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- 1 Intraguild aggressiveness between an alien and a native predatory mite
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 3 Escudero-Colomar, L.A. ⁽¹⁾*; Creus, E. ⁽¹⁾; Chorąży, A. ⁽²⁾; Walzer, A. ⁽³⁾
- 4
- 5 ⁽¹⁾ IRTA-Sustainable Plant Protection (Entomology). Mas Badia Experimental
- 6 Station. Canet de la Tallada S/N- 17134 Girona, Spain.*email:
- 7 <u>adriana.escudero@irta.cat</u>
- 8 0000-0002-2451-962X
- 9 ⁽²⁾ Department of Applied Entomology, Warsaw University of Life Sciences –
- 10 SGGW, Warsaw, Poland
- 11 ⁽³⁾ Division of Plant Protection, Department of Crop Sciences, University of
- Natural Resources and Life Sciences, Peter Jordan Strasse 82, 1190 Vienna, Austria
 0000-0002-8346-581X

14 Abstract

The predatory mite Amblydromalus limonicus, non-native in Europe, can be used 15 legally in several European countries as greenhouse biocontrol agent against thrips 16 species, although this species is also able to feed on whiteflies and gall mites. The first 17 18 record of the unintended occurrence of A. limonicus in apple orchards in Europe comes from Catalonia (Spain), where A. limonicus is well established in the native predatory 19 mite community since 2011. The dominant species in this community is Amblyseius 20 21 andersoni, which has a similar life-style as A. limonicus (large, aggressive predator with broad diet range) making intraguild (IG) interactions between the two predators likely. 22 Thus, we tested the IG aggressiveness of native and alien female predators, when 23 provided with IG prey (larvae). Alien females of A. limonicus proved to be highly 24 aggressive IG predators against native larvae of A. andersoni, which were attacked 25 earlier and more frequently than alien larvae by the native predator. Nearly all attacks 26 by the alien predator resulted in the death of native IG prey, whereas about 10% of the 27 alien intraguild prey escaped the attacks of the native predator. Additionally, native IG 28 prey is smaller than alien prey, which should facilitate the overwhelming by the alien 29 predator. We argue that the strong aggressive intraguild behavior of A. limonicus is 30 31 contributing to its establishment success in the native predatory mite community.

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33 Keywords: Biological control, *Amblydromalus limonicus*, *Amblyseius andersoni*,

34 intraguild predation, invasive species, Phytoseiidae

35 Introduction

36

37 Life-history traits, habitat and diet range of alien species are commonly used to predict their establishment opportunities in novel environments (Kolar and Lodge 2001; 38 Colautti et al. 2006; Hayes and Barry 2008), whereas behavioral traits have been 39 40 seldom considered (Chapple et al. 2011). Behavior, however, mediates how an alien species interact with novel resources and native species and should therefore play a 41 pivotal role in the establishment opportunities of alien species (Holway and Suarez 42 1999; Chapple et al. 2011). A characteristic feature of successful established alien 43 species is their highly aggressive behavior against native species (Chapple et al. 2011) 44 resulting in the superiority of alien species in competitive interactions such as resource 45 competition [alien Argentine ant Linepithema humile Mayr versus native tyrant ant 46 Iridomyrmex bicknelli Emery (Rowles and O'Dowd 2007; Carpintero and Reyes-López 47 2008)], shelter competition [alien crayfish Pacifastacus leniusculus Dana versus native 48 crayfish Cambaroides japonicus (De Haan) (Usio et al. 2001)], and nest site 49 50 competition [alien European starling Sturnus vulgaris Linnaeus versus native house wrens Troglodytes aedon (Vieillot) (Weitzel 1988)]. 51

When an alien predator encounters a native predator guild sharing similar prey 52 53 resources, intraguild (IG) aggressiveness among the invader and its native opponents is a frequent observed event. The degree of IG aggressiveness strongly influences not only 54 the outcome of resource competition, but also the strength and direction of intraguild 55 56 predation (IGP) (Polis et al. 1989; Snyder and Evans 2006; Wang et al. 2013). For example, the Asian ladybeetle Harmonia axyridis Pallas, initially used as alien 57 biological control agent control agent against greenhouse aphids in France, has 58 immigrated and established in native European ladybeetle communities (Slogett 2012). 59 The alien ladybeetle exhibited high IG aggressiveness against the native ladybeetles 60 Coccinella septempuncata (Linnaeus) and Adalia bipunctata (Linnaeus). Therefore, H. 61 axyridis was also superior in IGP, which is assumed a major cause for the successful 62 establishment of H. axyridis in European ladybeetle communities (Raak-van den Berg 63 et al. 2012). 64

The plant-inhabiting predatory mite Amblydromalus limonicus (Garman and 65 66 McGregor) (Acari: Phytoseiidae), alien in Europe, is used in some European countries such as Belgium, France, Germany, The Netherlands, Poland and Austria, as natural 67 enemy against greenhouse thrips species (Garman and McGregor 1956; Moraes et al. 68 69 1994; Chant and McMurtry 2005; Knapp et al. 2013). In 2011, the alien predator was detected for the first time on apple trees in Catalonia (Spain) (Escudero-Colomar and 70 Choraży 2012), where the commercial use of A. limonicus as biocontrol agent is not 71 permitted. In this region the native predatory mite guild in apple orchards frequently 72 consists of the species Amblyseius andersoni (Chant), Neoseiulus californicus 73 McGregor and Euseius stipulatus (Athias-Henriot) (Acari: Phytoseiidae) sharing 74 herbivorous mites, small insects and pollen as food resource (Costa-Comelles et al. 75 1986, Costa-Comelles et al. 1990, Avilla et al. 1993). These native predators provide an 76 important ecosystem service by the natural control of pest species such as spider mites 77 78 (Tetranychus urticae Koch, T. turkestani Ugarov y Nikolski, Panonychus ulmi (Koch)) (Costa-Comelles et al. 1994,) and gall mites (Aculus schlechtendali (Nalepa)) (Duso C 79 and Pasini M 2003). Although all three species are able to use a broad range of food 80 items allowing development, reproduction and survival, their different feeding 81 preferences result in resource partitioning and reduce intraguild aggression among them. 82 *Neoseiulus californicus* is a selective predator of spider mites producing dense webbing, 83 which can also serve as shelter against IG predators. Euseius stipulatus is a pollen-84

feeding generalist predator, whereas A. andersoni is a generalist predator, feeding and 85 reproducing on a wide range of animal and non-animal food without specific 86 87 preferences (McMurtry et al. 2013). The numerical dominant species in this guild on apple trees is A. andersoni, a large, aggressive predator (Walzer and Schausberger 88 2011a, b, 2013), which is responsible for the successful suppression of the spider mite 89 90 Panonychus ulmi (Koch) (Vilajeliu et al, 1994). The now established alien predator A. limonicus, however, has a very similar life-style as the native predator A. andersoni 91 (McMurtry et al. 2013). Thus, the two predator species are currently sharing both space 92 and prey increasing the likelihood of aggressive IG encounters. Consequently, the 93 objective of this study is to evaluate the IG aggressiveness potential between them. 94 These data should provide first indications, whether the native or alien predator is 95 superior in IGP over the other. 96

96 97

98 Material and methods 99

100 *Mite cultures*

101 The predatory mites were collected in September 2015 from apple leaves of organically managed orchards located at the Mas Badia Experimental Station, in La Tallada 102 103 d'Emporda (42.0541°N, 3.0614°E), near Girona, Spain. About 100 specimens of 104 Amblydromalus limonicus and Amblyseius andersoni were used to initiate lab populations, which were reared on separate arenas consisting of plastic tiles (13 cm x 13 105 106 cm) resting on water-saturated foam cubes (14 cm x 14 cm x 3.5 cm) in plastic boxes (20 cm x 20 cm x 4 cm) half-filled with water. The edges of the arenas were covered 107 with filter paper strips. Stripes of cloth of 1 cm width were placed on the filter paper in 108 109 such a way that they had in direct contact with the water of the plastic box to keep the filter paper moisturized. A barrier of fruit tree grease (Vitax Ltd, Leicestershire, UK) 110 was placed around the complete perimeter of the arena. Consequently, the mites were 111 provided with access to water without a risk of escaping from the arena. Small, 112 113 transparent plastic pieces and cotton wool threads were distributed on the surface of the arena to provide shelter and egg deposition sites for the predatory mites. The predatory 114 mites were fed with a mix of stages of T. urticae and Typha latifolia pollen ad libitum 115 three times per week. Rearings of A. limonicus and A. andersoni were kept in climatic 116 chambers at 25°C±1°C, 70±10% RH and 16:8 L:D (Light:Dark). 117

- 118
- 119 *Experimental units*

120 The experiment was conducted using lockable cages consisting of acrylic plates (80 x 121 35 x 3 mm) with a 15 mm diameter hole. Each hole was closed in one side with a fine 122 mesh screen at the bottom, which provided mites with adequate ventilation. The other 123 side of the cage was covered with a microscope slide secured with rubber bands 124 (Schausberger 1997). Cages were kept in a climatic chamber SANYO at $25^{\circ}C \pm 1^{\circ}C$, 125 $70 \pm 10\%$ RH and a photoperiod of 16:8 L:D (Light:Dark).

- 126
- 127 *IG predator and prey behavior*

IGP is a common interaction among phytoseiid mites, which is asymmetric with respect to size. Small/younger juveniles are usually preyed upon by larger/older juveniles and/or adult females, whereas adult females and eggs are relatively invulnerable to IGP (Walzer and Schausberger 2011a). Consequently, large females and small larvae were

used as IG predators and IG prey, respectively.

133 Single IG predator females of *A. limonicus* and *A. andersoni* were isolated and starved 134 for 24 hours in the lockable cages. Only females that laid eggs during this period were

used in the experiments. A single IG prey was placed in the cage together with a single 135 IG predator. 67 to 71 replicates per treatment were conducted. The behavior of IGP 136 137 predator and IGP prey was observed and recorded every 10 minutes for 6 hours (360 minutes in total) using a chronometer Fisherbrand[™] (Fisher Scientific SL, Madrid) to 138 evaluate the following parameters: (1) attack time of the IG predator, (2) the attack 139 140 success (if the attack results in the death of the prey or not), (3) activity of the IG predator and IG prey (moving or not). 141

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143 *IG predator and prey body sizes*

All females used in the experiment were measured and 50 larvae of each species were 144 randomly caught from the rearing boxes. Females and larvae were mounted on 145 microscopic slides using Heinze polyvinyl alcohol (PVA Heinze) (Heinze 1952; Colloff 146 2009). The distances between the bases of eight and six setae, i.e. j3, s4, S4, Z5 and j3, 147 s4 and Z5 at both sides of the dorsal shield were measured for IG predator females and 148 IG prey larvae of both predatory mites, respectively (Croft et al. 1999) (Fig. 1). The 149 calculated perimeter is a suitable approximation to body size, because it integrates both 150 body length and width and corresponds to 85-90% of the total perimeter of the dorsal 151 The pictures of the figures were taken using a Leica DMRXA Direct 152 shield. 153 microscope equiped with a digital camera Pointgrey Flea3 of 12MPx. The 154 measurements were done using the Nikon Eclipse 50i microscpe using the software NIS-D Elements 3.20 (Nikon Corporation, Japan). 155

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Fig 1 Body size measurements a) females (A. limonicus) and b) larvae (A. andersoni). 159 The distances between the bases of eight (j3-j3, j3-s4, s4-S4, S4-Z5, Z5-Z5) and six (j3-160 161 13, 13-s4, s4-Z5, Z5-Z5) setae on the dorsal shield were measured for the females (IG predators) and larvae (IG prey), respectively. The calculated perimeter of the dorsal 162 shields was used as proxy for body size 163

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167 *Data analysis*

SPSS 21.0.1 (SPSS Inc., 2012) was used for all statistical analyses. Kaplan-Meier 168 analysis with successive Breslow tests was used to compare the IG aggressiveness 169 functions of the predators (combination of cumulative attacks and attack time) (Bewick 170 171 et al. 2004). Generalized linear models (GLMs) with chi-square statistics (Wald-tests) 172 were used to analyze the effects of IG predator species on the predator activity, prey activity (moving or not), IGP success (prey killed or not), IG prey survival (yes/no) 173 174 (binomial distribution, logit link function). Additionally, the differences between the 175 two species with respect to the predator body size (females), the prey body size (larvae) and the IG predator/prey body size ratio were analyzed using GLM's (normal 176 distribution, identity link function). 177

178 179

180 Results

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182 *IG predator and prey behavior*

The IG aggressiveness functions of the IG predators differed significantly between the two predator species (Kaplan-Meier analysis, Breslow tests: $\chi^{2}_{1} = 10.834$, p = 0.001). The alien IG predator *A. limonicus* was more aggressive than the native predator *A. andersoni*, which was reflected in earlier attacks (mean attack times of *A. limonicus*: 48.31 min ± 5.55 SE versus *A. andersoni*: 103.78 ± 5.55) and more frequent attacks on IG prey larvae (Fig. 2).

Both IG predator ($\chi^2_1 = 0.142$, p = 0.706) and prey activity ($\chi^2_1 = 0.723$, p = 189 0.395) were not influenced by species affiliation (Fig. 3). Contrary, IGP success (χ^2_1 = 190 191 4.569, p = 0.033) of the alien predator was higher than for the native predator. First, almost all alien IG predators were observed to attack the IG prev within 360 min in 192 contrast to the native IG predators (alien A. limonicus: 69 out of 71; native A. andersoni: 193 58 out of 67). Second, nearly all attacked native IG prey larvae were killed by the alien 194 predator females, whereas about 10% of the alien IG prey larvae survived the attacks of 195 the native IG predator females. Thus, the survival probabilities of the native IG prey 196 were negligible in contrast to the alien IG prey ($\chi^2_1 = 9.500$, p = 0.002) (Fig. 3). 197

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199 *IG predator and prey body sizes*

The IG predator females of *A. limonicus* and *A. andersoni* were similarly sized ($\chi^{2}_{1} = 0.402$, p = 0.526); however, IG larval prey sizes differed between the alien and native species ($\chi^{2}_{1} = 13.343$, p < 0.001). The alien IG prey was larger than the native IG prey (Fig. 3). Consequently, also the IG predator/prey body size ratio of the alien predator and native prey was larger than the corresponding data of the native predator and alien prey (Fig. 4).

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Fig 2 Attack functions (combination of the proportion of attacking predators and the
time elapsed until an attack occurred) of singly caged native IG predators females of *A*. *limonicus* (black line) and *A. andersoni* (grey line), when offered single IG prey larvae
over 360 min

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Fig 3 IG predator activity (A), IG prey activity (B), IG predation success (C) and IG prey survival (D) of *A. limonicus* (black bars) and *A. andersoni* (grey bars), when acting as IG predator (female) or IG prey (larva) over 360 min. Different lower-case indicate significant effects between the species based on pairwise LSD-tests



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Fig 4 Body size of *A. limonicus* (black bars) and *A. andersoni* (grey bars) as IG predator (females) or IG prey (larvae), and the predator/prey body size ratios. Different lowercase indicate significant effects between the species based on pairwise LSD-tests

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226 **Discussion**

Alien females of *A. limonicus* proved to be highly aggressive IG predators against native larvae of *A. andersoni*, which were attacked earlier and more frequently than alien IG prey by the native predators. Additionally, nearly all attacks by the alien predator resulted in the death of native IG prey, whereas about 20% of the alien IG prey survived in the presence of native predators.

Both predator and prey traits may explain the proximate reasons for these

results. A specific feature of A. limonicus females, not known for A. andersoni females, 234 is their capability to overwhelm and kill 2nd instar thrips larvae (Van Houten et al. 235 1993), which are about double as large as the predator (Schuster and Pritchard 1963; 236 Van Houten et al. 1993). In concert with our results, these findings indicate that alien A. 237 limonicus females have a higher tendency to attack heterospecific individuals than 238 239 native A. andersoni females. Second, the IG predators are similarly sized, but not IG prey. The native IG larval prey is smaller than the alien IG prey, which facilitates the 240 overwhelming of the former by the alien predator. Additionally, the alien larvae are 241 obligatory feeders (Walzer, personal observation), whereas native larvae are facultative 242 feeders (Schausberger and Croft 1999a). Thus, alien A. limonicus larvae might be more 243 successful in defending themselves against IG predator attacks than the native A. 244 245 andersoni larvae.

High interspecific aggressiveness is often correlated with foraging behavior (Sih 246 et al. 2004), whereas more aggressive species are strong food competitors or superior 247 IG predators compared to less aggressive species (Schausberger and Croft 1999b, 248 2000a; Drescher et al. 2011). Dominance in IGP offers three adaptive benefits for the 249 alien A. limonicus: (1) the killing of juvenile native IG prey reduces the number of 250 potential adult IG predators and consequently increases the survival probabilities of 251 252 alien offspring; (2) the consumption of native predatory mites probably offers an 253 additional high-quality food resource for the generalist predator in the novel habitat (Schausberger and Croft 2000b); and (3) the decimation of a native food competitor 254 255 alleviates the access to extraguild prey.

We assume that the implications of the high interspecific aggressiveness of A. 256 *limonicus* are strongly dependent on the context. For example, acting as biocontrol 257 258 agent against greenhouse thrips species, the aggressive predation behavior of A. *limonicus* observed in the present study should have benefits, because A. *limonicus* can 259 260 kill a higher proportion and larger individuals of the target pest than other phytoseiid mites (Van Houten et al. 1995). However, when acting as alien IG predator in the case 261 of escape from greenhouses to novel environments, high IG aggressiveness may allow 262 A. limonicus to become established in native predatory mite communities. Additionally, 263 the capacity for population increase of A. limonicus is among the highest values 264 reported for phytoseiid mites (Sabelis 1985; Steiner et al. 2003) and clearly exceeds the 265 corresponding values of A. andersoni (Lorenzon et al. 2012). Nonetheless, so far the 266 alien predator A. limonicus was not able to eliminate the native predator A. andersoni on 267 268 apple trees in Catalonia. In contrast, alien and native predatory mites still co-exist since the detection of A. limonicus in 2011. Five non-mutually exclusive causes may be put 269 forward to explain these findings. First, the availability of alternative food resources and 270 high structural habitat complexity may lower the potential detrimental effects of IGP on 271 the native species, as it was demonstrated in two other phytoseiid mite species 272 (Pozzebon et al. 2015). Second, strong egg cannibalism of the alien predator species is a 273 well-documented phenomenon, which seems not dependent on the quality of available 274 food resources (Vangansbeke et al. 2014). Such a behavior should reduce the capacity 275 for population increase, so that A. limonicus is not able to build up sufficient high 276 277 population densities to outcompete native species. Third, the local climatic conditions may negatively influence the population growth and dispersal abilities of the alien 278 predator. The relative humidity can be very low during the vegetation period in the 279 apple producing area of the extreme NE of Catalonia (Ninyerola et al. 2001). 280 Amblydromalus limonicus is a species with high moisture requirements, being a 281 determining factor for the hatching success of their larvae (Bakker et al. 1993; Van 282 Houten et al. 1995). For example, in contrast to A. andersoni, the densities of A. 283

limonicus were very low on apple trees in the dry-hot summer 2012 in Catalonia 284 (Escudero-Colomar, personal observation), which could be caused by high egg 285 mortality of the alien predator. In California, native A. limonicus populations were 286 found only along the shore, but not in the drier inland regions (McMurtry and Scriven 287 1965; McMurtry et al. 1971). Along the same line, alien A. limonicus populations were 288 289 only sampled in high densities near to the seacoast in Catalonia (Choraży et al. 2016). Thus, the sensitivity of A. limonicus against dry-hot conditions may restrict its dispersal 290 potential to the coastal areas in Mediterranean countries. Fourth, the leaf morphology of 291 plants shared by A. limonicus and the native predatory mites may affect the strength of 292 their IG interactions (Seelmann et al. 2007). The alien predator A. limonicus is known to 293 avoid pubescent leaves (Lee and Zhang 2018), whereas A. andersoni is often found on 294 pubescent leaves (Overmeer and Van Zon 1984; Duso and Pasini 2003). Such 295 differential habitat selection should reduce encounters between the native and alien 296 predator. Finally, our results suggest that potential IG interactions between the alien 297 predator A. limonicus and the native predator A. andersoni could be mainly 298 unidirectional with A. limonicus as predator and A. andersoni as prey. Thus, the 299 selection pressure to cope with IG interactions should be lower on predator than on 300 prey, because it is less costly to miss a meal (in the case of the predator) as to be one (in 301 302 the case of the prey). Therefore, the native predator A. andersoni may have evolved 303 adaptive anti-predator strategies to cope with the presence of the aggressive alien IG predator. A common behavior in predatory mites is the spatiotemporal avoidance of 304 305 leaves occupied by high-risk IG predators (Walzer et al. 2006; Walzer et al. 2009), which is often optimized by learning (Walzer and Schausberger 2011a, 2012). It 306 remains an open question, whether experienced A. andersoni females respond in a 307 308 similar manner to alien IG predator cues.

309

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