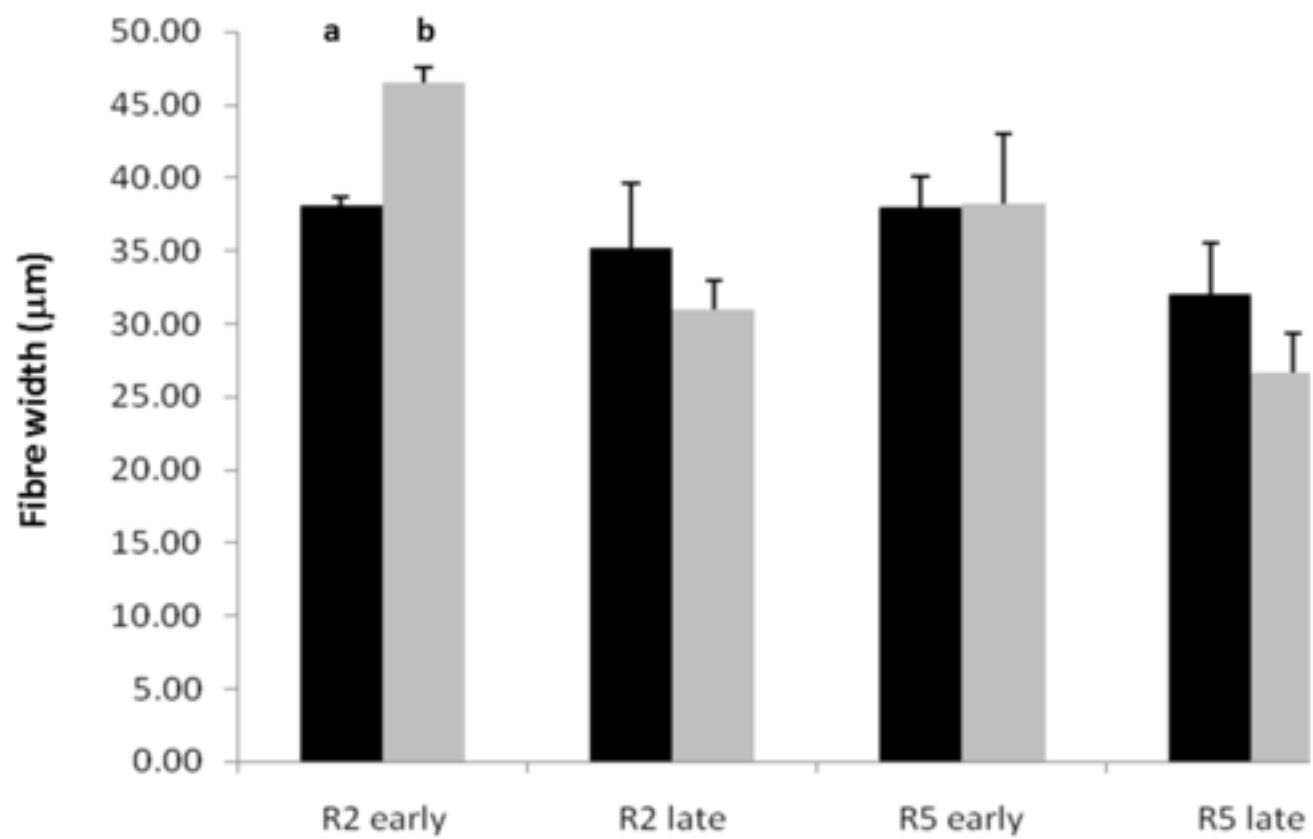


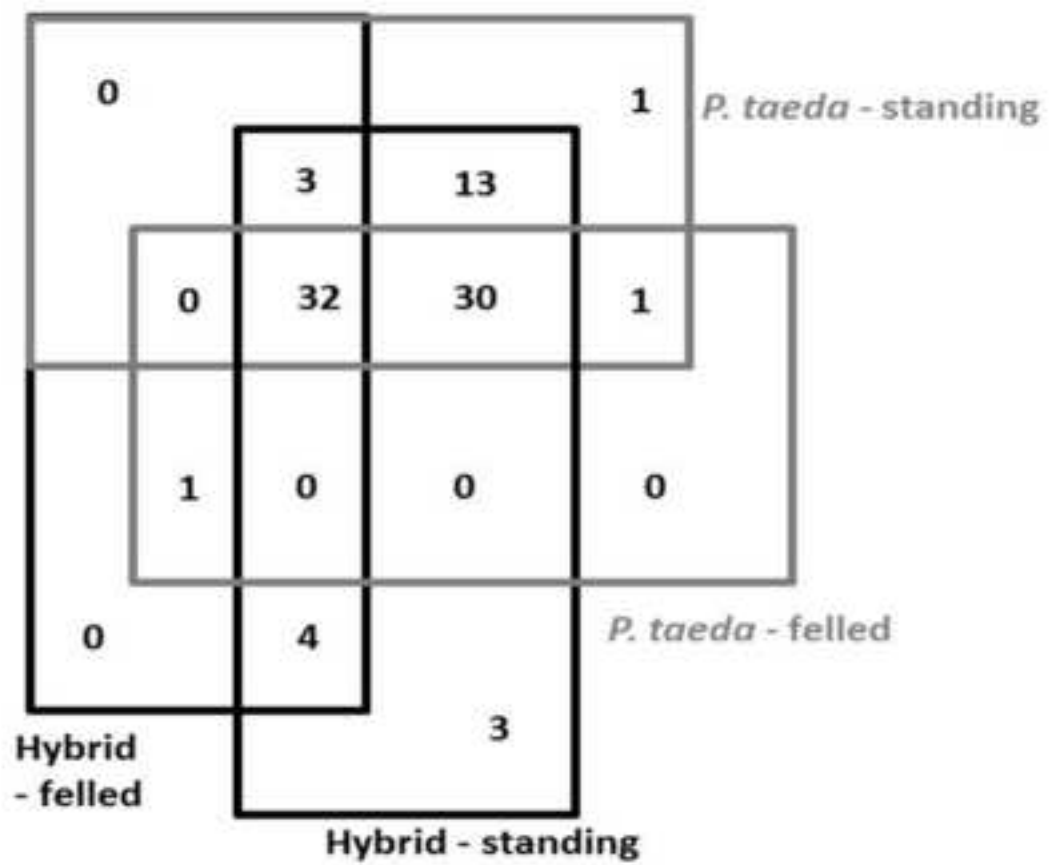












559 Table 1

	Field - standing	Field - felled	Laboratory
<i>Pinus taeda</i> (i)	0.57 ± 0.01	0.58 ± 0.02	0.62 ± 0.01
Hybrid (i)	0.55 ± 0.01	0.57 ± 0.02	0.61 ± 0.01
t-test	t ₈ = 1.86 P = 0.1	t ₈ = 0.1 P = 0.92	t ₈ = 0.31 P = 0.76
<i>Pinus taeda</i> (3)	0.45 ± 0.01	0.31 ± 0.2	0.33 ± 0.02
Hybrid (3)	0.39 ± 0.02	0.30 ± 0.03	0.34 ± 0.02
t-test	t ₈ = 10.26 P = 0.05	t ₈ = 0.06 P = 0.95	t ₈ = 0.46 P = 0.66

560

561 Table 2

Ret. Time (min)	Name	RI	Mean area hybrid	Mean area <i>P. taeda</i>	t-test
22.104	dehydroabiatic acid (DHA)	2463	23.79 ± 1.2	32.11 ± 15.0	n.s
21.768	palustic acid	2421	11.55 ± 1.7	3.15 ± 2.2	t=3.03, P=0.039
21.016	torulosol	2273	7.33 ± 1.1	0.774 ± 0.42	t=5.52, P=0.005
13.055	8-camphenemethanol	1513	1.95 ± 0.59	4.74 ± 2.5	n.s
9.205	verbenone	1218	1.61 ± 0.54	3.74 ± 1.6	n.s
20.233	unknown (MW 286)	2233	2.62 ± 0.52	0.515 ± 0.099	t=4.00, P=0.016
22.507	resin acid	2514	2.16 ± 1.0	2.46 ± 1.5	n.s.

562

563 Table 3

Ret. Time (min)	Name	RI	Mean area hybrid	Mean area <i>P. taeda</i>	t-test
22.535	callistic acid	2518	18.26 ± 18.3	25.17 ± 15.4	n.s.
22.104	dehydroabiatic acid (DHA)	2463	6.58 ± 6.6	7.76 ± 4.0	n.s.
15.527	methyl 4-hydroxy-2-methoxy-3,5,6-trimethyl benzoate	1735	4.96 ± 3.0	1.92 ± 1.3	n.s.
24.485	unknown	2786	6.90 ± 1.3	5.43 ± 2.7	n.s.
17.695	2,5-bis-(3-hydroxypropylamino)- <i>p</i> -benzoquinone	1950	5.06 ± 4.8	0.568 ± 0.34	n.s.
13.768	1,2-longidone	1576	0 ± 0	5.23 ± 2.7	n.s.
21.016	torulosol	2273	5.77 ± 1.7	1.47 ± 0.53	n.s.
23.106	methyl 7-oxodehydroabietate	2597	3.54 ± 2.6	2.12 ± 0.56	n.s.

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- 571
- 572
- Growth of *Amylosteterum areolatum*, the fungal symbiont of *Sirex noctilio*, was lower on hybrid pine substrate than on known host *Pinus taeda*.
 - Survival of *Deladenus siricidicola*, the most important biocontrol agent of *S. noctilio*, was also lower in hybrid pine than *P. taeda*.
 - Differences in secondary plant compounds may explain the observed differences in fungal growth and nematode survival.

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