Variation in levels of acceptance, developmental success, and abortion of *Ha-lyomorpha halys* eggs by native North American parasitoids

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### 28 Abstract

29 In order for native parasitoids to adapt to an unsuitable invasive host, there must be intraspecific variation in traits that could be selected upon to enable exploitation or avoidance. Using a-native 80 North American parasitoid species (Hymenoptera: Scelionidae) that often unsuccessfully attacking Β1 the eggs of the invasive brown marmorated stink bug Halyomorpha halys (Hemiptera: B2 33 Pentatomidae) as a case study, we assessed the variation in traits that may determine the parasitoids' future ability to adapt to the invasive host by either exploiting or avoiding *H. halys* eggs 34 (acceptance, developmental success). The aim of surveying these We also assessed variation in the 85 parasitoids' ability to induce *H. halys* host egg abortion, which may contribute to biological control 86 of *H. halvs* in invaded areas. parasitoid's variations is to assess their, and their current contribution 87 to theits biological control (ability to induce host egg abortion) of Halvomorpha halys. The first set 38 of experiments evaluated intra- and interspecific variation using standardized laboratory tests with 39 iso-female lines of Telenomus podisi and Trissolcus euschisti that included matching of detailed 40 behavioural observations of acceptance with developmental outcomes. In a second set of 41 experiments, we assessed how variation in developmental ability and abortion induction may affect 42 levels of biological control by indigenous parasitoid species. We examined a broader sample of 43 parasitoids that emerged from field collections of egg masses of an indigenous north American stink 44 bug Podisus maculiventris in a region newly invaded by H. halys. Results from the first set of 45 experiments showed high levels of acceptance of *H. halys* eggs among iso-female lines of 46 parasitoids, but offspring development success was almost zero. H. halys egg abortion due to 47 unsuccessful parasitism was often very low and varied among iso-female lines only for T. podisi. In 48 the second set of experiments we never observed increases in abortion levels of Halyomorpha halys 49 eggs above natural levels, even for the two species (T. euschisti and T. podisi) that were observed to 50 oviposit in and abort *H. halys* eggs in the first set of experiments. We conclude that while there may 51 be some variation in behavioural and physiological parameters mediating acceptance and abortion 52 of *H. halys* eggs by native North American egg parasitoids, there does not appear to be significant 53 54 variation in developmental success. Moreover, and that their current biological control impact of H. halys eggs via host egg abortion is likely very low. 55

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Keywords: Intraspecific variation, biological control, acceptance behaviour, evolutionary trap,
Scelionidae, *Telenomus podisi*, *Trissolcus euschisti*

## 61 1.1 Introduction

The introduction of invasive species exposes native species to novel biotic selective pressures. The 62 establishment, spread, and proliferation of invasive species in new environments can have important 63 ecological consequences such as niche displacement, competition and in some cases, extinction of 64 native species (Mooney and Cleland, 2001; Sax and Gaines 2008; Ekesi et al., 2009; Wilson and 65 Holway, 2010; San Sebastián et al., 2015; Kenis et al., 2017). However, new and frequent 66 interactions with invasive species may lead to novel behavioural responses and trophic interactions. 67 For example, native natural enemies may begin to exploit the invasive species as a prey resource 68 (Caroll and Dingle, 1996, Mooney and Cleland, 2001). In some cases, invasive species may initially 69 be unsuitable resources for native natural enemies that commonly attempt to exploit them, resulting 70 71 in evolutionary traps (Schlaepfer et al., 2005). Nevertheless, the adaptation of native species to successfully exploit invasive species can occur over a range of timescales. Native species can adapt 72 73 either by avoiding the unsuitable invasive species or by evolving traits that increase the suitability 74 of the invasive species as prey (reviewed in Berthon, 2015).

75 For insect egg parasitoids to adapt to an initially unsuitable invasive host (i.e., an escape from the evolutionary trap), there needs to be genetic variation in traits that mediate the parasitoid's 76 77 levels of host acceptance (oviposition inside the eggs of the invasive host) or the ability of the parasitoid's offspring to develop once accepted. Selection on variation in behaviours that mediate 78 acceptance could lead to avoidance of the unsuitable host, while selection on variation in 79 developmental success could lead to increased host suitability (Keeler and Chew, 2008; Nelson et 80 al., 2011; Abram et al., 2014). In addition, when an invasive species that acts as an evolutionary 81 82 trap for native parasitoids is the target of biological control programs, it may be important to consider, variation in the ability of parasitoids to kill the invasive host when they unsuccessfully 83 attack it ("non-reproductive effects"; reviewed in Abram et al., 2019a) may be important to 84 consider. The mortality of the invasive pest resulting from non-reproductive effects could 85 theoretically contribute to biological control of the invasive pest via both direct and indirect 86 87 ecological effects (Kaser et al., 2018; Abram et al., 2019a; Stahl et al., 2019a). Assessing 88 intraspecific variation in native parasitoid acceptance, developmental success, and the ability to induce host mortality can provide insight into the potential for the adaptation of native species to 89 90 new invasive hosts, while also forming and form the basis for genetic improvement through breeding programs for native biological control agents (Lommen et al., 2017; Kruitwagen et al., 91 2018). 92

93 The brown marmorated stink bug, *Halyomorpha halys* (Stål) (Hemiptera: Pentatomidae), is
94 a polyphagous insect pest native to Asia that has rapidly spread throughout North America and

Europe (Haye et al., 2015a; Leskey and Nielsen, 2018, Maistrello et al., 2018), causing economic 95 losses in a number of different crops (Leskey et al., 2012; Maistrello et al., 2017; Bosco et al., 96 2018). Many studies have been conducted in invaded areas to survey for native parasitoid and 97 98 predator species with the potential potentially able to contribute to the biological control of H. halvs biological control (e.g. Ogburn et al., 2016; Dieckhoff et al., 2017; Haye et al., 2015b; Herlihy et 99 al., 2016; Roversi et al., 2016; Cornelius et al., 2016a, b; Abram et al., 2017a; Costi et al., 20198). 100 Overall, egg parasitoids in the family Scelionidae (mostly species of Telenomus Haliday and 101 Trissolcus Ashmead) are the most common group of natural enemies attacking H. halys in invaded 102 103 areas. Several studies have found that these parasitoids readily accept H. halys eggs for oviposition, but their offspring rarely develop successfully (Abram et al., 2014; Haye et al., 2015b; Konopka et 104 105 al., 2018, 2019, 2020). This combination of high levels of acceptance and low developmental suitability suggests that *H. halvs* represents an evolutionary trap for these native scelionid 106 107 parasitoids (Abram et al., 2014; Schlaepfer et al., 2005). However, a few studies have shown that between 15-25% of attacked H. halys eggs abort development as a result of unsuccessful parasitoid 108 109 attack by parasitoids in invaded areas, potentially as a result of a lack of shared evolutionary history (Abram et al., 2014, 2016; Haye et al., 2015b). Recent studies using molecular diagnostic markers 110 111 have confirmed that the unsuccessful attack of *H. halys* eggs by native scelionid egg parasitoids is common in nature and have shown that observed H. halys egg mortality that could be due to 112 parasitoid-induced host egg abortion (Konopka et al., 2018; Gariepy et al., 2019). Thus, aborted 113 host egg development induced by native parasitoids may contribute to biological control of H. halys 114 in invaded areas, although the magnitude of the effect is unclear. 115

Some variation in the ability of indigenous parasitoids to develop in and abort *H. halys* eggs 116 has previously been observed in a number of field and laboratory studies (Abram et al., 2014; 117 Cornelius et al., 2016a, b; Herlihy et al., 2016; Ogburn et al., 2016; Dieckhoff et al., 2017; Konopka 118 et al., 2019; Tognon et al., 2017, 2019; Abram et al., 2014; Haye et al., 2015b; Abram et al., 2016; 119 Abram et al., 2017b). However, laboratory studies have either been restricted in taxonomic breadth, 120 and within-species genetic variation (e.g., Abram et al., 2014, 2016; Konopka et al., 2018, 2020; 121 122 Haye et al., 2015b), or have lacked direct behavioural observations of parasitoids and unexposed control egg masses to relate levels of acceptance to developmental success and egg abortion (e.g., 123 Tognon et al., 2017, 2019). In field studies, because most ovipositions by native parasitoids are 124 125 unsuccessful (Abram et al., 2014, 2016, 2017b), the true incidence of *H. halys* egg attack by genetically variable pools of native parasitoids and how it might explain variation in *H. halys* egg 126 mortality is still mostly unclear (Haye et al., 2015b). Previous studies have suggested that native 127 128 parasitoids might eventually adapt to be able to develop in *H. halys* eggs (Abram et al., 2014, 2016;

Tognon et al., 2017) but none have assessed whether there is enough intraspecific variation in relevant traits <u>that would be needed</u> for natural selection to proceed. Standardized laboratory studies that consider the role of intra- and inter-specific variation in acceptance, offspring development, and host egg abortion induction in field populations of parasitoids could improve understanding of the range of impacts that indigenous parasitoids may have on *H. halys* in different invaded regions, both in the short and long terms.

The goal of this study is to understand intra and interspecific variation in egg parasitoid 135 traits that may determine their ability to adapt to exploiting or avoiding *H. halys* eggs (acceptance, 136 developmental success), and their current contribution to its biological control (ability to induce 137 host egg abortion). We took two approaches. First, we conducted standardized tests with narrow 188 189 inter-lines genetic samples comparing among (iso-female lines) of two common native North 140 American parasitoid species, -attacking *H. halvs* that included matching of detailed behavioural observations of acceptance with developmental outcomes. Second, we screened a broader 141 taxonomic and genetic sample of parasitoids that emerged from field collections of egg masses of 142 143 an indigenous stink bug to assess how variation in developmental ability and abortion induction may affect levels of biological control by indigenous parasitoid species in a region newly invaded 144 by *H. halys*. 145

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## 147 **1.2 Material & Methods**

## 148 *1.2.1 Insect colonies*

The spined soldier bug, Podisus maculiventris (Say) (Hemiptera: Pentatomidae), a predatory 149 species native to North America, was used to produce egg masses for rearing native parasitoids and 150 151 "sentinel" egg masses to collect parasitoids in the field. The P. maculiventris colony, established from individuals collected in Ontario and Quebec, Canada in 2015, was reared continuously in 152 mesh cages with access to potted fava bean plants (Vicia faba L.) and mealworms (Tenebrio molitor 153 L.). Egg masses laid on the sides of cages and on plants were collected daily and glued to strips of 154 filter paper with white non-toxic glue (Elmer's school glue) and offered to parasitoid females to 155 156 maintain parasitoid colonies. Egg masses laid on strips of Reemay fabric (Avintiv, USA), which were provided as additional oviposition substrate, were stored for up to 4 days at 8°C before 157 deployment as sentinel egg masses. This short period of cold storage does not affect parasitism or 158 parasitoid offspring emergence (PKA, unpublished observations). 159

*Trissolcus euschisti* (Ashmead) (Hymenoptera: Scelionidae) (native to North America) and
 *Telenomus podisi* Ashmead (Hymenoptera: Scelionidae) (native to Southeastern USA) are solitary,
 sexually reproducing idiobiont and generalist parasitoids of stink bugs species of economic

importance and egg parasitoids distributed throughout North America that both have broad host 163 ranges within the Pentatomidae. Tthey are also the most common scelionids attacking eggs of the 164 brown marmorated stink bug in North America (Abram et al., 2017b). -We established five strains 165 of Tr. euschisti (provided by Tara Gariepy, London Research and Development Centre, Agriculture 166 and Agri-Food Canada; Jacques Brodeur and Mathilde Gaudreau, Université de Montréal) and five 167 strains of T. podisi (provided by Jacques Brodeur and Mathilde Gaudreau, Université de Montréal; 168 James Walgenbach and Emily Ogburn, North Carolina State University). Strains tested in our study 169 170 likely represent a small sample of the range present in North America, as our collections are based on a small number of localities. However, our goal was not to be comprehensive. Rather, we aimed 171 to conduct a preliminary assessment of whether there could be inter-line variation in the parameters 172 173 we measured. Single females from each collection were used to establish an iso-female lines for each strain, which were reared in climate-controlled rooms for at least 5 generations of sib-mating 174 before testing. Using this iso-female line approach, any phenotypic differences among lines should 175 be due mostly to genetic differences (Beukeboom and Zwaan, 2007), although we cannot exclude 176 177 the possibility that other differences such as microbiomes (e.g. facultative or obligate microbial 178 endosymbionts), long-term rearing effects, or maternal effects -could also be a factor, especially 179 because detailed genetic analyses of these lines has not been performed. Table 1 shows the parasitoid lines tested and their origins. Fresh (<24h since laying) P. maculiventris eggs were 180 offered to the parasitoids on a regular basis for colony maintenance. To collect parasitoid females 181 for use in experiments, parasitized egg masses were kept in 1.5 mL plastic microtubes and observed 182 daily for newly emerged female parasitoids. Females were always paired with males from the same 183 strain. 184

Halyomorpha halys individuals were originally collected from two locations in Chilliwack,
 British Columbia, in 2016 and were reared thereafter at the Agassiz Research and Development
 Center (BC). Stink bugs were maintained in mesh cages provided with potted pepper (*Capsicum annum* L.) plants and store-bought corn and carrots as food sources. Eggs laid by females on
 Reemay fabric, plant leaves, paper towels, and the sides of the cages were collected daily for
 laboratory experiments.

- 191 All insect colonies were maintained at  $23 \pm 2^{\circ}$ C, 40-60% relative humidity, and a 192 photoperiod of 16L:8D.
- 193

194 *1.2.2 Intraspecific variation in acceptance and development* 

195 The goal of the first experiment was to compare the level, latency, and rate of behavioural

acceptance of *H. halys* eggs among strains of *T. euschisti* and *T. podisi*. Prior to use in experiments,

newly emerged parasitoids from each species and strain were isolated daily in 1.2 mL Eppendorf 197 tubes with a drop of honey water (70% honey and 30% water) for food, kept with a male for 24 198 hours to allow mating, and then kept alone for one week to allow time for ovariole maturation. For 199 each trial, individual female parasitoids were transferred to small petri dishes (diameter: 5.0 cm, 200 height: 1.0 cm) where a fresh *H. halvs* egg mass containing 28 eggs was secured to the bottom of 201 the dish with a small amount of white, non-toxic glue. The trials were recorded for 5 h using a mini-202 button CCD camera connected to a digital microscope (Dino-Lite, London, Ontario, Canada). Each 203 treatment conducted in a climatic chamber (25°C and 16L:8D) was replicated between 17 and 22 204 205 times for each strain of each parasitoid species. The experiment was performed in 42 testing days 206 over a total period of 129 days. For each set of egg masses exposed to parasitoids (i.e. on each day of the experiment), unexposed egg masses (controls) were subjected to the same conditions as the 207 treatment egg masses except that they were not exposed to a parasitoid. These unexposed egg 208 209 masses (n=85) allowed the measurement of baseline levels of *H. halvs* egg abortion that was not due to parasitoids (see Abram et al., 2016). 210

After the recording period, parasitoids were removed, and both exposed and unexposed egg masses were maintained under standard rearing conditions for at least 30 days, observing them daily to record the number of host nymphs and parasitoids emerging. After 30 days, any unhatched eggs were dissected and classified as: (i) emerged or fully developed but unemerged *H. halys* nymphs or (ii) aborted, containing only undifferentiated liquid contents. The dissections yielded no fully developed parasitoid adults that failed to emerge.

When analyzing each video (n=198), a single observer recorded the occurrence and timing 217 of ovipositions, indicated by the parasitoid inserting its ovipositor into a host egg followed by 218 marking behaviour (see Abram et al., 2014). This allowed us to determine the level of acceptance of 219 the host eggs (the total number of eggs accepted), the latency to the first successful oviposition, and 220 the total time period over which parasitism took place (time between the first and last ovipositions). 221 Self-superparasitism was relatively uncommon in our trials (5.2% of all ovipositions for T.222 euschisti; 6.8% of all ovipositions for T. podisi), and the total number of ovipositions was almost 223 perfectly correlated with the number of eggs accepted for both T. euschisti (r = 0.99) and T. podisi 224 (r = 0.99). Thus, superparasitism events were disregarded in the analysis, and the number of eggs in 225 which at least one oviposition was observed was used as the metric of behavioural acceptance. 226

We expected the level of acceptance to indicate the readiness of native parasitoids to oviposit in *H. halys* eggs. We anticipated that the time until the first oviposition occurred could be a proxy for the time necessary for native parasitoids to recognise the eggs of the host and accept it as a resource. Finally, we expected acceptance rate, calculated as the total time to complete parasitism

of the *H. halys* egg mass divided by the number of eggs attacked, to measure the efficiency with which the different strains of the two species of parasitoids exploited *H. halys* eggs. Moreover, the acceptance rate would be driven by other behaviours that decrease this efficiency, such as increased time required to drill into eggs, host rejections, superparasitism, and short patch leaving bouts (Field 1998).

Next, we determined the level of *H. halys* egg abortion associated with unsuccessful ovipositions by comparing the number of host eggs the parasitoids oviposited into <u>and</u> the developmental outcomes of eggs. The number of eggs aborted due to oviposition in each replicate was calculated with the following formula (see also Abram et al. 2016):

240

241 eggs aborted due to oviposition =  $AbE - \left(\frac{AbC * TotE}{TotC}\right)$ 

242

Where AbE and AbC are, respectively, the number of aborted eggs observed in the exposed and
control egg masses, and TotE and TotC are the total number of eggs in the exposed and control,
respectively. <u>Abortion levels in each exposed egg mass (AbE and AbC) were adjusted using pooled</u>
values of TotE and TotC from paired egg masses set up on the same date.

(eqn. 1)

Two factors contribute to the numbers of aborted *H. halys* eggs among parasitoid strains. 247 The first is variation in the number of eggs attacked; the second is variation in the per-attack 248 probability of aborting host development. To investigate whether parasitoid strains differed with 249 respect to the per-attack probability of inducing H. halys egg abortion, we applied the following 250 formula to calculate abortion efficiency, calculating the number of eggs aborting due to oviposition 251 (for eggs accepted by parasitoids only), and then dividing it by the total number of accepted 252 eggs,:first calculated the corrected number of eggs aborted in each replicate (eqn. 1), considering 253 only accepted eggs. 254

$$\underline{Abortion efficiency} = \frac{\underline{AbE_AAbE}}{\underline{TotE_AT}} = \frac{\underline{AbE_AAbE}}{\underline{TotE_AT}}$$

256

255

257 Where AbE<sub>A</sub> and TotE<sub>A</sub> are the number of aborted eggs accepted by parasitoids, and the total
 258 number of eggs accepted by parasitoids, respectively.

259 First, we calculated the corrected number of eggs aborted in each replicate (eqn. 1), considering

260 <u>only accepted eggs</u> We then <u>we</u> divided this by the total number of eggs accepted by the parasitoid

261 (TotA), giving the corrected number of aborted eggs per host accepted, which we term "abortion"

- 262 efficiency"...
- 263

264 1.2.3 Laboratory tests of field-collected parasitoids on H. halys eggs

To complement detailed laboratory estimates of parasitoid developmental success and abortion 265 induction with narrow inter-linesgenetiinter-line comparisonse samples withinof two species, we 266 267 investigated variation in these parameters with a larger number of parasitoid species. We assumed that these encompassed a broader genetic diversity, because individuals were collected from sites 268 across a relatively wide geographic region, although we did not directly assess the true amount of 269 genetic variation present with molecular tools. The goal of these experiments was to explore to what 270 extent the evolutionary trap posed by H. halys for native parasitoids generally applies across species 271 present in British Columbia, and to what degree egg abortion caused by native parasitoids may be 272 contributing to biological control of *H. halvs*. 273

We used sentinel egg mass surveys, deploying eggs of the native species *P. maculiventris*, to 274 investigate the diversity of egg parasitoids of stink bugs present in British Columbia and obtain 275 276 parasitoids for laboratory screening against *H. halvs*. From the beginning of May to the end of August of 2017, 1,264 *P. maculiventris* egg masses (21,310 eggs) were set out and retrieved at 10 277 278 sites in coastal (Fraser Valley) and interior (Okanagan Valley) British Columbia (Table 2), on between 2 and 8 different dates per site through the sampling period, on an opportunistic basis. All 279 280 sites were semi-urban, urban, or suburban areas with mixed woody and herbaceous vegetation 281 where breeding populations of *H. halys* had previously been detected (Abram et al., 2017a8; Abram et al., 2019b). Fabric squares containing egg masses (see Insect Colonies above) were stapled to the 282 undersides of leaves of a variety of woody trees (e.g., Prunus spp., Davidia spp., Acer spp., 283 Gleditsia spp., Ailanthus spp., Sorbus spp.) and shrubs (e.g., Rubus sp., Mahonia spp., 284 Symphoricarpos spp., Rosa spp.). The egg masses were then retrieved before nymphs emerged, 3-7 285 days after being set out. Because of the variable retrieval intervals, parasitism rates measured here 286 are intended only to determine which parasitoid species were present and to serve as sources of 287 parasitoids for laboratory tests. 288

Retrieved egg masses were placed in Petri dishes (50 mm diameter, 9 mm depth) and kept 289 under ambient laboratory conditions until parasitoid emergence. Emerging parasitoids were given 290 291 cotton wicks with 10% sucrose as a food source. Newly emerged parasitoids from the Okanagan Valley collections were shipped overnight in a cooler with an ice pack (temperature  $\sim 8-10^{\circ}$ C) to the 292 Agassiz Research and Development Centre (Agassiz, BC) where samples from the Fraser Valley 293 were located, and where all subsequent tests took place in the laboratory. Emerging parasitoids from 294 295 a subset of 139 parasitized egg masses were separated into 1.5 ml plastic Eppendorf tubes at least 296 one day before they were tested. A subset of females was tested (n = 139), with the subset

297 <u>determined by the availability of fresh *H. halys* eggs when parasitoids were of the correct age for</u>

testing (see Table 2 and Figure 4 for per-species sample sizes). All females used in tests emerged in 298 the presence of males from the same egg mass and were thus assumed mated. In all cases, 299 parasitoids were 7-14 days old when used in these tests, enough time for post-emergence ovariole 300 301 maturation to take place but well before the end of their typical lifespan, which is underin the 302 laboratory conditions is (greater than >40 days) (see Abram et al., 2016). For each parasitoid species aAt least 5 replicates were conducted for each parasitoid species. Individual females were 303 transferred from plastic tubes to Petri dish arenas (50 mm diameter, 9 mm depth) containing an H. 304 305 halys egg mass of between 6 and 32 eggs (mean  $\pm$  SE: 20.6  $\pm$  0.52, n=191). The size of the egg masses used depended on the availability of fresh H. halys eggs on each testing day; egg masses 306 were divided into smaller patches when availability of host material was limited, depending on the 307 relative availability of *H. halvs* eggs and adequately aged parasitoids. However, we accounted for 308 variation egg mass size by including it as a covariate in the statistical analysis (see below). In all 309 replicates, parasitoids were removed from dishes after 24 h. Egg masses were then incubated under 310 standard rearing conditions and monitored for the emergence of host nymphs and parasitoids. After 311 being incubated for at least 1 month, they were dissected to record unemerged parasitoids and 312 313 aborted eggs, using the same classification criteria as the first experiment (see above). Throughout the period of the tests, a set of similarly-sized egg masses  $(24.3 \pm 1.0 \text{ eggs}, n=29)$  were set up in 314 315 Petri dish arenas but not exposed to parasitoids as controls, inclubated, and then dissected to record 316 levels of nymph emergence and egg abortion. These trials were run concurrently with the isofemale line experiments described above, under the same abiotic conditions and using H. halys egg 317 318 masses of the same age from the same laboratory colony.

All parasitoids emerging from sentinel egg masses, including those used in laboratory tests
 with *H. halys* eggs, were subsequently preserved in 95% EtOH, point-mounted, and identified based
 on morphological characters (Talamas et al., 2015). Vouchers are deposited in the Florida State
 Collection of Arthropods and Nematodes and the Canadian National Collection of Insects,
 Arachnids, and Nematodes.

324

# 325 1.2.4 Statistical analysis

For laboratory trials where *H. halys* eggs were exposed to different iso-female lines of *T. podisi* and *T. euschisti*, the number of eggs accepted, acceptance latency, acceptance rate, the corrected number of eggs aborted, and abortion efficiency were compared among iso-female lines with Kruskal-Wallis tests followed by Dunn's Kruskal-Wallis multiple comparisons with Benjamini-Hochberg adjustments. -These non-parametric tests were selected after finding that the response variables did not meet the assumptions of generalized linear models (with either Ggaussian or Ppoisson error

distributions) in most cases. To test whether exposure to parasitoids from each iso-female line of the
two species was associated with higher-than-baseline *H. halys* egg mortality, the corrected number
of eggs aborted when exposed to each iso-female line was compared to zero with a Wilcoxon test.

The statistical approach was modified for the experiment measuring levels of egg abortion 335 in *H. halvs* eggs exposed to parasitoids emerging from field-collected *P. maculiventris* egg masses. 336 First, because egg mass size varied among replicates, the proportion (rather than number) of aborted 337 eggs was used as the response variable, and egg mass size (number of eggs) was included as a 338 covariate in the analysis. Second, because multiple parasitoids were sometimes tested from the 339 same sites and natal egg mass (which could result in pseudoreplication if not accounted for), "natal 340 egg mass" and "collection site" were included in the statistical model as random effects. Third, 341 342 because paired unexposed control egg masses could not be run for each every exposed egg mass due to limited H. halys egg availability on some days, the global level of abortion in unexposed egg 343 344 masses set up over the course of the experiment was compared to levels of abortion in egg masses 345 exposed to each parasitoid species. Thus, the proportion of *H. halys*-egg eggs that aborted when 346 exposed to different species of parasitoids or were not exposed to parasitoids, was compared with a generalized linear mixed model (GLMM) with a binomial error distribution, with parasitoid species 347 and egg mass size as fixed effects and natal egg mass as a random effect. Observation-level random 348 effects (Harrison, 2014) were also included to account for overdispersion. 349

All statistical analyses were analyzed using R version 3.6.0 (R <u>Development</u> Core Team, 2019).

352

## 353 **1.3 Results**

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# 355 *1.3.1 Intraspecific variation in acceptance and development*

All iso-female lines of both T. euschisti and T. podisi oviposited in H. halvs eggs, but the number of 356 H. halys eggs in which parasitoids oviposited varied significantly among lines for both T. euschisti 357  $(\chi^2_4 = 26.19, p < 0.0001)$  and T. podisi  $(\chi^2_4 = 29.32, p < 0.0001)$ , with the median number of eggs 358 359 accepted ranging from 15 to 27 for T. euschisti and 7 to 28 eggs for T. podisi, out of the 28 H. halvs eggs offered (Figure 1). Median acceptance latency by T. podisi ranged between 0.34 and 1.68 360 hours but did not differ among lines ( $\chi^2_4 = \frac{26.197.42}{7.42}$ , p = 0.11 < 0.0001) (Figure 2). However, 361 acceptance latency did vary significantly among lines for *T. euschisti* ( $\chi^2_4 = 10.80$ , p = 0.029), with 362 median time until the first observed oviposition ranging from 0.09 to 0.67 hours (Figure 2). The 363 oviposition rate did not clearly differ among the lines of *T. podisi* ( $\chi^2_4 = 8.00$ , p = 0.092; medians 364

365 7.8 to 12.0 ovipositions/hour) or *T. euschisti* ( $\chi^2_4 = 8.81$ , p = 0.066; medians 5.6 to 10.0 366 ovipositions/hour).

There was no evidence in variation in the developmental success of parasitoid offspring among iso-female lines of either parasitoid species. In fact, only a single parasitoid offspring, a *T. euschisti* (line Te\_3) emerged from one of the parasitized *H. halys* egg masses, despite a total of 1,907 eggs being oviposited in by *T. euschisti* across all replicates. We never observed the emergence of *T. podisi* offspring from *H. halys* eggs, which were observed ovipositing in a total of 1,588 *H. halys* eggs across all replicates.

After being corrected for natural levels of *H. halys* egg mortality, the number of eggs 373 aborted by T. podisi varied among iso-female lines ( $\chi^2_4$  =12.22, p = 0.016), with all lines causing at 374 least some *H. halys* egg abortion; median corrected numbers of aborted eggs ranged from 0.5-4.0 375 (Figure 3). These differences in *H. halys* egg abortion induction among lines were maintained when 376 accounting for the number of ovipositions; abortion efficiency also varied among lines of T. podisi 377  $(\chi^2_4 = 18.11, p = 0.0012)$  (Figure 3), with per-strain medians of between 1.7% and 40.6% of 378 attacked eggs aborting. For *T. euschisti*, numbers of aborted eggs did not vary among lines ( $\chi^2_4$  = 379 2.67, p = 0.61), and medians were generally lower (range: 0.5-1.0 eggs), with only three out of five 380 381 lines causing abortion levels significantly above zero (Figure 3). Likewise, abortion efficiency did not vary significantly among the lines of *T. euschisti* ( $\chi^2_4$ =3.43, p = 0.49), with an intraspecific 382 mean of 6.0% of attacked *H. halys* eggs aborting. 383

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# 385 1.3.2 Laboratory tests of field-collected parasitoids on H. halys eggs

Out of the total of 1,264 P. maculiventris sentinel egg masses deployed in the field, 15.3% were 386 parasitized by scelionid egg parasitoids (Table 2). The percentage of egg masses with parasitoid 387 emergence varied among sites from 8.4–24.0%. Of the five scelionid species emerging from 388 sentinel eggs, T. euschisti was present at the greatest number of sites (10/10 sites), followed by Te. 389 390 podisi (9/10 sites), Trissolcus- cosmopeplae (Gahan) (3/10 sites), Tr.- utahensis (Ashmead) (2/10 sites), and Tr. hullensis Harrington (1/10 sites). The latter three Trissolcus spp. were only found at 391 sites in the Okanagan Valley; T. euschisti and T. podisi were found in both the Fraser Valley and 392 Okanagan Valley regions. The only non-scelionid parasitoid reared from P. maculiventris eggs was 393 a single individual in the family Encyrtidae (Table 2). 394

A total of 191 parasitoids, representing the five scelionid species emerging from *P*. *maculiventris* eggs, were tested on *H. halys* eggs in the laboratory. Oviposition was commonly observed at the beginning of these trials by all species tested, and post-oviposition brood guarding was often observed afterwards (P. Abram, personal observations). These informal observations,

399	together with the direct observations on two of these species (T. podisi and T. euschisti) in the iso-
400	female lines experiment under similar conditions, suggest that host acceptance occurred in many of
401	these trials. However, of the 3,935 eggs exposed to these parasitoids, only a single T. euschisti
402	individual emerged from an <i>H. halys</i> egg.
403	Levels of <i>H. halys</i> egg abortion in these experiments were very low, often with a median of
404	zero (Figure 4). The proportion of <i>H. halys</i> egg abortion was not significantly higher in eggs
405	exposed to any of the parasitoid species than in unexposed egg masses and did not vary
406	significantly among parasitoid species to which they were exposed (GLMM, $\chi^2_{4,81}$ = 2.92, p= 0.57)
407	(Figure 4). Levels of egg abortion in exposed egg masses were not associated with variation in <i>H</i> .
408	<i>halys</i> egg mass size ( $\chi^2_{1,81}$ = 0.013, p= 0.91) or its interaction with exposure treatment ( $\chi^2_{4,81}$ = 2.60,
409	p= 0.63).
410	

### 411 1.4 Discussion

Our study adds to a growing body of evidence from laboratory and field studies indicating that 412 native scelionid parasitoids in North America readily oviposit in the eggs of *H. halvs* but rarely 413 develop successfully (Abram et al., 2014; Haye et al., 2015b; Ogburn et al., 2016; Cornelius et al., 414 415 2016a, b; Dieckhoff et al., 2017; Abram et al., 2017b; Gariepy et al., 2019; Konopka et al., 2018, 2019, 2020). In fact, under our standardized laboratory conditions, the developmental success of 416 parasitoid offspring occurred only twice out of the thousands of *H. halvs* eggs that we exposed to 417 parasitoids. In addition, we provide the first evidence that, in at least two native parasitoid species, 418 behavioural parameters related to acceptance of *H. halvs* eggs can vary intraspecifically. Finally, 419 our results suggest that the biological control effects of parasitoid-induced abortion of H. halvs 420 421 eggs, when considering a broader range of native parasitoid genotypes than tested in past studies, is minimal. These findings have implications for the short-term biological control impact of native 422 423 parasitoids on *H. halvs* as well as their longer-term potential to adapt to this invasive host.

Several studies have suggested that native North American parasitoids may eventually adapt 424 to be able to develop in *H. halys* eggs (Abram et al., 2014; Konopka et al., 2018; Tognon et al., 425 2019), and there are other examples where native natural enemies have eventually adapted to 426 successfully exploit invasive prey or hosts (Keeler and Chew, 2008; Berthon et al., 2015). However, 427 in order for natural selection to promote this shift in physiological host range, there would need to 428 be genetic variation present in populations of native parasitoids for traits that influence their ability 429 to successfully complete development in *H. halys* eggs. The low levels of emergence of native 430 parasitoids, including T. podisi and T. euschisti, observed in past field studies using sentinel H. 431 halys egg masses (e.g. Dieckhoff et al., 2017; Ogburn et al., 2016; Herlihy et al., 2016; Cornelius et 432 al., 2016a, b), including in the area where parasitoids were collected for our second experiment 433 434 (Abram et al., 2019b), indicates that this variation may exist. However, it is also possible that other abiotic and biotic factors in the field (e.g., unfertilized eggs, extreme temperatures, desiccation) 435 436 could kill *H. halys* eggs at variable levels, making them more suitable for parasitoid development (similar to freeze-killing; e.g. Haye et al., 2015b), which would overestimate the ability of native 437 parasitoids to develop in viable H. halys eggs. In addition, what proportion of H. halys eggs are 438 located and unsuccessfully parasitized by native parasitoids in the field is unknown unless 439 molecular diagnostic tools are used (Gariepy et al., 2019; Stahl et al., 2019b). Thus, to properly 440 assess intraspecific variation in developmental success, standardized laboratory studies that observe 441 442 parasitoid behaviour, and using *H. halys* eggs with low levels of baseline mortality, are needed in conjunction with H. halys eggs with low levels of baseline mortality. In the current study, we found 443 no evidence to suggest that there is significant intraspecific variation in the ability of T. podisi, T. 444

euschisti, or any of the other scelionid species we tested, to complete development in viable H. 445 446 halvs eggs. Across both experiments, only two individuals of T. euschisti emerged. It is, of course, possible that the hypothesized variation in developmental success exists in some populations of 447 448 North American scelionids; our study did not include any such strains. EHowever, even if initially very rare, after several years of strong selection, parasitoid these variants capable of developing in 449 *H. halvss* eggs would be expected to spread rapidly in areas where *H. halvs* is common. But also, in 450 terms of variation, it's possible that some traits of an appropriate strains may needs different years 451 of strong selections before showing their adaptation or avoidance to the target host. Even if initially 452 rare, these variants would be expected to spread rapidly in areas where H. halys is common. 453 However, Dduring the period of 2013–2016, when our parasitoid lines were established, there is no 454 evidence to suggest that there is significant intraspecific phenotypic variation in the ability of native 455 North American scelionid parasitoids to consistently develop in viable *H. halvs* eggs. However, the 456 data and methodology presented here provide a basis for assessing how this situation could change 457 in the future, and when considering more comprehensive samples of parasitoid strains present in 458 459 areas invaded by H. halvs.

There may be a significant fitness penalty for native parasitoids when they expend energy, 460 time, and eggs while unsuccessfully exploiting H. halys eggs (Abram et al., 2014). Thus, natural 461 selection could act on variation in parasitoid behaviours that mediate acceptance of *H. halys* eggs, 462 leading to avoidance of the evolutionary trap by native parasitoids. We did observe variation in 463 acceptance levels, measured as the number of eggs parasitized in 5 h, of both T. podisi and T. 464 euschisti. Likewise, we observed significant variation in the time it took different iso-female lines 465 466 of *T<sup><i>r*</sup>. euschisti, but not *T. podisi*, to begin ovipositing in *H. halys* eggs. We did not observe any variation in the speed at which either parasitoid species oviposited in *H. halys* eggs once parasitism 467 began. On the one hand, these results could provide preliminary evidence that intraspecific variation 468 exists in behaviours or physiological variables that determine acceptance of *H. halvs* eggs by native 469 parasitoids. For example, greater latency to the first oviposition could indicate that native 470 parasitoids are less able to locate *H. halys* egg masses at a short range or are more likely to delay 471 472 oviposition once the egg mass is encountered. Likewise, fewer total ovipositions by females from some iso-female lines of the two species could indicate that some strains had a greater tendency to 473 leave H. halys egg masses before fully exploiting them. On the other hand, the variation in 474 acceptance and oviposition latency we observed could be underpinned by variation in other aspects 475 of parasitoid behaviour or physiology that are unlikely to be acted upon by natural selection to 476 reduce the level of exploitation of *H. halvs* by native parasitoids. For example, some iso-female 477 478 lines may simply have lower fecundity than others leading to fewer ovipositions before their egg

load is exhausted; or lower locomotor activity leading to a delay in the amount of time required to 479 find and begin ovipositing in *H. halys* eggs. We would not anticipate natural selection to act on 480 these types of traits specifically to avoid the evolutionary trap posed by *H. halys*, as they would also 481 affect their ability to exploit more suitable native host species. Any eventual behavioural avoidance 482 or rejection of *H. halvs* eggs by native parasitoids would need to be mediated by modified 483 sensitivity to specific semiochemical, visual, or physical cues that are specific to *H. halvs* eggs. 484 Future studies that examine intraspecific variation in parasitoid behaviour throughout the process of 485 host searching are needed to better understand the potential for the evolution of behavioural 486 avoidance of *H. halys* eggs. In particular, experiments comparing the behaviour of different strains 487 on a suitable, native host species in addition to *H. halys* (e.g., Abram et al. 2014) could help to 488 determine whether behavioural variation of parasitoids in response to *H. halys* is due to specific 489 parasitoid avoidance or increased acceptance of H. halvs specifically, versus non-specific parasitoid 490 life history variation. 491

Although almost none of the oviposition events by T. podisi and T. euschisti we observed in 492 493 this study resulted in offspring production, some of them caused increases in the number of aborted 494 H. halys eggs. In our first experiment, all five iso-female lines oif T. podisi (1.7-40.6% of attacked 495 eggs aborted) and three out of five lines of T. euschisti (7.4 - 10.0%) of attacked eggs aborted) caused a significant number of H. halys eggs to abort above natural levels (Figure 3). Thus, the 496 unsuccessful attack of *H. halys* eggs by *T. podisi* and *T. euschisti* in some invaded areas may be 497 contributing to increasing its egg mortality. Past studies using a single laboratory strain of T. podisi 498 have recorded increased abortion levels of between approximately 24 and 30% (Abram et al., 2014; 499 Abram et al., 2016) in parasitized eggs, while one other study on two native European Trissolcus 500 spp. estimated abortion rates of about 10-15% on unsuccessfully attacked H. halvs eggs (Haye et 501 al., 2015b). Abortion levels observed in past studies are thus within the range measured in our first 502 experiment. Additionally, our results demonstrate that abortion levels vary between species, and, in 503 some cases, within species. The number of *H. halys* eggs aborted, as well as abortion efficiency, 504 varied among our iso-female lines of *T. podisi*. In contrast, there was no clear intraspecific variation 505 in *H. halys* egg abortion caused by *T. euschisti*, with only three out of five lines causing at least 506 some H. halys eggs to abort and two lines causing no abortion whatsoever. In combination with past 507 studies, the results of our first experiment imply that for *H. halys*, the biological control effect of 508 egg abortion by native parasitoids is likely to vary widely in time and space depending not only on 509 parasitoid species composition but also on the relative frequency of different intraspecific variants 510 causing different levels of egg abortion. 511

Although some parasitoid lines we tested aborted a significant number of *H. halys* eggs, 512 after taking into account variation in acceptance levels and intraspecific variation in abortion 513 efficiency, the absolute numbers of eggs aborted were low overall. While the most efficient strain of 514 T. podisi aborted, on average, about 40% of attacked H. halvs eggs (= 11.3 eggs per completely 515 exploited egg mass of 28 eggs) (Figure 3), many egg masses were not completely exploited (Figure 516 1), and it was more typical for *T. podisi* and *T. euschisti* to abort fewer than four eggs per exposed 517 egg mass (Figure 3). Considering that less than 50% of H. halys egg masses are located and 518 oviposited within by native parasitoids in the field (Gariepy et al., 2019; Konopka et al., 2018), our 519 results indicate that the biological control effect of egg abortion induced by native parasitoids on H. 520 halys in North America is probably limited or even negligible. In fact, in our second set of 521 experiments testing the capacity of native parasitoids emerging from sentinel egg masses of P. 522 maculiventris in British Columbia, we never observed increases in abortion levels of H. halys eggs 523 above natural levels – even for the two species (T. euschisti and T. podisi) that were observed to 524 oviposit in and abort *H. halvs* eggs in the first laboratory experiment. This result could indicate that 525 526 the populations we sampled in British Columbia are composed of strains of native parasitoids that are ineffective at aborting *H. halys* eggs. Because we did not conduct detailed, comprehensive 527 528 behavioural observations in this experiment, however, we cannot rule out that acceptance levels of *H. halys* eggs may have been lower than in the first experiment, and that this may be responsible for 529 the low abortion levels observed. This would be unexpected, however, given that (i) qualitative 530 behavioural observations (oviposition, patch guarding) indicated frequent acceptance; and (ii) the 531 exposure period in the second experiment (24h) was almost five times longer than the first 532 experiment (5h), giving parasitoids much more time to exploit the available eggs under similar 533 abiotic conditions. In any case, our results suggest that native scelionid parasitoids are probably not 534 currently contributing significantly to biological control of *H. halys* in newly invaded areas of 535 British Columbia, either through host egg abortion or offspring development. 536

The susceptibility of *H. halys* to parasitism and egg abortion would depend not only on 537 intraspecific variation in the attacking parasitoid but also potentially on phenotypic variation in 538 589 hosts that mediates their susceptibility -((Kraaijeveld et al., 1998)Kraaijeveld and Godfray, 1999). There is some evidence for genetic differences among geographic populations of *H. halys*, 540 541 including between *e*Eastern and *w*Western North America (Gariepy et al., 2014; Valentin et al., 2017; Abram et al., 2017a8; Cesari et al., 2017), so it is possible that there are differences in the 542 susceptibility of these different populations to parasitoid attack. Variation in host susceptibility is 543 unlikely to be responsible for the significant differences in the traits we measured among lines of 544 545 parasitoids, as host egg masses were randomly assigned among iso-female line treatments, and

546 experiments were run in parallelduring the same time period. However, it is possible that some differences between our study (which used we we stern North American H. halys) and past studies 547 548 (Abram et al. 2014, 2016, which used e Eastern North American H. halys), could be due to the different population of hosts used. The fact that one of the *T. podisi* strains used in our study (Tp 1) 549 is the same as those used in Abram et al. (2014, 2016) allows an informal comparison. In both 550 551 studies, there was no successful -development of this strain in *H. halvs* eggs. In the current study, this strain aborted an average of 12% of attacked eggs, which is considerably lower than the 24-552 30% estimates from past studies. As most aspects of the methodology of the two studies were 553 554 similar, this comparison could indicate a difference in the susceptibility of wWestern North American *H. halvs* to abortion by native parasitoids. However, as some parts of the methodology 555 differed between the two studies – most importantly egg mass size – further research that includes 556 direct comparisons is needed to investigate this matter. 557

Adaptation of native parasitoids to successfully recognize and successfully exploit invasive 558 hosts is typically considered to occur on timescales of multiple decades (Cornell and Hawkins, 559 560 1993). Given our results as well as those of past studies (e.g., Dieckhoff et al., 2017), it appears that for native scelionid parasitoids exploiting *H. halys* in North America, the variation in traits required 561 562 for this adaptation is not yet clearly present, as w<u>We found no evidence of variation in the</u> developmental success of these <u>native</u> parasitoids. Our results do provide some evidence to 563 suggest that variation in behaviours mediating acceptance of *H. halys* may exist, although it is 564 unclear whether avoidance of *H. halvs* eggs by these parasitoids could feasibly be selected for given 565 566 this variation. We found no evidence of variation in the developmental success of the native parasitoids. Determining whether these native parasitoids will adapt to the presence of *H. halvs* by 567 avoiding its eggs, or alternatively by gaining the capacity to exploit them successfully, will 568 requirecould include long-term studies in several areas invaded by *H. halys*, as well as studies of 569 parasitoid behaviour and development across gradients of time since the invasive stink bug has 570 established... 571

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# 573 Credit authorship contribution statement

574 Elena Costi: Validation, Conceptualization, Data curation, Formal analysis, Writing - Original

- 575 Draft, Writing Review & Editing
- 576 Warren Wong: Validation, Data curation, Writing Review & Editing
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- 578 Susanna Acheampong: Review & Editing, Supervision, Resources, Funding acquisition
- 579 Tim Haye: Writing Review & Editing, Supervision, Funding acquisition

580 Elijah J. Talamas: Writing - Review & Editing, Data curation, Funding acquisition

- 581 Lara Maistrello: Writing Review & Editing, Supervision, Funding acquisition
- 582 Paul Abram: Conceptualization, Formal analysis, Resources, Writing Original Draft, Writing -

583 Review & Editing, Visualization, Project administration

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607 **Conflict of Interest** 

All authors declare that they have no conflict of interest.

# 609 Informed consent

610 Informed consent was obtained from all individual participants included in the study

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Figure 1. The number of *H. halys* eggs in which different iso-female lines of *Trissolcus euschisti* 785 and Telenomus podisi oviposited when observed on H. halys egg masses of 28 eggs for 5 h. For 786 lines within each parasitoid species, boxes not labeled with the same letter are significantly different 787 (p < 0.05; Dunn's Kruskal-Wallis multiple comparison test). In boxplots, horizontal lines show 788 medians, boxes contain the 25th-75th percentiles, whiskers show the upper and lower deciles, and 789 points show outliers. 790

791





**Figure 2.** Acceptance latency (time until first oviposition) of different iso-female lines of *Trissolcus euschisti* and *Telenomus podisi* on *H. halys* egg masses. For parasitoid lines within each species, boxes not containing the same letter are significantly different (p < 0.05; Dunn's Kruskal-Wallis multiple comparison test). In boxplots, horizontal lines show medians, boxes contain the 25<sup>th</sup>-75<sup>th</sup> percentiles, whiskers show the upper and lower deciles, and points show outliers.



Figure 3. (A) The corrected number of eggs aborting The number of eggs aborting due to 800 parasitism, and (B) corrected abortion efficiency (number of aborted eggs per parasitoid 801 oviposition) when *H. halys* egg masses were exposed to different iso-female lines of *Trissolcus* 802 803 euschisti and Telenomus podisi, after correcting adjusting for baseline levels of H. halys egg 804 mortality. Negative values for some observations result from higher mortality in unexposed egg masses than exposed egg masses. In both panels, for lines within each parasitoid species, boxes not 805 labeled with the same letter are significantly different (p < 0.05; Dunn's Kruskal-Wallis multiple 806 807 comparison test). In (A), symbols underneath the boxes indicate whether the corrected abortion levels are significantly different from zero (n.s. -p > 0.05; \*-p < 0.05; \*\*\*-p < 0.001; Wilcoxon 808

- test). In boxplots, horizontal lines show medians, boxes contain the 25<sup>th</sup>-75<sup>th</sup> percentiles, whiskers
- show the upper and lower deciles, and points show outliers.



**Figure 4.** Percentage of *H. halys* eggs aborting in unexposed control egg masses (UC), compared to egg masses exposed to one of five parasitoid species in the laboratory for 24 h: Th – *Trissolcus hullensis* (n = 5); Tu – *Trissolcus utahensis* (n = 9); Tp – *Telenomus podisi* (n = 36); Tc – *Trissolcus cosmopeplae* (n = 46); Te – *Trissolcus euschisti* (n = 95). Horizontal lines show medians, boxes contain the 25<sup>th</sup>-75<sup>th</sup> percentiles, whiskers show the upper and lower deciles, and points show outliers. Abortion levels did not vary significantly among treatments (see the text of Results for statistical information).

Species	Line name	State/Province, Country of Collection:	Original	Year of
	Line nume	GPS	emergence host	collection
Telenomus podisi	Tp-1	Quebec, Canada: 45.556, -73.557	P. maculiventris	2013
	Tp-2	Ontario, Canada: 43.040, -81.219	P. maculiventris	2016
	Tp-3	Ontario, Canada: 43.028, -81.213	P. maculiventris	2016
	Tp-4	North Carolina, USA: 35.428, -82.564	H. halys	2016
	Tp-5	Ontario, Canada: 43.074, -81.337	P. maculiventris	2016
Trissolcus euschisti	Te-1	Ontario, Canada: 43.028, -81.213	P. maculiventris	2016
	Te-2	Ontario, Canada: 43.030, -81.209	P. maculiventris	2016
	Te-3	Ontario, Canada: 43.030, -81.209	P. maculiventris	2016
	Te-4	Ontario, Canada: 43.030, -81.209	P. maculiventris	2016
	Te-5	Quebec, Canada: 45.499, -73.346	P. maculiventris	2016

# **Table 1.** Sources of parasitoid iso-female lines used in experiments.

821

**Table 2.** Scelionid parasitoids emerging from *P. maculiventris* egg masses set out and retrieved

from different sites in British Columbia, Canada, and the number of each species from each site that

- 825 was subsequently offered *H. halys* eggs in the laboratory: Te *Trissolcus euschisti*; Tp *Telenomus*
- 826 *podisi*; Tc *Trissolcus cosmopeplae*; Tu *Trissolcus utahensis*; Th *Trissolcus hullensis*.
- 827

	Site Name (GPS)	# sentinel egg masses set out (total # eggs)	% egg masses parasitized per species <sup>a</sup>					
Region			[# individuals lab-tested against <i>H. halys</i> ]					
Region			Те	Тр	Тс	Tu	Th	Total <sup>b</sup>
	Chilliwack #1 (49.158, -122.003)	107 (1,635)	4.67 [1]	3.74 [1]	0.00	0.00	0.00	8.41
	Chilliwack #2 (49.159, -121.997)	55 (807)	14.55 [5]	9.09 [2]	0.00	0.00	0.00	21.82
Fraser Valley	Rosedale (49.184, -121.800)	53 (721)	3.77 [0]	7.55 [3]	0.00	0.00	0.00	11.32
	Abbotsford (49.003, -122.263)	25 (364)	20.00 [3]	4.00 [1]	0.00	0.00	0.00	24.00
	Langley (49.122, -122.657)	10 (138)	10.00 [0]	10.00 [0]	0.00	0.00	0.00	20.00
	Penticton <sup>c</sup> (49.483, -119.603)	579 (10,208)	4.49 [21]	1.04 [7]	8.29 [46]	2.76[9]	0.69 [5]	14.51
	Kelowna #1 (49.885, -119.484)	115 (1,958)	20.00 [31]	0.00	0.00	0.00	0.00	20.00
Okanagan Valley	Kelowna #2 (49.884, -119.456)	106 (1,821)	4.72 [5]	15.09 [15]	0.94[0]	0.00	0.00	18.87
	Kelowna #3 (49.880, -119.485)	104 (1,804)	16.35 [20]	2.88 [2]	0.96[0]	0.96[0]	0.00	20.19
	Kelowna #4 (49.871, -119.490)	110 (1,854)	6.36 [9]	2.73 [5]	0.00	0.00	0.00	9.09
	Total	1,264 (21,310)	7.75 [95]	3.32 [36]	3.96[46]	1.34[9]	0.32[5]	15.34

828 <sup>a</sup> Percentage of egg masses from which at least one individual of <u>a given each</u> species emerged

<sup>b</sup>Percentage of egg masses parasitized. This value is sometimes lower than row totals due to multiparasitism; i.e.,

830 multiple parasitoid species emerging from a single egg mass (this occurred in a total of 18 egg masses).

<sup>c</sup>One individual of an unidentified Encyrtidae emerged from a single egg mass at this site.

# 832 Credit authorship contribution statement

- 833 Elena Costi: Validation, Conceptualization, Data curation, Formal analysis, Writing Original
- 834 Draft, Writing Review & Editing
- 835 Warren Wong: Validation, Data curation, Writing Review & Editing
- 836 Joan Cossentine: Review & Editing, Supervision, Resources, Funding acquisition
- 837 Susanna Acheampong: Review & Editing, Supervision, Resources, Funding acquisition



Iso-female line

Te\_1

n = 20

Te\_2

n = 21

Te\_3

n = 21

Te\_4

n = 20

Te\_5

n = 20

Tp\_5

n = 19

Tp\_4

n = 17

856

0

Tp\_1

n = 20

Tp\_2

n = 17

Тр\_3

n = 22





