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1	Defence responses of native and invasive plants to the native
2	generalist vine parasite <i>Cassytha pubescens</i> – Anatomical and
3	functional studies
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5	Evelina Facelli ^{AB} , Noriko Wynn ^{AC} , Hong T Tsang ^{AD} , Jennifer R Watling ^{AE} and José
6	M Facelli ^A
7	^A Ecology & Environmental Science, The University of Adelaide. Adelaide SA 5005,
8	Australia
9	^B Agriculture, Food and Wine, The University of Adelaide. Adelaide SA 5005,
10	Australia
11	^C Aurecon, 850 Collins St, Docklands 3008, Australia
12	^D State Key Laboratory in Marine Pollution, City University of Hong Kong, Hong
13	Kong, China
14	^E All Saints Building, Manchester metropolitan University, Manchester, M15 6BH,
15	UK
16	
17	RUNNING TITLE: Native and invasive plants' responses to a vine parasite
18	
19	Corresponding author:
20	Evelina Facelli evelina.facelli@adelaide.edu.au

21 Summary

22	We investigated the responses of two invasive and two native host species to the
23	parasitic vine Cassytha pubescens using glasshouse experiments. We assessed growth
24	of the parasite and its hosts, and anatomy and functionality of haustoria. Target hosts
25	were infected using C. pubescens already established on a donor host. This enabled
26	measurement of growth in target hosts that were detached (parasite connection
27	severed) or not from the donor host. Haustorial connections to hosts were investigated
28	using histological methods. We tested the functionality of haustoria in one invasive
29	and one native host using radiolabelled phosphorus (³² P).
30	After it was severed from the donor host, C. pubescens grew poorly on the native
31	host, Acacia myrtifolia. This was likely due to a lack of effective functional haustorial
32	development: while haustoria were firmly attached and morphologically alike those
33	formed on the other hosts, their anatomy was different: their connections with the
34	vascular system were not developed and there was no transfer of 32 P from A.
35	myrtifolia to the parasite. In contrast, the other three host species supported the
36	growth of the parasite and had fully developed haustoria. Effective transfer of ^{32}P
37	from the invasive host to the parasite confirmed this. Our results suggest a range of
38	defence mechanisms in C. pubescens hosts and are consistent with reports of strong
39	detrimental effects on invasive hosts. Further, they amount to evidence for the
40	potential use of a native parasite as biological control for invasive species.
41	
42	Keywords: parasitic plants, ³² P tracer, histology, biological control, Acacia myrtifolia,

43 Leptospermum myrsinoides, Cytisus scoparius, Cassytha pubescens

44 Introduction

45	Parasitic plants are significant components of natural vegetation worldwide.
46	They affect biodiversity and ecosystem processes and services through their negative
47	effects on native and invasive species. However, the differential responses between
48	native and invasive host species may contribute to changes in plant community
49	structure, and may be particularly useful to control invasive host species if they are
50	differentially impacted (Yu et al. 2009; Yu et al. 2011; Těšitel et al. 2020).
51	While host range in parasitic plants is well documented, variation in host
52	responses to generalist parasites has only been well studied for a few species, but has
53	been shown for both stem and root parasites (Cameron et al. 2006). Differential
54	infection rates seem to be a function of either active host selection by the parasite
55	(Hart 1990; Kelly 1992; Callaway and Pennings 1998), or differences in the
56	resistance/tolerance of hosts (Cameron et al. 2009). Despite a large host range,
57	generalist parasites tend to preferentially utilise a subset of the species available. In
58	the field this is most commonly observed as the disproportionate use of host species
59	relative to species abundance (Kelly et al. 1988; cf. Koch et al. 2004) and is
60	considered to indicate host preference by the parasite.
61	Resistance to parasitic plants includes several different mechanisms that
62	generally act to prevent establishment of a functional haustorial connection between
63	host and parasite. The extent to which haustorial development and functionality are
64	impaired varies. Host defence responses range from full resistance (where penetration
65	is prevented or impeded), to a continuum (high to nil) of tolerance responses (hosts
66	traits that reduce the effect of the parasite on host fitness) (Koskela et al. 2002;
67	Gurney et al. 2003). For example, full xylem-xylem continuity with the host is
68	achieved by Striga hermonthica attached to the tolerant host Tripsacum dactyloides,

69 while some cereal cultivars can prevent effective haustorial development of the 70 parasite (Gurney et al. 2003). Similarly, Rhinanthus minor haustoria are prevented 71 from penetrating host xylem in *Plantago lanceolata* and *Leucnathemum vulgare* 72 because of extra lignification or hypersensitive responses in the hosts (Cameron et al. 73 2006; Cameron and Seel 2007). Use of isotope tracing showed that R. minor had only 74 very limited access to nutrients from these hosts, confirming the lack of full 75 functionality of the haustoria (Cameron and Seel 2007). 76 The Australian parasitic vine Cassytha pubescens R.Br. is a generalist that 77 grows on a wide range of species, usually spreading and attaching to a large number 78 of individuals of different species. Field surveys in areas with native and invasive 79 species, demonstrated that infection by C. pubescens was somewhat disproportionate 80 to species availability, indicating slight or no host preference by the parasite (Prider et 81 al. 2009; Supplementary Material Table S1; Figure S1). Pot experiments showed that 82 when placed between a known host, an artificial plant and an empty space C. 83 pubescens did not grow preferentially in any direction (Noriko Wynn unpublished 84 data). This suggests that unlike other parasitic vine species (e.g. Cuscuta spp, Kelly 85 1992; Runyon et al. 2006), C. pubescens does not appear to detect the presence of nearby hosts. 86

We investigated the associations between *C. pubescens*, two invasive hosts
(*Cytisus scoparius* (L.) Link and *Ulex europaeus* L.) and two native hosts (*Acacia myrtifolia* (Sm.) Wild. and *Leptospermum myrsinoides* Schltdl.). We examined
growth of both the parasite (host use) and its hosts (host responses), and the anatomy
of haustoria on each host. Further, we tested the functionality of the haustorial
connections in one invasive (*C. scoparius*) and one native species (*A. myrtifolia*)
using radiolabelled soil phosphorus (³²P).

94 Materials and Methods

95 *Plant species*

Cassytha pubescens (Lauraceae) is a perennial, rootless, stem-twining, hemi-parasitic 96 97 vine native to southern Australia. Its leaves are reduced to scales, but the stem 98 contains chlorophyll and is capable of photosynthesis (Abubacker et al. 2005; Prider 99 et al. 2009). Cassytha pubescens is an obligate parasite, and has to attach to a host 100 within 6 weeks of germination to survive (McLuckie 1924). It has a wide host range 101 including many native Australian woody perennials and also non-native invasive 102 perennial shrubs (Prider et al. 2009; Supplementary Material Table S1). Although 103 morphologically similar to the well-studied parasitic vine Cuscuta spp. 104 (Convolvulaceae), the life strategy is quite different. Whereas Cuscuta is a genus of 105 annual holoparasites, in which the stem contains little or no chlorophyll (Kuijt 1969; 106 Allen and Allen 1990), C. pubescens is a perennial hemiparasite that spreads mostly 107 through vegetative growth, growing across branches within a host and spreading from 108 one plant to another, often connected to several individuals of different species. 109 The woody perennial hosts tested in different experiments were two invasive shrubs, 110 Cytisus scoparius (Fabaceae) and Ulex europaeus (Fabaceae), and two native shrubs 111 Acacia myrtifolia (Fabaceae) and Leptospermum myrsinoides (Myrtaceae). Cytisus 112 scoparius and U. europaeus were apparently introduced in the early 1800 as hops 113 substitute (the former) and garden plants (Waterhouse 1988; Ireson et al. 2003). Both 114 species are listed as Weeds of National Significance (Australian Weeds Committee 115 2012). The distribution of the four species overlaps with that of the parasite in South 116 Australia in the open sclerophyll woodlands of the Mt Lofty Ranges around Adelaide. 117 In these woodlands, we found C. scoparius, A. myrtifolia and L. myrsinoides to be 118 amongst the species on which C. pubescens was most abundant and its haustoria were 119 firmly attached (Supplementary Material Figure S1). In field and glasshouse studies, 120 C. pubescens has been shown to have strong negative effects on the growth of U. 121 europaeus and C. scoparius but not on the native shrub L. myrsinoides (Prider et al. 122 2009; Cirocco et al. 2016, 2017, 2018). Presently there is no information about the 123 ecophysiological responses of A. myrtifolia to the parasite. Field observations 124 (summarised in Supplementary Material) report haustoria (morphologically alike 125 those formed on other species) firmly attached, and large amounts of the parasite 126 growing on it. However, the surveys did not determine if the parasite was also 127 connected to other surrounding hosts that could have been supporting its growth. A 128 greenhouse experiment (Tsang 2010) found that shortly after the connections of C. 129 pubescens with the donor host were severed, the parasite growing on A. myrtifolia 130 died. 131 Unless otherwise stated, all plant material (seeds, collected plants etc.) used in 132 our study came from the same area in the Mt Lofty Ranges. The native host species 133 were sourced from a local nursery (Native Flora, SA) and the invasive species 134 obtained from stock grown by the Terrestrial Plant Ecology Laboratory, The

135 University of Adelaide.

136

137 Experiment 1 – Growth of parasite and hosts

138 Experimental set up

139 Twenty-four individuals each of *L. myrsinoides*, *A. myrtifolia*, *U. europaeus* and *C.*

140 *scoparius* were grown in 140 mm pots filled with native potting mix and a slow

141 release native fertiliser (Osmocote, Scotts-Sierra Horticultural Products, Marysville,

142 OH, USA), supplied at the recommended dosage, in a greenhouse in Adelaide.

143 Sixteen individuals of each species (target hosts) were infected using tendrils from C.

144 pubescens growing on eight C. scoparius plants (donor host) (Shen et al. 2010). Two 145 individuals from each species were placed randomly around each infected C. scoparius donor plant and C. pubescens tendrils were trained onto the new host. Eight 146 147 uninfected individuals of each target host species acted as controls. Plants were misted 148 twice daily for ten minutes and temperatures within the greenhouse maintained at 149 approximately 23°C. After three months, the connection between C. pubescens 150 growing on the donor host and one of the target hosts of each species was severed. 151 The target hosts by then had well established growth of C. pubescens with well 152 attached haustoria. This created three treatments: detached (parasite connected to 153 target host only), connected (parasite connected to donor and target hosts) and control 154 (uninfected target hosts). The detached treatment examined the growth of C. 155 pubescens (and corresponding host) when growing on a single host. The connected 156 treatment examined parasite growth (and corresponding host) when utilising the 157 resource from two hosts: C. scoparius-A. myrtifolia, C. scoparius-C. scoparius, C. 158 scoparius-L. myrsinoides and C. scoparius-U. europaeus. 159 160 Data collection and analyses 161 After five months the shoot biomass of all host plants and the parasite was harvested. 162 When C. pubescens was separated from the host plants, the total number of haustoria

163 formed and the number of haustoria with firm connection to the host stem were

164 recorded. Parasite biomass was separated into dead and living material. Host and

165 parasite tissue were dried for 96 hours at 80 °C then weighed. ANOVAs were applied

166 to parasite biomass (species, four levels; treatment, two levels: connected and

167 detached) and host biomass (species, four levels; treatment: three levels: connected,

168 detached and control) using JMP 7 (SAS Institute). The Tukey-Kramer HSD test was

169 used to compare means where the effects of treatments were significant.

170

171 Experiment 2 - Haustoria formation – histology

The anatomy of haustoria of C. pubescens growing on the four different host species 172 173 was studied using light microscopy. Haustoria from stems with a minimum infection time of ten weeks and a maximum stem diameter of 3 mm were harvested from three 174 175 healthy individuals of U. europaeus, C. scoparius, A. myrtifolia and L. myrsinoides 176 grown as described in experiment 1. Specimens were preserved in 2% glutaraldehyde 177 and 2.5% paraformaldehyde in 0.1 M phosphate buffer (pH 7.2), at 4 °C for four 178 weeks to allow the fixative to penetrate the plant tissue. Specimens were then washed 179 in 100% ethanol and dehydrated in a graduated ethanol series for 40 minutes in each 180 70%, 90% and 100% ethanol under vacuum. The haustoria were left under vacuum 181 for 12 hours in a 1:1 solution of 100% ethanol and LR-White resin. Samples were 182 embedded in 100% LR-White resin after being placed in resin for 84 hours under 183 vacuum with resin changes every 12 hours and then set in gelatine capsules for 48 184 hours at 80 °C. Three haustoria from each species were cut into sections transverse to 185 the stem of the host, 2 to 4 µm thick (Leica Ultracut E Ultramicrotome). Sections 186 were floated onto slides, placed on an 80 °C hotplate and stained on the hotplate using 187 1 % Toluidine blue O in boric acid. Sections were examined under a light microscope 188 (Olympus BX51) fitted with a camera (Colorview III Camera).

189

190 Experiment 3 - Functionality of haustoria – Transfer of radiolabelled P

191 To test functionality of firmly attached haustoria of A. myrtifolia and C. scoparius we

192 compared transfer of 32 P between pairs of hosts connected by *C. pubescens* (Fig 1).

195 Ten seedlings of C. scoparius were collected from a field site near Adelaide (35°) 196 0'58.08"S, 138°45'58.45"E), South Australia. The seedlings were placed in 1.5 L pots 197 with sandy loam soil, in a greenhouse for two months until established. Ten seedlings 198 of A. myrtifolia were grown in 1.5 L pots in a greenhouse for six months. All plants 199 were watered as required. The C. scoparius plants were infected with C. pubescens by placing them next to an already infected C. scoparius and directing the tendrils of the 200 201 parasite to the stem of the target seedlings (as described above; Shen et al. 2010). 202 After approximately three months, the connections between the donor host and the 203 target seedlings were severed and the 10 newly infected C. scoparius plants used to 204 similarly infect one plant each of A. myrtifolia. The pots containing A. myrtifolia 205 plants were left for 10 weeks next to the infected C. scoparius plants to allow the 206 haustoria of C. pubescens to develop. All plants were watered with 250 mL of reverse 207 osmotic (RO) water three times a week and received 290 mL of full strength Hoagland's solution in the 4th week. To increase the phosphorous requirements in the 208 209 hosts, in the 8th week all pots received the same amount of Hoagland's solution but with only one fifth the amount of phosphate. In the 11th week, the 10 pairs of hosts, all 210 211 having several haustoria of the parasite firmly attached to both plants, were randomly assigned to two treatments (five pairs per treatment): 1) radioactive phosphate (^{32}P) 212 injected into the soil of pots containing the C. scoparius host or 2) ³²P injected into 213 214 the soil of pots with the A. myrtifolia host (Fig. 1). Each injected pot received 6 MBq of radioactive phosphate (carrier-free H₃³²PO₄) dissolved in 125 mL of RO water, 215 216 divided into 5 aliquots of 25 mL each. Each aliquot was injected using a syringe with 217 a 10 cm needle into 5 different locations in each pot to maximize the chance of it 218 being absorbed by the host. Two weeks after injection, each pair of plants and their

219	parasite were harvested and divided into the following components: 1) host shoot
220	from the pot injected with ³² P, 2) <i>C. pubescens</i> growing on the radio-labelled host, 3)
221	C. pubescens spanning between the two hosts, 4) C. pubescens on the non-labelled
222	host, 5) infected shoot of the non-labelled host, and 6) uninfected shoot of the non-
223	labelled host (Fig. 1). Plant material was dried for 2 days at 70 °C and then ground to
224	a fine powder. For each replicate, 5 mL of nitric acid was added to 0.5 g of ground
225	plant material in a test tube, and digested overnight in a heat block at 140 °C (Hanson
226	1950). Acacia myrtifolia digests were centrifuged at 2000 rpm for 10 minutes to
227	remove a milky gelatinous residue. Radioactivity was determined using 2 mL aliquots
228	of the digests in a liquid scintillation counter (Wallac 1215 RackBeta II) by measuring
229	the Cerenkov radiation produced by beta particles without any scintillation fluor
230	cocktail and corrected for decay (L'Annunziata 1997).
231	
232	Data analysis
233	One-way ANOVAs were performed using Graphpad Prism 5 for Windows, GraphPad
234	Software, La Jolla California USA, <u>www.graphpad.com</u> .
235	
236	Results
237	Experiment 1 – Growth of parasite and hosts
238	The amount of live biomass of C. pubescens was influenced by both treatment
239	and species (ANOVA _{interaction} : $F_{3, 32} = 2.93$, $P = 0.049$). Live parasite biomass was
240	significantly lower growing on a single A. myrtifolia individual than when growing on
241	C. scoparius and A. myrtifolia simultaneously (Fig. 2). The growth of the parasite in
242	the detached treatment was greatest on C. scoparius, and significantly higher than on
243	either A. myrtifolia or U. europaeus but not L. myrsinoides (Fig. 2). Live C. pubescens

- biomass supported by two hosts was greatest on *A. myrtifolia*, followed by *C*.
- scoparius, L. myrsinoides and U. europaeus. Only the live biomass on U. europaeus
- 246 was significantly different from A. myrtifolia (Fig. 2). Treatment did not influence the
- amount of dead parasite biomass (ANOVA: $F_{1,32} = 1.07$, P = 0.31), however C.
- 248 *pubescens* growing on *A. myrtifolia* had more dead tissue than any of the other species
- 249 (ANOVAspecies: $F_{3,32} = 14.16, P \le 0.0001$; Fig. 2).
- Host biomass differed between species (ANOVA: $F_{3, 48} = 128.0, P \le 0.0001$).
- 251 A. myrtifolia had the highest biomass followed by C. scoparius, L. myrsinoides and U.
- 252 europaeus (Fig. 3). Plants in the connected treatment had lower biomass than plants in
- either the detached or control treatments (ANOVA: $F_{2,48} = 7.48$, P = 0.002).
- 254 No differences were observed between treatments or species for either total
- number of haustoria on each host (ANOVA_{species}: $F_{3, 72} = 1.61$, P = 0.194;
- 256 (ANOVA_{treatment}: $F_{1, 72} = 1.93$, P = 0.17), or the proportion of haustoria attached to the
- 257 host stems (ANOVA_{species}: $F_{3, 72} = 1.61$, P = 0.3448; ANOVA_{treatment}: $F_{1, 72} = 1.93$, P =
- 258 0.45). Cassytha pubescens biomass was correlated with the proportion of haustoria
- that were considered to be well attached and therefore viable ($R^2 = 0.22$, Pearson two
- 260 tailed test, P = 0.001; Fig. 4).
- 261
- 262 *Experiment 2 Haustoria formation histology*

263 Representative sections from the sectioned haustoria from each species are presented.

- All sections from the three plants per species showed the same anatomical
- 265 characteristics. The haustoria formed on the two invasive species, U. europaeus, and
- 266 C. scoparius had endophytes capable of penetrating host tissue. Parasite tissues are
- 267 clearly observed entering the host and growing in close contact with host vascular
- structures (Fig. 5). Endophyte of C. pubescens growing on C. scoparius widens after

269 penetrating the host forming an oval like structure within host tissue (Fig. 5*b*, E). A 270 large proportion of the endophyte tissue is in close contact with the host xylem. The 271 early stages of a vascular core are evident, running through the middle of endophyte 272 into the haustorial tissue (Fig. 5*a*, IV). It appears that growth of the endophyte 273 structure has spread increasing the surface area in contact with host vasculature (Fig. 274 5*b*, I).

The anatomy of endophytes formed on *U. europaeus* was different for each of the haustoria sectioned. Yet all were able to penetrate host tissues and contact host vascular structures (Fig. 5c, d, I). As with the haustoria formed on *C. scoparius*, there was evidence of the formation of a vascular core in dense differentiating parenchyma cells running through the central body of the endophyte (Fig. 5c, IV). The cells of the endophyte were darkly stained and appeared to form dense tissue (Fig. 5d, DT).

When grown on native host species, *C. pubescens* was able to form apparently functional haustoria on *L. myrsinoides* (Fig. 6*a*) but was prevented from entering host tissues when growing on *A. myrtifolia*. In the haustoria formed on *L. myrsinoides* the endophyte had clearly penetrated the host tissues and formed direct luminal contact with host xylem via the differentiation of xylem (Fig. 6*b*, PX). There is also evidence of a hyaline rich body of cells located in the centre of endophyte tissue. In contrast, *C. pubescens* growing on *A. myrtifolia* was prevented from entering host

tissue at the cortex, although an endophyte is present (Fig. 6c, d). There was evidence

289 of thickening host tissue where the endophyte attempted to enter the host tissue (Fig.

 $290 \quad 6c, d, T$). At the interface between host and parasite (Fig. 6d, I), there are darkly

stained tissues; these clearly delineate the barrier between host and parasite tissues.

292 There is no evidence of a vascular core or differentiated xylem in the body of the

293 haustoria.

294 Experiment 3 - Functionality of haustoria – Transfer of radiolabelled P

295 There were significant differences in the radioactivity of plant components between the two treatments. When ³²P was injected into pots containing C. scoparius, the same 296 297 level of radioactivity was detected in both C. scoparius and in C. pubescens, but only trace amounts were detected in the paired A. myrtifolia (ANOVA: $F_{1,2} = 12.17$, P =298 0.001; Fig. 7*a*). This contrasted with the distribution of 32 P when it was injected into 299 300 pots containing A. myrtifolia. In this case, radioactivity was detected in A. myrtifolia 301 but only traces were detected in C. pubescens and C. scoparius (ANOVA: $F_{1,2}$ = 302 10.07, *P* = 0.003; Fig. 7*b*). 303 304 Discussion 305 Regardless of the presence of attached haustoria and the growth of the parasite on A. 306 *myrtifolia*, this native host resisted penetration by the parasite. In contrast, haustoria 307 on the invasive species and on the other native species (L. myrsinoides) were able to penetrate host tissues successfully and, in C. scoparius, supported transfer of ³²P 308 309 between host and parasite. Importantly, the relative lack of severe or lethal negative 310 effects on L. myrsinoides (compared with invasive species) (Prider et al. 2009; 311 Cirocco et al. 2015) occurs in spite of the fully developed anatomical connections we 312 documented. This suggests that there is a range of defence mechanisms amongst hosts 313 of C. pubescens. 314 315 Growth of C. pubescens on A. myrtifolia

316 Field studies have reported that *C. pubescens* is able to successfully grow on *A*.

317 *myrtifolia*, and even that this is one of the species on which the parasite is more

318 abundant (Supplementary Material Table S1). In our experiments as in field

observations we found that *C. pubescens* haustoria were as firmly attached to *A. myrtifolia* as to the other hosts. However, *C. pubescens* did not grow in high densities
on *A. myrtifolia* unless it was also still attached to the donor host. Further, there was
large accumulation of dead biomass on the detached plants. These results, indicate
that the parasite was unable to effectively use *A. myrtifolia* as a host.

324 The anatomical studies showed that A. myrtifolia exhibited resistance by 325 preventing the penetration of the parasitic endophyte. The localisation of the defence 326 response indicates resistance is induced by contact and attempted penetration of host 327 tissues by the parasite. During haustorial formation C. pubescens excretes a fluid 328 which helps the parasite invade host tissues by the formation of an adhesive disk 329 (Heide-Jørgensen 1991). This attachment mechanism is also observed in the 330 formation of prehaustoria by Cuscuta spp. (Kaiser et al. 2015). Contact with this fluid 331 may trigger the thickening of the cortical tissue in A. myrtifolia stems at the site of 332 attempted parasite penetration. The parasitic vine *Cuscuta pentagona* was similarly 333 prevented from penetrating the cortex of tomato varieties (Goldwasser et al. 2017). 334 Resistance in tomato has been since attributed to hormonal signalling triggered by the 335 parasite (Runyon et al. 2010). Studies of the root parasite, Orobanche spp., which is also prevented from penetrating tissues of resistant hosts beyond the cortex, show 336 337 that the production of toxic phenols (Serghini et al. 2001), reinforcement of host cell 338 walls, deposition of callose and suberisation (Perez-de-Luque et al. 2005; Echevarría-339 Zomeño et al. 2006) contribute to host resistance.

341 *pubescens* was grown on *A. myrtifolia*, probably explains the inability of the parasite

The lack of well-developed haustorial structure that we observed when C.

342 to acquire 32 P from this host. This confirms that *A. myrtifolia* prevents the

340

343 development of functional connections by the parasite. Our results are similar to those

reported for the root hemiparasite *R. minor*, which absorbed different amounts of ¹⁵N when grown on hosts with different degrees of defence responses (Cameron and Seel 2007). Similar to our results, host resistance mechanisms prevented the parasite from establishing functional connections with host vascular tissues. Further, the concentration of ¹⁵N taken up from tolerant hosts was positively correlated with parasite biomass, providing additional evidence of the importance of functional haustorial connections for parasite growth (Cameron and Seel 2007).

351 Biomass of C. pubescens was higher when growing on A. myrtifolia still 352 connected with the donor host, than on the detached plants. Given the lack of 353 functional haustoria when growing on A. myrtifolia, the parasite must have been 354 mostly relying on resources from the donor host, C. scoparius. This characteristic 355 complicates the study of host use by C. pubescens, because potentially masks native 356 host resistance or tolerance as it gives C. pubescens the appearance of an ability to 357 form functional haustoria and grow on resistant species such as A. myrtifolia. As a 358 result resistance or tolerance to C. pubescens may be more widespread than the host 359 range of the parasite suggests. Some native species, like A. myrtifolia, which could be 360 considered 'pseudo-hosts', may only provide physical support for the parasite, while 361 it moves between gaps of suitable hosts (Marquardt and Pennings 2011). While C. 362 *pubescens* possibly obtains little or no nutrients from these 'pseudo-hosts', they may 363 provide physical support to photosynthetic stems and facilitate its dispersal by 364 vegetative means to suitable hosts.

365

366 *Growth of* C. pubescens *on* C. scoparius, U. europaeus *and* L. myrsinoides

367 Comparable amounts of dead and live parasite tissue in the detached and connected

368 treatments on C. scoparius, U. europaeus and L. myrsinoides, demonstrates similar

369 parasite performance on these species. This corresponds with the anatomical

similarities we observed in the development of the haustoria on these hosts. Further,
the transfer of ³²P through the haustoria from the host *C. scoparius* to *C. pubescens*confirmed the physiological functionality of these haustoria. Generally, there is a
strong association between biomass of the parasite and the transfer of resources and/or
number of haustoria attached (Kelly 1992; Cameron and Seel 2007) as we observed in
our first experiment (but see discussion about *A. myrtifolia* above).

376 Cassytha pubescens formed fully developed haustoria on the infected native L. 377 myrsinoides, which also had lower biomass when infected by the parasite. Previous 378 studies have also reported lower biomass and even some negative physiological 379 effects on L. myrsinoides but detrimental effects of C. pubescens have been always of 380 lower magnitude than on invasive hosts in glasshouse and field conditions (Cirocco et 381 al. 2016, Prider et al. 2009). These effects could be attributed to incomplete haustorial 382 connections (Cameron and Seel 2007) and/or adaptive tolerance mechanisms 383 (Mutikainen et al 2000). Our results allow us to rule out the first alternative. Cirocco 384 et al. (2015) proposed that the ability of L. myrsinoides to maintain photoprotective capacity/engagement when infected by C. pubescens, thereby preventing 385 386 photodamage, could explain this host's tolerance. Its adaptations to low availability of 387 water and nutrients, characteristic of plants in the sclerophyll woodlands of South 388 Australia which contrast with the higher resource requirements of invasive species, 389 may also contribute to its higher tolerance to reduction in resources produced by the 390 parasite (Li et al. 2012). Another native host, Acacia paradoxa, also shows tolerance 391 to C. pubescens; it supports parasite growth but host photosynthesis is not affected 392 (Cirocco et al. 2017). Other native species have been observed to support the parasite 393 (Prider et al. 2009; Supplementary Material Table S1, Figure S1). On the other hand,

394 our results on *A. myrtifolia* open the possibility that some of those species may

395 partially or completely prevent formation of functional of haustoria by the parasite,

and thus also be 'pseudo hosts'. Further research is required to determine the

397 functionality of haustoria, and parasite performance on these species, along with host

398 physiological responses to infection. This would inform our understanding of

399 ecological responses of the parasite and its many hosts (or pseudo hosts).

400

401 *Overall implications*

402 Our results suggest that the parasite does not selectively utilise invasive species over
403 natives. This generalist strategy allows the parasite to become established on host
404 species with which it has not coevolved (Koch *et al.* 2004). Importantly, however,
405 differences in resistance or tolerance of the native and invasive hosts to the parasite
406 could then induce changes in plant community structure and diversity (Yu *et al.* 2011;
407 DiGiovanni *et al.* 2017).

408 The differences in defence responses between the native and invasive hosts 409 reported here, albeit based on a small number of species, are overall consistent with 410 the prediction of the biotic resistance hypothesis (Těšitel et al. 2020). According to 411 this interpretation, we could speculate that the two native hosts have evolved in the 412 presence of the parasite and over time have developed suitable and different, 413 mechanisms to resist/tolerate infection (Li et al. 2012; Cirocco et al. 2016). In 414 contrast, the two invasive hosts, which were introduced to Australia less than 200 415 years ago, have not evolved defence mechanisms capable of resisting infection by the 416 novel enemy. Our results suggests a broad spectrum of responses of the native plants 417 to the native parasite. Confirming this will require a more comprehensive assessment 418 of anatomy and function of haustoria formed on native and invasive hosts, which was 419 beyond the scope of our study. In addition, it will be important to determine if 420 resistance/tolerance is variable at several levels, i.e. individuals and populations, and 421 if this variation is associated with previous coexistence, and hence coevolution, of the 422 parasite and the host (e.g. Jerome and Ford 2002). 423 If differential responses between native and invasive species are proven valid 424 for this type of vegetation, C. pubescens could be used as an important agent for 425 biological control in the area (Li et al. 2012; Těšitel et al. 2020). Species used for 426 biological control generally have high host specificity so that only the target pest is 427 affected by the introduction of the species into a system (Myers and Bazely 2003). 428 However, this is generally applied when introducing a further non-indigenous species 429 into a system. The use by augmentation of a native parasite already present in the 430 system provides a novel way to aid in control of introduced species, because infection 431 by C. pubescens of invasive species has a greater effect on host health, biomass and 432 fecundity than on the native species so far tested (Prider et al. 2009; Cirocco et al. 433 2016, 2018). This suggests that if used as a biological control the parasite will have 434 little or no significant effects on native species within the system (Heer et al. 2018). Further, our ³²P tracer technique enabled us to assess the degree of host 435 436 defence responses to C. pubescens (similarly to the study on a root parasite of 437 Cameron et al. 2006), but could also be extended for similar experiments with other 438 stem parasites, such as the economically important Cuscuta. This technique also 439 provides the potential to determine the relative contribution of multiple hosts 440 simultaneously parasitised by twining stem parasites such as C. pubescens, by 441 applying different tracers to the various hosts. Conversely, the impact of the parasite 442 on its multiple hosts could also be determined.

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453 **Conflicts of Interest**

- 454 The authors declare no conflicts of interest.
- 455

456 **References**

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585 Figure legends

586 Figure 1. Experimental design showing the pot containing either *Cytisus scoparius* or

587 *Acacia myrtifolia* injected with ³²P (radiation symbol) and the various components

- 588 harvested separately for 32 P analyses: (1) host shoot from the pot injected with 32 P, (2)
- 589 Cassytha pubescens on the radio-labelled host, (3) C. pubescens spanning the two
- bosts, (4) C. pubescens on the non-labelled host, (5) infected shoot of the non-labelled
- 591 host and (6) uninfected shoot of the non-labelled host.
- 592 Figure 2. Live (a) and dead (b) biomass (g) of Cassytha pubescens when grown on
- 593 Acacia myrtifolia (Acacia), Cytisus scoparius (Cytisus), Leptospermum myrsinoides
- 594 (Leptospermum) or *Ulex europaeus* (Ulex) and exposed to two treatments, connected
- to or detached from donor host. Mean + s.e. (n = 8). Different letters indicate means

596 are significant different. Tukey-Kramer HSD, $\alpha = 0.05$.

- 597 Figure 3. Shoot biomass (g) of Acacia myrtifolia (Acacia), Cytisus scoparius
- 598 (Cytisus), *Leptospermum myrsinoides* (Leptospermum) and *Ulex europaeus* (Ulex)
- after infection by *Cassytha pubescens* for five months in the following treatments:
- 600 connected to donor host (filled bars), detached from donor host (hatched bars) and
- 601 control, non-infected (clear bars). Mean + s.e. (n = 8). Different letters indicate
- 602 significant differences between species. * connected treatment significantly different
- from detached and control. Tukey-Kramer HSD, $\alpha = 0.05$.
- 604 Figure 4. Relationship between *Cassytha pubescens* biomass and the percentage of
- 605 viable haustoria over total haustoria when grown on Acacia myrtifolia (Acacia,
- 606 circles), Cytisus scoparius (Cytisus, squares), Leptospermum myrsinoides
- 607 (Leptospermum. triangles) or *Ulex europaeus* (Ulex, diamonds) and exposed to two
- treatments, connected (black symbols) to or detached (white symbols) from donor
- 609 host.

- 610 Figure 5. Light microscopy of *Cassytha pubescens* haustoria on (a) Cytisus scoparius
- at x 4 magnification, (b) C. scoparius at x 10 magnification, (c) Ulex europaeus at x

612 10 magnification and (d) U. europaeus at x 20 magnification. H, haustoria, HS, host

- 613 stem, PS, parasite stem, E, endophyte, HX, host xylem, PX, parasite xylem, I,
- 614 interface between host and parasite, IV, initial vascular core formation, DT, darkly
- 615 stained tissue, CL, collapsed layer, HB, hyaline body. Slides stained with 1 %
- 616 Toluidine blue O solution. Scale bars equal 1000 μm at x 4 magnification, 500 μm at
- 617 x 10 magnification and 200 μm at x 20 magnification.
- 618 Figure 6. Light microscopy of *Cassytha pubescens* haustoria on *(a) Leptospermum*
- 619 myrsinoides at x 10 magnification, (b) L. myrsinoides at x 20 magnification, (c)
- 620 Acacia myrtifolia at x 4 magnification and (d) A. myrtifolia at x 10 magnification. H,
- haustoria, HS, host stem, PS, parasite stem, E, endophyte, HX, host xylem, PX,
- 622 parasite xylem, T, thickening of tissue, I, interface between host and parasite, IV,
- 623 initial vascular core formation, DT, darkly stained tissue, CL, collapsed layer, HB,
- 624 hyaline body. Slides stained with 1 % Toluidine blue O solution. Scale bars equal
- 625 1000 μm at x 4 magnification, 500 μm at x 10 magnification and 200 μm at x 20
- 626 magnification.
- 627 Figure 7. Radioactivity (kBq gP⁻¹) in the various plant components (see Figure 1 for
- 628 details of the experimental setup) when the pot containing either *Cytisus scoparius (a)*
- 629 or Acacia myrtifolia (b) was injected with 32 P. Means + s.d. (n=5). Different letters
- 630 indicate significant differences between plant components ($P \le 0.05$). Note different
- 631 scales for both graphs.





















654 Figure 7