

Effect of humidity and temperature on the performance of three strains of Aphalara itadori, a biocontrol agent for Japanese knotweed

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- 1 Effect of humidity and temperature on the performance of three strains of Aphalara
- 2 itadori, a biocontrol agent for Japanese Knotweed
- 3
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35 Highlights

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- Three strains of *Aphalara itadori* were tested under two environmental conditions
- More stressful environmental conditions slowed down psyllid development
- 39
- Biocontrol effectiveness was similar among strains, with no clear hybrid advantage
- 40

41 Abstract

Japanese knotweed (Fallopia japonica) is a highly damaging invasive species affecting UK 42 infrastructure and biodiversity. Under laboratory conditions, the psyllid Aphalara itadori has 43 demonstrated its potential to be a successful biocontrol agent for F. japonica. However, this 44 45 potential has not materialised in the field where long-term establishment of A. *itadori* has been unsuccessful and faces the added challenge of climate change. Intraspecific variation (variation 46 47 among individuals of a species) has been shown to support establishment in alien species and improve resilience to changing environmental conditions, here we propose it could improve 48 the performance of biocontrols. To test this possibility we compared the performance and 49 impact on F. japonica of three strains of A. itadori with different genetic backgrounds, 50 51 including a newly created hybrid. We hypothesize that genetic variability would be increased in hybrids resulting in greater biocontrol effectiveness (greater impact on plant growth). We 52 also explored the potential influence of changing climate in performance, testing all strains 53 54 under two humidity conditions (with the same temperature). Contrary to our expectation, the hybrid strain had the worst performance (slowest development rate and lower survival from 55 egg to adult emergence) under both environmental conditions. Exposure to different strains of 56 A. *itadori* did not result in consistent differences in plant growth, suggesting similar biocontrol 57 effectiveness among strains. Under the drier, more stressful, conditions plants exposed to A. 58 itadori had fewer leaves and accumulated less above-ground biomass. Overall, our results 59 suggest that genetic variability may not be the key to improve A. itadori biocontrol 60 effectiveness, but that predicted climate change, which anticipates drier and hotter summers in 61 the UK, could reduce the growth potential of F. japonica when exposed to A. itadori. 62

63

Keywords: Biological Control; Climate change; *Fallopia japonica*; Intraspecific Variation;
Invasive Species; Japanese Psyllid; Saturation Deficiency Index.

66 Abbreviations

67 LTLR: long-term laboratory-reared strain

- 68 STLR: short-term laboratory-reared
- 69 SDI: Saturation Deficiency Index
- 70

71 **1. Introduction**

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73 Invasive species are a significant problem in the United Kingdom, where they are estimated to 74 cost the economy approximately £1.7 billion per annum (Booy et al., 2008; Williams et al., 2010). Invasive species are both damaging to the UK's infrastructure and to the native 75 76 biodiversity. One of the most problematic invasive weeds in the UK is Japanese knotweed (Fallopia japonica [Houttuyn] Ronse Decraene), a species native to Japan. The lack of fertile 77 F. japonica males in Britain, as determined from Random Amplified Polymorphic DNA 78 (RAPDs) analysis, suggests that all F. japonica in the UK is derived from a single clonal 79 individual that has reproduced through vegetative propagation (Hollingsworth and Bailey, 80 2000). This low genetic diversity however, has not hindered its invasive ability. Fallopia 81 *japonica* has become established in a wide-range of habitats, and grows asexually from small 82 83 fragments of underground root networks – rhizomes, weighing less than a gram (Bashtanova et al., 2009; Hollingsworth and Bailey, 2000). These features, as well as its rapid growth rate, 84 85 make *F. japonica* highly invasive in the UK.

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87 There have been varying attempts to eradicate or control F. japonica. Manual or chemical removal can work at a local scale; however, the costs and time requirements make these 88 89 methods unfeasible as long-term or large-scale management solutions. Herbicide use in parks 90 and riparian areas where the plant is most prevalent has become less acceptable (Forman and 91 Kesseli, 2003). Biological control is often proposed as an effective alternative tactic for invasive species, such as F. japonica. Reuniting an introduced weed with its host-specific 92 natural enemies from their country of origin has resulted in successful suppression of many 93 invasive weeds worldwide (Clewley et al., 2012; Schwarzländer et al., 2018). In comparison 94 to other control methods, biocontrol can be used everywhere and is generally cost effective and 95 environmentally friendly (Wittenberg and Cock, 2001). 96

97

98 The use of biocontrol agents for *F. japonica* in the UK has been explored by the non-profit 99 organisation CABI, UK, since 2003. Initially, candidate species were identified from the 100 Kyushu Island of Japan, the region from where the UK invasive *F. japonica* clones are thought 101 to have originated (Djeddour and Shaw, 2010). Out of the 186 candidate arthropod species

considered, Aphalara itadori Shinji (Hemiptera: Aphalaridae), otherwise known as Japanese 102 knotweed psyllid, was found to be the best agent, since laboratory studies showed it to be host-103 specific (i.e. not affecting native plants) and highly damaging to F. japonica. However, despite 104 its effectiveness under laboratory conditions (Grevstad et al., 2013), the establishment of viable 105 populations in the field has been largely unsuccessful. A possible explanation for why field 106 releases have failed is a lack of genetic and phenotypic variability in the batches of A. *itadori* 107 that were released. Genetic bottlenecking is commonly implicated in the establishment failure 108 of biocontrol agents (see review by Fauvergue et al., 2012). It is not unusual in biocontrol 109 110 programs for host-range testing for specificity and safety to require a long period of laboratory rearing. Indeed, in the UK, A. itadori was maintained in the laboratory from 2004 until its 111 approval for release in 2010 (Shaw et al., 2009). Because the released A. itadori came from 112 populations maintained under Japanese summer conditions at 22°C 13:11 hours day:night 50-113 85% humidity for at least six years (~66 generations), they may have become conditioned to 114 115 the controlled environment room, as well as have potentially lost genetic diversity. This 'colony effect' of laboratory reared animals has been seen in other insect species, such as in Drosophila 116 117 when undergoing laboratory selection experiments (Harshman and Hoffmann, 2000) and when comparing wild to laboratory cultures of Drosophila (Sgrò and Partridge, 2000), and also in 118 Anopheles gambiae (Huho et al., 2007). As a result, the long-term laboratory-reared A. itadori 119 could have been ill-prepared for dealing with the variability in the natural environmental 120 121 conditions in the UK.

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Intraspecific variation — the diversity of characteristics amongst individuals of a species 123 (Cianciaruso et al., 2009) — can be an important factor aiding in the establishment of alien 124 species (Forsman, 2014), but as mentioned above variability may be reduced in laboratory-125 reared organisms. Plant and animal species with higher levels of intraspecific genetic and 126 phenotypic variation are more likely to establish successfully in new environments under 127 laboratory, semi-natural and natural conditions, with the largest effects seen in natural 128 experiments (Forsman, 2014). In addition, intraspecific variability can provide resilience to 129 changes in climatic conditions (Reusch et al., 2005; Sgrò and Hoffmann, 2004). Under climate 130 change, more variable populations are predicted to have an increased chance of containing 131 individuals with genotypes that allow population persistence (Oliver et al., 2015) whereas 132 locally adapted, less diverse populations are vulnerable because they have evolved traits to 133 suite only local stress factors (Benito Garzón et al., 2011). 134

The establishment of A. *itadori* may also have been affected by the interaction of different 136 climatic conditions. Hodkinson (2009) and pilot field experiments (CABI, unpublished data) 137 have shown that A. *itadori* population dynamics, and therefore their potential for establishment 138 in the UK, can be affected by expected rising temperatures and declining relative humidity. In 139 the UK, under climate change, conditions are likely to become more stressful due to a predicted 140 141 increase in temperature and decrease in humidity in the spring and summer (Murphy et al., 2010) when A. *itadori* are most active after hibernation (Hodkinson, 2009). Therefore, effective 142 biocontrol requires consideration of how different environmental conditions could affect 143 144 effectiveness and resilience to future changes in climate.

145

For this study we compared the performance of the strain used in historic biocontrol releases 146 to two other strains with different genetic backgrounds. The first genetically different strain we 147 tested was from the same locality as original strain (Kyushu in South Japan) but had a shorter 148 149 laboratory-rearing history (2 years compared to 13 years). Using a newly collected wild type strain would have been desirable but was not possible due the timing and cost of a new 150 151 collection and quarantine space. The second genetically different strain tested was a new hybrid strain created from two distinct provenances of A. *itadori*. To create the hybrid we combined 152 153 males from Kyushu and females from Hokkaido (North Japan; Grevstad et al., 2013). The Kyushu and Hokkaido strains of A. itadori are genetically distinct and both strain, as well as 154 155 the hybrid, can be distinguished using neutral molecular markers (Andersen et al., 2016). We tested a hybrid as a potential approach to increase genetic variability and vigor (Birchler et al., 156 2006; Szűcs et al., 2012). However, hybridization can also have negative effects which could 157 reduce the potential of this new hybrid strain (Heinze et al., 2019; Peer and Taborsky, 2005). 158 The performance and impact on F. japonica of the three strains was tested under two 159 environmental conditions that reflected standard laboratory growing conditions and a drier 160 environment reflective of climate change predictions. 161

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- 163 **2. Material and methods**
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- 165 *2.1. Aphalara itadori strains*
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We used three *Aphalara itadori* strains. Two, the LTLR and STLR strains, were established
using adults collected from Kyushu, Japan (taken in 2004 and 2015 respectively). The hybrid
strain was created by mating LTLR strain males with females from a *A. itadori* line collected

in 2007 in Hokkaido, Japan and reared since that date at the Agriculture and AgriFood Centre

- 171 (AAFC) in Lethbridge, Canada. The crossing of lines was completed in December 2016 at
- 172 AAFC-Lethbridge under 16L:8D laboratory conditions. Second generation adult hybrids
- 173 (N~200) were shipped to the UK and reared in CABI under standard laboratory conditions

174 (see below). We used fourth generation hybrids for oviposition during the experiment. All

three strains were reared on knotweed in 100 x 90 x 100cm Perspex cages (average \pm SD:

- 176 $16.9 \circ C \pm 3.8 \circ C$, 47.2% ± 10.7 % RH and 14L:10D) in CABI's Egham quarantine greenhouse
- 177 facility.
- 178

179 2.2. Experimental design and conditions

180

181 We tested two environmental conditions that we then characterized using empirical estimates of Saturation Deficiency Index (SDI), a measure of climate severity (Samways, 1987). In its 182 183 simplest form, SDI it is the difference between the saturation vapour pressure (SVP) at maximum temperature, and the actual vapour pressure of a volume of air at maximum 184 temperature (Green and Catling, 1971; Samways, 1987). The value of SDI increases with rising 185 temperature and/or decreasing relative humidity. For our experiment, treatments were created 186 by changing humidity within experimental cages. Plants under high SDI conditions, reflective 187 of climate change predictions (hotter and drier), had dry capillary matting for the base of the 188 cage and a 40 x 50cm gauze covered hole at the back of the cage to increase ventilation. Plants 189 in low SDI conditions had wet capillary matting for the base of the cage, watered with 800ml 190 tap water every week, reflecting the standard laboratory growing conditions. We calculated 191 empirical SDI values for each treatment cage following Abtew and Melesse, (2013) and 192 Samways (1987): 193

194 $SDI = SVP\left(\frac{100 - RH}{100}\right)$ (Equation 1)

where RH is relative humidity, and SVP is saturation vapour pressure calculated based ontemperature (T) as below:

197 SVP = 0.611 $e^{\left(\frac{17.27 \times T}{T+237.7}\right)}$ (Equation 2)

Humidity and temperature were recorded during the experiment at 30-minute intervals using
LogTag Haxo-8 dataloggers placed inside the sleeve of one randomly selected plant per cage.
We estimated SDI using the humidity and temperature recorded at each 30-minute interval. For
each day we then identified the three highest SDI values and calculated the arithmetic mean

per cage of those maxima over the duration of the experiment. This resulted in six SDI values
(one per cage). We averaged the three highest values instead of using the single highest value
to control for potential outliers. There are alternative methods of calculating SDI (see Green
and Catling, 1971), but we found results were equivalent with all methods (Table S1, Figure
S1).

207

Fifty-five days prior to the start of the first experimental batch, the rhizomes of 71 young *F*. *japonica* of uniform genetic stock (collected from a single *F. japonica* patch with vegetative reproduction) were cleaned and wet rhizome weights for each plant were obtained (average \pm SD: 75.85g \pm 36.06g). Each rhizome was potted in an individual plastic pot (14.7cm diameter) with a saucer (16.5cm diameter) and left to grow in a greenhouse under natural conditions (average \pm SD: 21.0°C \pm 4.5°C, 51.6% \pm 12.4% RH and 14L:10D).

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215 All experimentation was performed in quarantine glasshouses (average \pm SD: 21.0°C \pm 4.5°C, $51.6\% \pm 12.4\%$ RH and 14L:10D). Due to space constraints in the glasshouses, the experiment 216 217 was completed in three sequential batches over four months. For each batch, 14-15 days before the start of the experiment, 18 plants were cut to the fourth node above ground on the main 218 219 stem and first node from the stem on branches, with additional stems cut to ground level. This allowed us to standardise above-ground measurements of biomass. Cut F. japonica material 220 was collected and frozen, and dry weights later obtained for before and after above-ground 221 weight comparisons. Plants were then randomly assigned a A. *itadori* strain, and six plants from 222 each strain were placed into designated chambers for up to 8 days with 150 A. *itadori* adults to 223 allow oviposition (n ≈ 25 A. *itadori* per plant). 224

225

After the oviposition period, the total number of eggs per plant was counted by searching the 226 top and bottom of all leaves and nodes using a hand lens. Plants with very high numbers of 227 eggs were removed from egging chambers earlier to avoid high egg density variation across 228 229 treatments (batch one: one STLR low SDI and one hybrid high SDI plant; batch two: one STLR low SDI plant). Egg counts are minimum estimates because total counts would have required 230 231 damaging the plant, which would have prevented the experiment. We make the assumption here that the number of visible eggs is proportionally related to the total number of eggs. Plants 232 were then randomly assigned to a low or high SDI treatment, resulting in three plant replicates 233 per strain per treatment per batch (experiment total: n = 9 plant replicates per strain per 234 treatment, total n = 54). We used 1m long insect sleeves supported by bamboo hoops for each 235

plant to prevent A. itadori from moving between plants (Figure S3). Each plant was placed in 236 a 16.5cm diameter saucer and irrigated twice a week manually to ensure F. japonica survival 237 irrespective of treatment. Total adult counts began 37 days after plants were placed in treatment 238 cages. Emergent adults were counted and removed using a manual aspirator every 6-7 days for 239 six weeks to allow all adults from the eggs laid prior to the experiment to emerge. Although 240 the nymphal stages cause the most damage to plants (Djeddour and Shaw, 2010), accurately 241 counting nymphs without removal is complicated, therefore we used adult counts to infer 242 survival to adult emergence. After all adults were counted, we obtained wet weights of above 243 244 ground and below ground plant biomass. Above ground plant material was then frozen and dry 245 weights were later obtained.

246

247 2.3. Response variables: A. itadori performance and plant growth

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We used survival to adult emergence (henceforth referred to as 'A. itadori survival') and 249 development rates to assess A. itadori performance. Aphalara itadori survival was adjusted for 250 initial egg density, and was calculated as $100 * \frac{Adults}{Eggs}$, where Eggs was the total number of 251 eggs counted before moving the plants to the experimental treatments, and Adults was the total 252 number of emerged adults counted over the entire experiment for each plant. Aphalara itadori 253 development rate was evaluated by comparing the number of adults for each plant (expressed 254 as percentage of the total), counted at 1, 2 and 3 weeks after the first adult survival in each 255 cage. Counts after week 3 were not considered to avoid counting second generation offspring 256 emerging. One STLR plant from the low SDI treatment was removed as it had extreme adult 257 A. *itadori* numbers emerging compared to initial eggs counted. 258

259

Due to space limitations in the quarantine glasshouses, we could not assess how SDI treatments 260 affected plants without A. itadori. We evaluated impacts of A. itadori on F. japonica by 261 262 measuring differences in above and below ground biomass, number of leaves and stem height. There was considerable variation in these traits between plants, thus, in the variables rhizome 263 weight, maximum height and leaf number, we did not compare absolute growth but instead 264 calculated relative growth as $100 * \frac{(Final - Initial)}{Initial}$, where *Final* was the measurement taken at 265 266 the end of the experiment and Initial was the measurement before the start of the experiment. For the variable above-ground weight, the Initial was taken as zero (plants were potted as 267 rhizomes, without above ground material), and the Final was calculated as the sum of the 268

material that had been removed just prior to the experiment (to standardize plant size) and the remaining material at the end. Both were measured as dry weights. Plant material was wrapped in foil and placed into an oven at 70-90°C for 48h or until dried. As it was not possible to dry rhizomes before the experiment without killing the plant, change in below ground biomass was calculated using wet weights. The number of leaves was counted at the start and the end of the experiment. Stem height was measured using a ruler from soil level to the tallest standing point on the plant.

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277 2.4. Data analysis

278

We evaluated the effect of strain and SDI on A. itadori survival, development and the four 279 measurements of F. japonica growth using linear mixed effect regression models fitted with 280 function 'lmer' from package *lme4* (Bates et al., 2015) in R version 3.4.3 (R Core Team, 2017). 281 282 Table S2 lists the fixed and random effects considered for each model. In summary, all models included as a random factor the batch number (one, two or three) and, for A. itadori survival 283 284 and development, also observer ID (authors CF and CP, and Kate Constantine contributed to egg counting). All models included SDI and strain as fixed predictors. In addition, models 285 286 assessing plant growth included as covariates: total number of adults to control for variation in insect densities, and rhizome weight to control for initial plant conditions (except when 287 modelling rhizome weight). Models of A. itadori survival also included the total number of 288 eggs as a covariate. To model A. *itadori* development we used a B-splines analysis based on 289 count week to allow for non-linear changes in development. We tested models with additive 290 effects only, as well as with interactions between strain and SDI treatment. In the case of 291 development, Week was also tested for interactions (Table S2). Models with interactions were 292 only considered to be supported if interaction terms were significant (p-value < 0.05). We 293 294 evaluated model assumptions (normality and heteroscedasticity) plotting residuals from tested models. We used post-hoc tests based on R function 'difflsmeans' and 'lsmeansLT' from 295 296 package *lmerTest* (Kuznetsova et al., 2017) to contrast among strains.

- 297
- 298 **3. Results**
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- 300 *3.1. Aphalara itadori performance*
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Aphalara itadori survival varied among strains ($F_{2, 36, 17} = 12.49$, P < 0.001, n = 18, 17 and 18 302 for LTLR, STLR and Hybrid strains respectively; Table 1, Figure 2A). In particular, survival 303 from egg to adult emergence was significantly lower in hybrids (predicted mean [95% 304 confidence intervals]: 26.00% [10.99 - 41.01]) compared to LTLR (57.72% [41.86 - 73.57]) 305 and STLR (54.79% [34.49 – 75.08]) strains, but LTLR and STLR did not differ (P = 0.68). 306 SDI did not significantly affect A. *itadori* survival ($F_{1, 46.86} = 1.66, P = 0.20$), but survival was 307 proportionally higher in plants with fewer eggs suggesting a density dependence effect ($F_{1, 46.82}$ 308 = 7.98, P = 0.007). 309

310

The proportion of adults emerging generally decreased from the first to the third week, with earlier emergence time under low SDI (higher humidity, $F_{I, 153} = 28.34$, P < 0.001; Table 1). The LTLR strain had the fastest development rates, with notable difference under high SDI, with the LTLR strain having peak emergence in the first week one compared to both the STLR and the hybrid strain which displayed peak emergence during the second week (Table 1, Figure 1B). There was an interaction between STLR and SDI, with the majority of STLR adults emerging sooner under lower SDI ($F_{2, 153} = 6.69$, P < 0.001).

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319 *3.2. Impacts on F. japonica*

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The considerable variation in plant growth recorded in all four traits was not consistently 321 associated with the A. itadori strains to which plants were exposed (Figure 2, Table 2). The 322 only significant effect of strain was detected in the change in plant height, where hybrids 323 (predicted mean [95% confidence intervals]: 352.21% [232.24 – 472.19]) had least effect in 324 suppressing plant growth (plants had greater percentage changes in height) compared to LTLR 325 (275.23% [155.23 - 395.22]) and STLR (286.40% [161.65 - 411.14]) which had similar 326 327 estimates ('lsmeansLT' estimates: P = 0.08 and 0.07, for hybrids vs LTLR and STLR respectively). For plant height, we also found evidence of a differential effect of A. itadori 328 strain conditional to SDI: the STLR strain was most effective at reducing maximum height 329 under high SDI values, but least effective under low SDI ($F_{2,44,16} = 4.08$, P = 0.019, N = 54, 18 330 plants per strain; Figure 2C, Table 2). SDI influenced leaf number and above ground weight, 331 with plants having fewer leaves ($F_{1,46.83} = 5.82$, P = 0.020) and smaller above ground weight 332 $(F_{1.38.06} = 5.87, P = 0.020)$ under higher SDI (low humidity). 333

Out of the four plant growth variables tested, leaf number was the only response variable which was influenced by another predictor besides strain and SDI (other predictors: total eggs, number of adults, week of emergence and initial rhizome weight; see Table A2 for when these predictors were included in our models), where higher rhizome weights at the start of the experiment were associated with more leaves ($F_{1, 47.01} = 9.29$, P = 0.004; Table 2). None of the variables we tested explained change in rhizome weight (Table 2; Figure 1A).

341

342 4. Discussion

343

Our study aimed to improve biocontrol of F. japonica by exploring the effectiveness of 344 different A. itadori strains. We hypothesised that strains which had spent less time in the 345 laboratory (STLR and hybrid strain) would have undergone less selection pressure to perform 346 better under standard laboratory conditions, and therefore would perform better under altered 347 348 climatic conditions. Previous studies have shown that laboratory rearing may lead to reduced genetic variability compared to wild stocks due to population bottlenecks and selection (Huho 349 350 et al., 2007; Sgrò and Partridge, 2000), and therefore laboratory stocks tend to become more stress sensitive as selection for stress-related traits is relaxed (Hoffmann and Ross, 2018). Our 351 352 results did not consistently support our predictions suggesting longer time in laboratory culture by itself is not affecting the performance of A. *itadori* biocontrol for F. *japonica*. 353

354

Among the strains, hybrids had lower survival and developed slower compared to the LTLR 355 strain. Although the hybrid was created from two genetically different strains (Andersen et al., 356 2016), differences in the single-nucleotide polymorphisms (SNPs) may not have matched 357 differences in functional gene regions linked to the traits we were assessing. In addition, 358 although there have been studies which have shown improved hybrid fitness, for example in 359 360 ornamental pear tree Pyrus calleryana (Culley and Hardiman, 2009), hybridisation in our study could have led to reduced, rather than improved, fitness. Between-population crosses from 361 Bremgartewald and Spilwald strains of the black timber bark beetle, Xylosandrus germanus, 362 were found to be less fit compared to inbred individuals (Peer and Taborsky, 2005). Hybrids 363 from populations of the intertidal copepod species Tigriopus californicus also exhibited the 364 negative effect of outbreeding depression, with hybrid fitness initially lower in terms of 365 survivorship and morphology (Hwang et al., 2011). In our study, the hybrid strain was created 366 from the combination of males from the Kyushu strain, which performs best on F. japonica 367 compared to other knotweeds, and females from the Hokkaido strain, which oviposit and 368

develop well on *R. sachalinensis* (Grevstad et al., 2013). It is possible that hybrid breakdown may have occurred whereby the Hokkaido strain's adaptation and preference to living on *R. sachalinensis* was expressed in the hybrids, explaining the low survival to adult emergence observed in the hybrid strain compared to other strains. However, it is important to note that the hybrid was equal to the other two strains observed in terms of reducing the plant growth predictors assessed, and future work assessing more traits would further aid in determining the performance of hybrid strains.

376

377 Our study found that Aphalara itadori development was slower under high SDI, which has also been found for other psyllid species (see Hodkinson, 2009). Slower development could explain 378 why plants exposed to A. itadori under stressful low humidity levels (high SDI) had lower 379 growth in above-ground weight, height and number of leaves, compared to plants under high 380 humidity levels. The more damaging nymphal stage of A. itadori is extended under slower 381 382 development (Djeddour and Shaw, 2010) and therefore the per capita impact of individuals is likely to increase, potentially making them more effective biocontrol agents under high SDI 383 384 conditions. Indeed, we found that the STLR strain developed slowest and had a greater impact on plant height under high SDI. However, this benefit could be offset by there being fewer 385 386 generations per season, something that will need to be confirmed in future studies.

387

The findings that *A. itadori* survival was not influenced by SDI contrasts with other studies on other *A. itadori* species that have shown that high SDI leads to lower survival (Hall and Hentz, 2001; Hodkinson, 2009; McFarland and Hoy, 2001). These differences may reflect variation among species, but it is also possible that our drier conditions were not sufficiently stressful to induce mortality. The experiments were done within a greenhouse where conditions limited our ability to strictly control temperature and humidity.

394

Due to space limitations in the quarantine area we could not assess how environmental conditions affected plants without *A. itadori*. However, the reduced above-ground biomass and number of leaves observed in plants under high SDI could reflect more stressful conditions for the plants, especially as all plants were regularly watered, so only ambient humidity changed. If plants by themselves were not affected by the more stressful ambient conditions in the experiment, this suggests that *A. itadori* could be even more damaging when plant do suffer from high stress conditions in the field.

Notably, we found no effects of strain or SDI on rhizome weight. This could be because both
insects and ambient humidity do not directly affect rhizomes, and nutrient availability in the
soil was sufficient to avoid rhizome depletion associated to above ground growth. Since *F*. *japonica* is mainly spread by pieces of rhizome this highlights the challenge in developing an
effective biological control to reduce the spread of this invasive plant.

408

409 Overall, our results do not support a beneficial role of intraspecific variation in the biocontrol effectiveness of A. itadori. Genetic work would be necessary to reveal if this is due to genetic 410 411 variability being different from our expectation (lower in LTLR and highest in hybrids). Additional work under laboratory and field conditions would also be necessary to test a wider 412 range of climate conditions (as responses are likely to be non-linear), to evaluate cross-413 generational changes including hybrid fitness after more generations, and to take into account 414 additional factors such as predator avoidance and overwintering performance. Effectively 415 controlling F. japonica, both above and below ground, is still the challenge ahead. 416





419 **Figure 1** Relationship between *Aphalara itadori* performance in terms of (A) percentage A.

- 420 *itadori* survival to adult emergence versus Saturation Deficiency Index (SDI) in treatment
- 421 cages and (B) A. *itadori* development rate per week. Data points show the observed survival
- 422 of three *A. itadorii* strains (LTLR = Long-term laboratory reared; STLR = short-term
- 423 laboratory reared and Hybrid strain) grown on *Fallopia japonica*. Lines show the predicted
- relationship with SDI from a linear mixed effects model with shaded areas indicating 95%
- 425 confidence intervals.
- 426



strain 📥 LTLR 📥 STLR 📥 Hybrid

429 **Figure 2** Relationship between growth of *F. japonica* versus Saturation Deficiency Index

- 430 (SDI) in treatment cages. Data points show the observed survival to adult emergence of three
- 431 *Aphalara itadori* strains (LTLR = Long-term laboratory reared; STLR = short-term
- 432 laboratory reared and Hybrid strain) grown on *Fallopia japonica*. Plant growth was
- 433 measured as either (A) rhizome weight, (B) above ground weight, (C) maximum height, and
- 434 (D) leaf number. Lines show the predicted relationship with SDI from a linear mixed effects
- 435 model with shaded areas showing 95% confidence intervals.
- 436

437	Table 1 Coefficient estimates for the model predicting Aphalara itadori adult survival to
438	adult emergence as a function of total number of A. <i>itadori</i> eggs, Saturation Deficiency Index
439	value (SDI), and A. <i>itadori</i> strain (LTLR = Long-term laboratory reared; STLR = short-term
440	laboratory reared and Hybrid strain); and adult development as a function of time (in weeks),
441	Saturation Deficiency Index value (SDI), and A. itadori strain. We report best parameter
442	estimates (β), their 95% confidence interval (CI), <i>P</i> -value, and the number of plants used in
443	each analyses (N). The strain reference level (e.g. 'LTLR') is indicated in parentheses. The
444	colon separating variable names indicates interaction terms. Significant variables are

445 highlighted in bold.

Variable	β	Lower 95% CI	Upper 95% CI	<i>P</i> -value	
Survival $(N = 53)^*$					
Intercept (LTLR)	66.05	50.23	81.88	< 0.001	
SDI	0.54	-0.28	1.37	0.204	
STLR	-2.93	-16.71	10.86	0.679	
Hybrid	-31.71	-45.67	-17.76	<0.001	
Total eggs	-0.02	-0.03	-0.01	0.007	
Development ($N = 54$)					
Intercept (LTLR: Low)	78.05	69.42	86.69	<.001	
STLR	-20.91	-30.29	-11.54	<.001	
Hybrid	-10.23	-19.67	-0.78	0.035	
High	-32.64	-40.26	-25.02	<.001	
LTLR: Week 1-2	-56.54	-67.23	-45.85	<.001	
LTLR: Week 2-3	-77.62	-88.31	-66.93	<.001	
STLR: Week 1-2	33.07	19.98	46.16	<.001	
STLR: Week 2-3	29.67	16.58	42.76	<.001	
Hybrid: Week 1-2	14.72	1.63	27.81	0.029	
Hybrid: Week 2-3	15.96	2.86	29.05	0.018	
High: Week 1-2	53.17	42.48	63.86	<.001	
High: Week 2-3	44.75	34.06	55.43	<.001	
Total Eggs	0	-0.01	0.01	1	

446 *One STLR plant had extreme adult *A. itadori* numbers emerging was removed from analysis

448	Table 2 Summary of models predicting percentage change in <i>Fallopia japonica</i> factors as a
449	function of Saturation Deficiency Index (SDI), Aphalara itadori strain (LTLR = Long-term
450	laboratory reared; STLR = short-term laboratory reared and Hybrid strain), the total number
451	of A. itadori adults produced at the end of the experiment, and the initial rhizome weight. We
452	report best parameter estimates (β), their 95% confidence interval (CI), <i>P</i> -value, and the
453	number of plants used in each analyses (N). The strain reference level (e.g. 'LTLR') is
454	indicated in parentheses. The colon separating variable names indicates interaction terms.
455	Significant variables are highlighted in bold.

Variable	ß	Lower 95%	Upper 95%	P-
Variabic	Р	CI	CI	value
Rhizome weight ($N = 54$)				
Intercept (LTLR)	27.32	-3.89	58.53	0.109
SDI	0.22	-1.03	1.47	0.729
STLR	7.29	-13.86	28.44	0.503
Hybrid	12.77	-12.45	37.98	0.326
Number of Adults	-0.01	-0.06	0.04	0.770
Above Ground Weight (<i>N</i> =				
46)*				
Intercept (LTLR)	57.97	27.22	88.73	0.020
SDI	-0.87	-1.58	-0.17	0.020
STLR	-2.01	-13.79	9.77	0.740
Hybrid	-4.00	-18.61	10.61	0.594
Number of Adults	-0.02	-0.05	0.00	0.106
Initial Rhizome Weight	-0.00	-0.13	0.12	0.940
Maximum Height ($N = 54$)				
Intercept (LTLR)	340.49	204.08	476.90	< 0.001
SDI	-1.14	-8.30	6.01	0.755
STLR	150.85	42.17	259.54	0.009
Hybrid	142.80	23.88	261.72	0.023
SDI: STLR	-16.27	-26.10	-6.45	0.002
SDI: Hybrid	-7.67	-17.33	2.00	0.127
Number of Adults	-0.15	-0.32	0.02	0.081
Initial Rhizome Weight	-0.12	-0.94	0.69	0.769

Leaf Number ($N = 54$)			
1622.59	493.61	2751.57	0.009
-51.75	-93.79	-9.71	0.020
-127.08	-833.21	579.04	0.726
-438.50	-1272.10	395.10	0.308
-0.26	-1.88	1.36	0.754
12.48	4.45	20.50	0.004
	1622.59 -51.75 -127.08 -438.50 -0.26 12.48	1622.59 493.61 -51.75 -93.79 -127.08 -833.21 -438.50 -1272.10 -0.26 -1.88 12.48 4.45	1622.59 493.61 2751.57 -51.75 -93.79 -9.71 -127.08 -833.21 579.04 -438.50 -1272.10 395.10 -0.26 -1.88 1.36 12.48 4.45 20.50

456 *Eight *F. japonica* (three LTLR, one STLR and four Hybrid) had weights missing and were

457 removed from analysis

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583 Appendix A

- **Table A1** Alternative methods of calculating Saturation Deficiency Index value (SDI)
- adapted from (Green and Catling, 1971). Maximum temperatures were the maximum
- temperatures across the whole experiment.

SDI Methods

- 1 Mean of 3 highest daily maximum temperatures with the mean of the three vapour pressures coinciding with the 3 highest maximum temperatures
- 2 Mean of the 3 SDI values (millibars) coinciding with the 3 highest maximum temperatures

587

- 589 **Table A2** All models tested to analyse the effects of strain and Saturation Deficiency Index
- value (SDI) on *Aphalara itadori* performance (survival to adult emergence and development)
- and *A. itadori* impact on *Fallopia japonica* growth (rhizome weight, above ground weight,
- 592 maximum height and leaf number. * indicates tested interactions and models used are
- 593 highlighted in bold.

Model	Fixed predictors	Random Factors		
A.itadori Performance				
Survival				
<i>S1</i>	Strain + SDI + total eggs	Batch + observer		
<i>S2</i>	Strain*SDI + total eggs	Batch + observer		
Development				
D1	Strain + SDI + week emerge + total eggs	Batch + observer		
D2	Strain*SDI + week emerge + total eggs	Batch + observer		
D3	Week emerge*(strain + SDI) + total eggs	Batch + observer		
D4	strain*(SDI + week emerge) + total eggs	Batch + observer		
Impact on F. japa	onica			
Rhizome Weight				
R1	Strain + SDI + total number of adults	Batch		
R2	Strain*SDI + total number of adults	Batch		
Above Ground W	⁷ eight			
A1	Strain + SDI + total number of adults + before	Batch		
	rhizome weight			
A2	Strain*SDI + total number of adults + before rhizome	Batch		
	weight			
Maximum Heigh	t			
H1	Strain + SDI + total number of adults + before rhizome	Batch		
	weight			
H2	Strain*SDI + total number of adults + before	Batch		
	rhizome weight			
Leaf Number				
L1	Strain + SDI + total number of adults + before	Batch		
	rhizome weight			

594

L2



595 Figure A1 Four methods chosen for calculating Saturation Deficiency Index value (SDI) 596 adapted from (Green and Catling, 1971). Points are the calculated SDI values of dataloggers 597 for each experimental batch. Each datalogger was placed in one sleeve within a treatment 598 599 cage.

Batch • 1 • 2 • 3

(1A) SDI was firstly calculated per day by taking the mean of the top three temperature 600 values and its corresponding relative humidity values (RH). The final SDI value assigned to 601 the batch was the average SDI for the whole experiment. (1B) SDI was firstly calculated per 602 603 day by taking the mean of the top three temperature values and corresponding RH values. The final SDI value was than assigned by calculating the mean of the top three SDI values for 604 605 the whole experiment. (2A) SDI values were calculated for each reading (30min) and the mean of the highest three SDI values was obtained. The final SDI value assigned to the batch 606 607 was the average SDI for the whole experiment. (2B) SDI values were calculated for each reading (30min) and the mean of the highest three SDI values was obtained. The final SDI 608 609 value assigned by calculating the mean of the top three SDI values for the whole experiment. 610



Figure A2 Experimental *Fallopia japonica* plants. a) For the experiment, plants were placed
in a 16.5cm diameter saucer within a humidity cage with capillary matting. They were
irrigated twice a week manually; b) after egg counts, plants were covered in 1m long insect
sleeves, tied with elastic bands and supported by bamboo halos to avoid *Aphalara itadori*escaping.