

# Host-mediated, cross-generational intraspecific competition in a multivoltine herbivore species

Bastien Castagneyrol, Inge van Halder, Yasmine Kadiri, Laura Schillé, Hervé

Jactel

## ▶ To cite this version:

Bastien Castagneyrol, Inge van Halder, Yasmine Kadiri, Laura Schillé, Hervé Jactel. Host-mediated, cross-generational intraspecific competition in a multivoltine herbivore species. 2021. hal-03276170

## HAL Id: hal-03276170 https://hal.inrae.fr/hal-03276170

Preprint submitted on 1 Jul 2021

**HAL** is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers. L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.



Distributed under a Creative Commons Attribution 4.0 International License

#### Host-mediated, cross-generational intraspecific competition in a multivoltine herbivore species 2 Bastien Castagnevrol<sup>1,\*</sup> Inge van Halder<sup>1</sup> Yasmine Kadiri<sup>1</sup> 3 Laura Schillé<sup>1</sup> Hervé Jactel<sup>1</sup> 4 <sup>1</sup> Univ. Bordeaux, INRAE, BIOGECO, F-33612 Cestas, France 5 6 \* Bastien Castagnevrol, INRAE UMR BIOGECO, 69 route d'Arcachon, FR-33612 Cestas 7 Cedex, France; bastien.castagneyrol@inrae.fr 8 9 August 9, 2020 10 Conspecific insect herbivores co-occurring on the same host plant interact both directly through 11 interference competition and indirectly through exploitative competition, plant-mediated interac-12 tions and enemy-mediated interactions. However, the situation is less clear when the interactions 13 between conspecific insect herbivores are separated in time within the same growing season, 14 as it is the case for multivoltine species. We hypothesized that early season herbivory would 15 result in reduced egg laying and reduced performance of the next generation of herbivore on 16 previously attacked plants. We tested this hypothesis in a choice experiment with box tree moth 17 females, Cydalima perspectalis Walker (Lepidoptera: Crambidae), to which box trees, Buxus 18 sempervirens L. (Buxaceae), were proposed that had or had not been previously defoliated by 19 BTM larvae earlier in the season. We then compared the performance of the next generation larvae 20 on previously damaged vs undamaged plants. Previous herbivory had no effect on oviposition 21 behaviour, but the weight of next generation larvae was significantly lower in previously damaged 22 plants. There was a negative correlation between the number of egg clutches per plant and larval 23 performance. Overall, our findings reveal that early season herbivory reduces the performance 24 of conspecific individuals on the same host plant later in the growing season, and that this 25 time-lagged intraspecific competition results from a mismatch between the oviposition preference 26 of females and the performance of their offspring. 27

## <sup>28</sup> 1 Introduction

Insect herbivores exploiting the same plant can compete for food, even when interactions among individuals 29 are separated in time (Kaplan & Denno, 2007). Insects may thus reduce the impact of inter- and intraspecific 30 competition by avoiding crowded plants, or plants that have been previously consumed by herbivores, which 31 supposes that they can detect competitors or their effects on plants (De Moraes et al., 2001; Shiojiri & 32 Takabayashi, 2003). For many species, the choice of the oviposition site by mated females is crucial in 33 this respect. The preference-performance hypothesis - aka the 'mother knows best hypothesis' - states that 34 female insects evolved host searching behaviour that leads them to oviposit on hosts where their offspring 35 do best (Gripenberg et al., 2010). A good match between the preference of a mated female for a given 36 plant and the performance of their offspring developing on the same plant implies that females can recognize 37 cues that correlate with larval performance, in particular those related to plant defenses and nutritional 38 quality. Yet, these cues can be largely modified by the simultaneous or sequential presence of other competing 39 herbivores (Bultman & Faeth, 1986; Nykänen & Koricheva, 2004; Abdala-Roberts et al., 2019; Visakorpi et 40 al., 2019). Therefore, initial herbivory may have time-lagged consequences of the preference and performance 41

<sup>42</sup> of herbivores that subsequently attack the same plant in the same growing season (Poelman et al., 2008;

43 Stam et al., 2014). However, while such time-lagged *interspecific* interactions between herbivores have long

<sup>44</sup> been documented (Faeth, 1986), surprisingly much less is known about delayed *intraspecific* interactions in
 <sup>45</sup> multivoltine species.

Previous herbivory can influence the oviposition preference of later herbivores. Several studies have demon-46 strated that mated females can discriminate host plants that have been previously attacked by insect herbivores 47 (Wise & Weinberg, 2002; Stam et al., 2014; Moura et al., 2017; Barnes & Murphy, 2018; Moreira et al., 48 2018; Weeraddana & Evenden, 2019). This behaviour involves the use of a mix of plant and herbivore 49 cues to detect the passage of competitors on potential hosts. Several mechanisms can explain such indirect 50 interactions between herbivores separated in time. First, mated females can directly detect the present, past 51 and possibly future presence of competitors. For instance, Averill & Prokopy (1987) showed that female 52 Rhagoletis pomonella Walsh (Diptera: Tephritidae) marks its oviposition site with an epideictic pheromone 53 that deter conspecific females from laying eggs, thus reducing intraspecific competition at the larval stage. 54 The frass of several Lepidoptera species was also found to act as an oviposition deterrent to subsequent 55 herbivores (Jones & Finch, 1987; Hashem et al., 2013; Molnár et al., 2017). Second, herbivory can induce 56

57 changes in the physical and chemical characteristics of attacked plants (Marchand & McNeil, 2004; Blenn et

al., 2012; Fatouros et al., 2012; Hilker & Fatouros, 2015; Abdala-Roberts et al., 2019), thereby reducing their

<sup>59</sup> attractivity to mated females.

Previous herbivory generally reduces the performance of later herbivores. A common mechanism is that 60 herbivory induces changes in plant quality and defenses that generally reduce the performance of late coming 61 herbivores (Wratten et al., 1988; Agrawal, 1999; Abdala-Roberts et al., 2019). For instance, prior damage by 62 the western tent caterpillar Malacosoma californicum Packard (Lepidoptera: Lasiocampidae) induces the 63 regrowth of tougher leaves acting as physical defenses and reducing the fitness of the next tent caterpillars 64 generation (Barnes & Murphy, 2018). However, several authors reported a mismatch between prior herbivory 65 effects on female oviposition preference vs larval growth, consumption or survival of their offspring (Wise 66 & Weinberg, 2002; Bergamini & Almeida-Neto, 2015; Martinez et al., 2017). For instance, Weeraddana 67 and Evenden (2019) found that herbivory by the diamondback moth, *Plutella xylostella* (L.) (Lepidoptera: 68 Plutellidae) on canola plants (Brassica napus L.) had no effect on subsequent oviposition by the bertha 69 armyworm, Mamestra configurata Walker (Lepidoptera: Noctuidae) whereas its larvae had reduced growth 70 on previously damaged plants. It follows that a sound understanding of the effect of prior herbivory on 71 subsequent herbivory requires studying its effect on the preference and performance of later herbivores 72 simultaneously. 73

The box tree moth (BTM) Cydalima perspectalis Walker (Lepidoptera: Crambidae) is a multivoltine moth 74 species introduced to Europe in 2007 from Asia (Wan et al., 2014). In its native range, BTM larvae can 75 feed on different host genera, whereas in Europe they feed exclusively on box trees (Buxus spp) (Wan et al., 76 2014). In the introduced area, the BTM overwinters at the larval stage, mainly in the third instar, such that 77 defoliation restarts at the beginning of the growing season. In Europe, damage is aggravated by the fact 78 that the BTM has 3-4 generations a year (Kenis et al., 2013; Matošević et al., 2017). When several pest 79 generations successively defoliate the same box tree, there are no leaves left to eat and the caterpillars then 80 feed on the bark, which can lead to the death of the host tree (Kenis et al., 2013; Wan et al., 2014; Alkan 81 Akıncı & Kurdoğlu, 2019). In the present study, we investigated the consequences of defoliation by the first, 82 spring generation of BTM larvae on the oviposition behaviour of the adults and the larval performance in 83 the next generation. We hypothesized that early herbivory would reduce oviposition on previously attacked 84 hosts and reduce the performance of the next generation larvae feeding on damaged hosts. By addressing the 85 above, our study brings new insights into the understanding of cross-generational intraspecific competition in 86 insect herbivores and further challenges the 'mother knows best hypothesis'. 87

## **2** Materials and methods

#### <sup>89</sup> 2.1 Biological material

In spring 2019, we obtained box trees from a commercial nursery and kept them in a greenhouse at INRAE Bordeaux forest research station. Box trees were on average 25 cm high and 20 cm wide. We transferred them into 5 L pots with horticultural loam. For two months, we watered them every four days from the above (*i.e.* watering leaves too) to remove any potential pesticide remain.

BTM larvae (L1-L3) overwinter in cocoons tied between two adjacent leaves. We initiated BTM larvae rearing with caterpillars collected in the wild in early spring 2019. We reared them at room temperature in 4320 cm<sup>3</sup> plastic boxes, and fed them *ad libitum*, with branches collected on box trees around the laboratory. At 25°C, the BTM achieves one generation in 45 days. The larval phase lasts for about 30 days. Adults live 12-15 days. A single female lays on average 800 eggs. We initiated the herbivory on box tree plants with larvae which were the offspring of the overwintering generation. Their adults were used in the preference test, and second generation larvae used in the performance test (see below).

#### <sup>101</sup> 2.2 Experimental design

On June 18<sup>th</sup> 2019, we haphazardly assigned box trees to *control* and *herbivory* experimental groups. The 102 herbivory treatment consisted of n = 60 box trees that received five L3 larvae each. Larvae were allowed to 103 feed freely for one week, after which we removed them all from plant individuals. In order to confirm that 104 the addition of BTM larvae caused herbivory, we visually estimated BTM herbivory as the percentage of 105 leaves consumed by BTM larvae, looking at every branch on every plant. We then averaged herbivory at the 106 plant level. Herbivory ranged from 2 to 18% and was on average 9%. The *control* group (n = 61) did not 107 receive any BTM larva. On July  $8^{th}$ , we randomly distributed plants of the herbivory and control treatments 108 on a  $11 \times 11$  grid in a greenhouse. We left 40 cm between adjacent pots, which was enough to avoid any 109 physical contact between neighbouring plants (Figure 1, Figure 2). 110



Figure 1: Photos illustrating the experimental design.



Figure 2: Experimental design. Pots were 40 cm appart. Circles and triangles represent control and non-attacked trees. Scale colour represents the number of egg clutches per box tree (log-transformed).

<sup>111</sup> The same day, we released *ca* 100 BTM moths that had emerged from chrysalids less than two days before

(i.e., an uncontrolled mix of males and females). Moths were allowed to fly freely within the greenhouse.

 $_{113}$   $\,$  They could feed on small pieces of cotton imbibed with a sugar-water solution, disposed on the ground in the

114 greenhouse.

It is important to note that at the time we released moths, there were no larvae feeding on experimental box 115 trees anymore. In addition, at this time, plants in the herbivory treatment had been cleared of caterpillars for 116 three weeks during which they were watered every two to three days from above. Although larval frass may 117 have been present in pots submitted to the herbivory treatment, it should have been washed out. Finally, we 118 carried out our experiment in an enclosed greenhouse in which the potential effect of natural enemies on 119 BTM behaviour can be neglected. The consequences are that any effect of prior herbivory on subsequent 120 oviposition behaviour and larval performance should have been independent of cues emitted by BTM larvae 121 themselves or by their frass (Sato et al., 1999; Molnár et al., 2017) and therefore were only plant-mediated. 122

#### 123 2.3 BTM host choice

<sup>124</sup> In order to test whether initial defoliation of focal plants influenced host choice for oviposition by BTM <sup>125</sup> females, we counted egg clutches on every branch of every box tree on July 17<sup>th</sup>. Once eggs were counted, we <sup>126</sup> moved box trees to another greenhouse. We installed box trees in plastic saucers. We interspaced plants and <sup>127</sup> filled saucers with a few centimeters of water (renewed regularly) to prevent larvae from moving from one <sup>128</sup> potted plant to another.

#### <sup>129</sup> 2.4 BTM growth rate

Fifteen days later (July 31<sup>st</sup>), we counted larvae on every plant and haphazardly collected five L3 BTM larvae per box tree. We kept them in Petri dishes without food for 24h and weighted them to the closest 10 µg. In some Petri dishes, we observed cases of cannibalism such that in some instances we could only weight two larvae. We therefore calculated the average weight of a L3 larva, dividing the total mass by the number of larvae. Because we did not record the day every single egg hatched, we could not quantify the number of days caterpillars could feed and therefore simply analysed the average weight of a L3 larva. <sup>136</sup> Larvae were allowed to complete their development on the potted box trees. After every larvae pupated, we <sup>137</sup> counted the number of chrysalis per box tree and weighted them to the closest 10 µg.

At this stage of the experiment, box trees that had hosted BTM were close to 100% defoliated so that it is possible that some larvae ran out of food before they pupated. In addition, we noticed that the number of chrysalis in 32 control plants (out of 60, *i.e.* 53%) was greater that the number of larvae (and only in 1 previously attacked plant) suggesting that in spite of our precautions some larvae could have moved from

142 attacked to control plots (Table 1). We therefore decided not to analyze data on chrysalis.

#### 143 2.5 Analyses

<sup>144</sup> All analyses were ran in R using libraries *nlme* and *car* (Fox et al., 2016; Team, 2018; Pinheiro et al., 2020).

<sup>145</sup> We first looked for spatial patterns in female BTM oviposition. We ran a generalized least square model

 $_{146}$  (GLS) testing the effect of x and y coordinates on the number of clutches per plant (*log*-transformed) from

which we explored the associated variogram using the functions gls and Variogram in the nlme library. There

was evidence that oviposition was spatially structured, with strong spatial autocorrelation between 1 and 3m (Figure S1).

We tested the effect of prior herbivory on female BTM oviposition while controlling for spatial non-independence using a GLS. We had no particular hypothesis regarding the shape of the spatial correlation structure. We

<sup>152</sup> therefore ran separate models with different spatial correlation structures (namely, exponential, Gaussian,

spherical, linear and rational quadratic), as well as a model with no correlation structure, and compared them based on their AIC (Zuur, 2009). For each model, we computed the  $\Delta AIC$  (*i.e.*,  $\Delta_i$ ) as the difference

between the AIC of each model i and that of the model with the lowest AIC (Burnham & Anderson, 2002).

<sup>156</sup> We report and interpret the results of the model with the lowest AIC (see *Results*).

We then tested the effect of prior herbivory on BTM performance using ordinary least square models, with the mean weight of five L3 larvae (*log*-transformed) as a response variable, the herbivory treatment (non-attacked *vs* attacked) as a two-levels factor and the number of egg clutches as a covariate.

## 160 **3** Results

We counted a total of 818 egg clutches and 593 larvae. At individual box tree level, the number of egg clutches varied from 0 to 25 (mean  $\pm$  SD: 6.76  $\pm$  5.11, **Figure 2**).

When modelling the effect of prior herbivory on the number of egg clutches, the best model (*i.e.*, model 6 with  $\Delta_i = 0$ , **Table 2**) was the model with a rational quadratic spatial correlation. It was competing with two other models with  $\Delta_i < 2$  (**Table 2**). The  $\Delta_i$  of the model with no correlation structure was > 13, confirming that accounting for spatial autocorrelation among plants improved model fit.

<sup>167</sup> Prior herbivory had no significant effect on the number of egg clutches per plant (model 6:  $\chi^2 = 2.91$ , P = 0.088, Figure 3A). Competing models 2 and 3 provided the same conclusion.

<sup>169</sup> The weight of BTM larvae varied from 6 to 54 mg (mean  $\pm$  SD: 20  $\pm$  9 mg). BTM larval weight was lower

<sup>170</sup> on box trees that had been previously defoliated (**Table 3**, **Figure 3B**). There was a significant, negative

Table 1: Repartition of egg clutches, larvae and chrysalis across box trees with or without prior herbivory. Numbers correspond to mean  $(\pm sd)$  and total number of egg clutches, larvae or chrysalis (n).

Response variable	Control	Herbivory treatment
Egg clutches	6.1 (4.87), n = 372	7.43 (5.3), $n = 446$
Larvae	$4.84 \ (0.61), n = 295$	4.97 (0.18), n = 298
Chrysalis	6.8 (5.78), n = 415	1.85 (1.79), n = 111

Table 2: Summary of AIC of GLS models testing the effect of prior herbivory on the number of egg clutches with different spatial correlation structures.

Model	Correlation structure	df	AIC	Delta
Model 1	None	3	262.5	13.1
Model 2	Exponential	5	249.8	0.4
Model 3	Gaussian	5	250.2	0.8
Model 4	Spherical	5	250.9	1.5
Model 5	Linear	5	255.1	5.7
Model 6	Rational quadratic	5	249.4	0.0

relationship between larval weight and the number of egg clutches on a box tree (Table 3, Figure 3B),
 suggesting intraspecific competition for food. There was no significant interaction between the herbivory
 treatment and the number of egg clutches, indicating that intraspecific competition was independent of prior
 herbivory (Table 3). The model explained 29 % variability in larval weight.



Figure 3: Effects of prior herbivory and conspecific density on (A) the number of egg clutches and (B) L3 larva weight. In A, grey dots represent raw data. Black dots and vertical bars represent raw means (+/- SE). In B, dots represent raw data. Black and grey curves represent model predictions for control and herbivory treatments, respectively.

## 175 4 Discussion

Our findings reveal that early season herbivory reduces the performance of conspecific individuals that subsequently attack the same host plant later in the plant growing season, and that this time-lagged intraspecific competition results from a mismatch between female oviposition preference and the performance of its offspring.

Prior herbivory had no effect BTM oviposition choice. Possible explanations are that prior herbivory
 had no effect on box tree characteristics, or that female BTM were indifferent to them.

Table 3:	Summary	of models	testing t	the effect	of prior	herbivory	and	initial	egg	$\operatorname{clutch}$	density	on	BTM	larval
weight														

Predictor	df	Estimate (SE)	F-value	P-value
Number of egg clutches	1,117	-0.03(0.01)	20.20	< 0.001
Herbivory	1, 117	-0.34(0.1)	20.30	< 0.001
Eggs x Herbivory	1, 117	$0.01 \ (0.01)$	0.73	0.396

The first explanation seems unlikely as numerous studies have clearly established that insect herbivory induces 182 changes in plant physical and chemical traits, which have profound consequences on herbivores or herbivory 183 on the same host plant latter in the season (Wise & Weinberg, 2002; Poelman et al., 2008; Stam et al., 2014; 184 Abdala-Roberts et al., 2019; but see Visakorpi et al., 2019). For instance, Barnes and Murphy (2018) recently 185 observed that the number of eggs laid by western tent caterpillar (Malacosoma californicum) females was 186 much lower on chokecherry trees (Prunus virginiana) that have been defoliated by conspecific caterpillars 187 earlier in the season, as a likely result of induced increase in leaf toughness. In our case, it is also possible 188 that induced defense reactions were delayed in box trees, or that they were already relaxed when we released 189 BTM moths three weeks after the herbivory treatment (Karban, 2011), which remains to be evaluated. 190

We cannot dismiss the second explanation that BTM females were indifferent to box tree cues related to 191 earlier herbivory. This may be particularly true in species whose females individually lay several hundred 192 eggs, thus corresponding to a within-generation bet-hedging strategy (Root & Kareiva, 1984; Hopper, 1999). 193 Consistently, Leuthardt and Baur (2013) observed that BTM females evenly distributed egg clutches among 194 leaves and branches, and that oviposition preference was not dictated by the size of the leaves. Assuming that 195 this behavior is reproducible, the close distance between box-trees that we used in the present experiment (40 196 cm) could explain the lack of effect of initial defoliation on BTM oviposition behavior. In addition, Leuthard 197 et al. (2013) showed that BTM larvae are able to store or metabolise highly toxic alkaloid present in box tree 198 leaves. Last, BTM larvae proved to be unable to distinguish between box tree leaves infected or not by the 199 box rust *Puccinia buxi*, while their growth is reduced in the presence of the pathogenic fungus (Baur et al., 200 2019). Altogether, these results suggest that BTM female moths are not influenced by the amount of intact 201 leaves and probably not either by their chemical quality when choosing the host plant, perhaps because of 202 their strong ability to develop on toxic plants. 203

Prior box tree defoliation by the spring generation of BTM larvae reduced the growth of the 204 next generation. Two alternative, non-mutually exclusive mechanisms can explain this pattern. First, 205 the reduced performance of individuals of the second generation can have been trait-mediated and have 206 resulted from induced plant defenses. This explanation is in line with studies that have documented in 207 several plant species reduced herbivore performance and changes in plant-associated herbivore communities 208 linked to induced defenses after prior herbivory (Nykänen & Koricheva, 2004; Karban, 2011; Stam et al., 209 2014). In the case of multivoltine species, negative relationship between prior herbivory and subsequent larva 210 growth rate would indicate intraspecific plant-mediated cross-generation competition between cohorts of 211 herbivores separated in time (Barnes & Murphy, 2018), which could influence herbivore population dynamics 212 and distribution across host individuals. However, this explanation is partially conflicting with previous 213 observations that BTM larva growth rate did not differ significantly among box-tree varieties, suggesting 214 broad tolerance to variability in host traits (Leuthardt et al., 2013). Alternatively, reduced performance on 215 previously defoliated plants may partly result from food shortage and increased exploitative competition 216 among larvae of the same cohort. Although free living mandibulate herbivores were described to be less 217 sensitive to competition (Denno et al., 1995), the effect of food shortage may have been exacerbated by the 218 small size of box trees and interference competition, for instance through cannibalism (Kaygin & Taşdeler, 219 2019) (Schillé and Kadiri, personal observation). 220

Herbivore feeding behavior is an important determinant of herbivore adaptive response to plant defenses (Karban, 2011; Weeraddana & Evenden, 2019). Yet, in our case, larvae were not allowed to move freely and leave the potted tree to find a more suitable host, or more generally escape competition. It is therefore possible that our experimental setup overestimates the effect of intraspecific competition. Whether this <sup>225</sup> finding is relevant to real world interactions remains to be tested.

We detected a negative relationship between the number of egg clutches laid by BTM female 226 moths and the subsequent growth of BTM larvae. This suggests the existence of intraspecific 227 competition for food with the same cohort. Such competition has already been reported, particularly in 228 leaf-miners (Bultman & Faeth, 1986; Faeth, 1992), which are endophagous insect herbivores whose inability 229 to move across leaves makes them particularly sensitive to the choice of oviposition sites by gravid female. 230 In our study, we prevented larvae from moving from one plant to another and noticed that some box trees 231 were completely defoliated by the end of the experiment. Although we did not record this information, it is 232 very likely that larvae first ran out of food in plants on which several egg clutches were laid. We are however 233 unable to determine whether the observed intraspecific competition in this cohort was determined by food 234 shortage, or by herbivore-induced changes in resource quality, or both. 235

## 236 5 Conclusion

Our greenhouse experiment provides evidence for negative interaction across and within BTM generation, 237 which are independent of BTM female choice for oviposition site. Such interactions may have consequences 238 on BTM population dynamics and damage on box-trees. On the one hand, the slow-growth-high-mortality 239 hypothesis states that any plant trait reducing the growth rate of herbivores can be seen as a resistance trait, 240 because slow-growing herbivores are longer exposed to their enemies (Benrey & Denno, 1997; Coley et al., 241 2006; Uesugi, 2015). It is therefore possible that a stronger top-down control can be exerted by generalist 242 enemies on BTM larvae feeding on previously defoliated hosts which could reduce damage on box-trees. On 243 the other hand, if herbivores take a longer time to complete development, they may be more damaging to 244 plants, in particular to plants with low nutritional quality as a result of compensatory feeding (Simpson 245 & Simpson, 1990; Milanovic et al., 2014). The consequences of time-lagged intraspecific competition on 246 the spread of and damage by BTM remain however to be investigated in the field. Particular efforts should 247 be dedictated to the identification of host traits controlling the performance of BTM larvae and the the 248 interaction between these traits and the higher trophic level. 249

### <sup>250</sup> 6 Acknowledgements

We thank Alex Stemmelen, and Yannick Mellerin for their help in BTM rearing and data collection. This
research was founded by the HOMED project, which received funding from the European Union's Horizon
2020 research and innovation program under grant agreement No. 771271.

### <sup>254</sup> 7 Data accessibility

Raw data as well as codes of statistic analysis are available in supplementary material and on the INRA dataverse: Castagneyrol, Bastien; van Halder, Inge; Kadiri, Yasmine; Schillé, Laura; Jactel, Hervé, 2020, "Raw data for the paper 'Host-mediated, cross-generational intraspecific competition in a multivoltine herbivore species', https://doi.org/10.15454/KMUX39, Portail Data INRAE, V1

## 259 8 References

Abdala-Roberts L, Reyes-Hernández M, Quijano-Medina T, Moreira X, Francisco M, Angulo DF, Parra-Tabla

V, Virgen A & Rojas JC (2019) Effects of amount and recurrence of leaf herbivory on the induction of direct

- <sup>262</sup> and indirect defences in wild cotton. Plant Biology 21:1063–1071.
- Agrawal A (1999) Induced responses to herbivory in wild radish: Effects on several herbivores and plant fitness. Ecology 80:1713–1723.
- <sup>265</sup> Alkan Akıncı H & Kurdoğlu O (2019) Damage Level of Cydalima perspectalis (Lepidoptera: Crambidae)
- on Naturally Growing and Ornamental Box Populations in Artvin, Turkey. Kastamonu Üniversitesi Orman
   Fakültesi Dergisi.
- Averill AL & Prokopy RJ (1987) Intraspecific Competition in the Tephritid Fruit Fly Rhagoletis Pomonella.
   Ecology 68:878–886.
- Barnes EE & Murphy SM (2018) Time-lagged intraspecific competition in temporally separated cohorts of a generalist insect. Oecologia 186:711–718.
- <sup>272</sup> Baur B, Jung J & Rusterholz H-P (2019) Defoliation of wild native box trees (Buxus sempervirens): Does box
- rust (Puccinia buxi) infection influence herbivory, survival and growth of the invasive Cydalima perspectalis? Journal of Applied Entomology 143:766–775.
- Benrey B & Denno RF (1997) The slow-growth-high-mortality hypothesis: A test using the cabbage butterfly.
  Ecology 78:987–999.
- 277 Bergamini LL & Almeida-Neto M (2015) Female Preference and Offspring Performance in the Seed Beetle
- <sup>278</sup> Gibbobruchus bergamini Manfio & Ribeiro-Costa (Coleoptera: Chrysomelidae): A Multi-Scale Comparison.
- 279 Neotropical Entomology 44:328–337.
- <sup>280</sup> Blenn B, Bandoly M, Küffner A, Otte T, Geiselhardt S, Fatouros NE & Hilker M (2012) Insect Egg Deposition
- Induces Indirect Defense and Epicuticular Wax Changes in Arabidopsis thaliana. Journal of Chemical Ecology
   38:882–892.
- Bultman TL & Faeth SH (1986) Experimental Evidence for Intraspecific Competition in a Lepidopteran Leaf
   Miner. Ecology 67:442–448.
- Burnham KP & Anderson DR (2002) Model selection and multimodel inference a practical information theoretic approach. Springer, New York.
- <sup>287</sup> Coley PD, Bateman ML & Kursar TA (2006) The effects of plant quality on caterpillar growth and defense
   <sup>288</sup> against natural enemies. Oikos 115:219–228.
- De Moraes CM, Mescher MC & Tumlinson JH (2001) Caterpillar-induced nocturnal plant volatiles repel
   conspecific females. Nature 410:577–580.
- Denno RF, McClure MS & Ott JR (1995) Interspecific Interactions in Phytophagous Insects: Competition
   Reexamined and Resurrected. Annual Review of Entomology 40:297–331.
- Faeth SH (1986) Indirect Interactions Between Temporally Separated Herbivores Mediated by the Host Plant.
   Ecology 67:479-494.
- Faeth SH (1992) Interspecific and Intraspecific Interactions Via Plant Responses to Folivory: An Experimental
   Field Test. Ecology 73:1802–1813.
- <sup>297</sup> Fatouros NE, Lucas-Barbosa D, Weldegergis BT, Pashalidou FG, Loon JJA van, Dicke M, Harvey JA, Gols
- R & Huigens ME (2012) Plant Volatiles Induced by Herbivore Egg Deposition Affect Insects of Different
   Trophic Levels. PLOS ONE 7:e43607.
- <sup>300</sup> Fox J, Weisberg S, Adler D, Bates D, Baud-Bovy G, Ellison S, Firth D, Friendly M, Gorjanc G, Graves S,
- <sup>301</sup> Heiberger R, Laboissiere R, Monette G, Murdoch D, Nilsson H, Ogle D, Ripley B, Venables W, Winsemius
- $_{302}~$  D, Zeileis A & R-Core (2016) Car: Companion to Applied Regression.
- Gripenberg S, Mayhew PJ, Parnell M & Roslin T (2010) A meta-analysis of preference-performance relationships in phytophagous insects. Ecology Letters 13:383–393.
- <sup>305</sup> Hashem MY, Ahmed AAI, Mohamed SM, Sewify GH & Khalil SH (2013) Oviposition deterrent effect of

- Spodoptera littoralis (Boisd.) larval frass to adult females of two major noctuid insect pests. Archives of 306 Phytopathology and Plant Protection 46:911–916. 307
- Hilker M & Fatouros NE (2015) Plant Responses to Insect Egg Deposition. Annual Review of Entomology 308 60:493-515.309
- Hopper KR (1999) Risk-spreading and bet-hedging in insect population biology. Annual Review of Entomology 310 44:535-560. 311
- Jones TH & Finch S (1987) The effect of a chemical deterrent, released from the frass of caterpillars of the 312 garden pebble moth, on cabbage root fly oviposition. Entomologia Experimentalis et Applicata 45:283–288. 313
- Kaplan I & Denno RF (2007) Interspecific interactions in phytophagous insects revisited: A quantitative 314 assessment of competition theory. Ecology Letters 10:977–994. 315
- Karban R (2011) The ecology and evolution of induced resistance against herbivores. Functional Ecology 316 25:339 - 347.317
- Kaygin AT & Taşdeler C (2019) Cydalima perspectalis (Walker) (Lepidoptera: Crambidae, Spilomelinae)'in 318
- Türkiye'de Coğrafi Yayılışı, Yaşam Döngüsü Ve Zararı. Bartın Orman Fakültesi Dergisi 21:833–847. 319
- Kenis M, Nacambo S, Leuthardt FLG, Domenico F di & Haye T (2013) The box tree moth, Cydalima 320 perspectalis, in Europe: Horticultural pest or environmental disaster? Aliens: The Invasive Species Bulletin:38-321 41. 322
- Leuthardt FLG & Baur B (2013) Oviposition preference and larval development of the invasive moth Cydalima 323 perspectalis on five European box-tree varieties. Journal of Applied Entomology 137:437-444. 324
- Leuthardt FLG, Glauser G & Baur B (2013) Composition of alkaloids in different box tree varieties and their 325 uptake by the box tree moth Cydalima perspectalis. Chemoecology 23:203–212. 326
- Marchand D & McNeil JN (2004) Avoidance of intraspecific competition via host modification in a grazing, 327 fruit-eating insect. Animal Behaviour 67:397–402. 328
- Martinez G, Finozzi MV, Cantero G, Soler R, Dicke M & Gonzalez A (2017) Oviposition preference but not 329
- adult feeding preference matches with offspring performance in the bronze bug Thaumastocoris peregrinus. 330
- Entomologia Experimentalis Et Applicata 163:101–111. 331
- Matošević D, Matošević D, Croatian Forest Research Institute, Cvjetno naselje 41, HR-10450 Jastrebarsko, 332
- Croatia, Bras A, INRA, UR633 Unité de Recherche de Zoologie Forestière, 2163 Avenue de la Pomme de 333
- Pin, CS 40001 ARDON45075 ORLEANS Cedex 2, France, Lacković N, Croatian Forest Research Institute. 334
- Cvjetno naselje 41, HR-10450 Jastrebarsko, Croatia, Pernek M & Croatian Forest Research Institute, Cvjetno 335 naselje 41, HR-10450 Jastrebarsko, Croatia (2017) Spatial Distribution, Genetic Diversity and Food Choice 336
- of Box Tree Moth (Cydalima perspectalis) in Croatia. South-east European forestry 8. 337
- Milanovic S, Lazarevic J, Popovic Z, Miletic Z, Kostic M, Radulovic Z, Karadzic D & Vuleta A (2014) 338
- Preference and performance of the larvae of Lymantria dispar (Lepidoptera: Lymantriidae) on three species 339
- of European oaks. European Journal of Entomology 111:371–378. 340
- Molnár BP, Tóth Z & Kárpáti Z (2017) Synthetic blend of larval frass volatiles repel oviposition in the 341 invasive box tree moth, Cydalima perspectalis. Journal of Pest Science 90:873–885. 342
- Moreira X, Abdala-Roberts L & Castagneyrol B (2018) Interactions between plant defence signalling pathways: 343
- Evidence from bioassays with insect herbivores and plant pathogens (M Rees, Ed. by). Journal of Ecology 344 106:2353-2364. 345
- Moura RR, Ribeiro PVA, Pereira BG, Quero A, Carvalho RL & Oliveira DC (2017) Food, shelter or 346 competitors? Overlapping of life stages and host plant selection in a Neotropical stink bug species. Journal
- of Plant Interactions 12:560-566. 348

347

Nykänen H & Koricheva J (2004) Damage-induced changes in woody plants and their effects on insect 349

- <sup>350</sup> herbivore performance: A meta-analysis. Oikos 104:247–268.
- Pinheiro J, Bates D, DebRoy S, Sarkar D & Team RC (2020) Nlme: Linear and Nonlinear Mixed Effects
   Models.
- <sup>353</sup> Poelman EH, Broekgaarden C, Loon JJaV & Dicke M (2008) Early season herbivore differentially affects
- plant defence responses to subsequently colonizing herbivores and their abundance in the field. Molecular
   Ecology 17:3352-3365.
- Root RB & Kareiva PM (1984) The Search for Resources by Cabbage Butterflies (Pieris Rapae): Ecological
   Consequences and Adaptive Significance of Markovian Movements in a Patchy Environment. Ecology
   65:147–165.
- 359 Sato Y, Yano S, Takabayashi J & Ohsaki N (1999) Pieris rapae</i> (Ledidoptera : Pieridae) females avoid
- oviposition on <i>Rorippa indica plants infested by conspecific larvae. Applied Entomology and Zoology 361 34:333–337.
- Shiojiri K & Takabayashi J (2003) Effects of specialist parasitoids on oviposition preference of phytophagous
   insects: Encounter-dilution effects in a tritrophic interaction. Ecological Entomology 28:573–578.
- Simpson SJ & Simpson C (1990) The Mechanisms of Nutritional Compensation by Phytophagous Insects.
   Insect-Plant Interactions (1990) : Volume II. CRC Press,
- Stam JM, Kroes A, Li Y, Gols R, Loon JJ van, Poelman EH & Dicke M (2014) Plant Interactions with
   Multiple Insect Herbivores: From Community to Genes. Annual Review of Plant Biology 65:689–713.
- <sup>368</sup> Team RC (2018) R: A language and environment for statistical computing.
- Uesugi A (2015) The slow-growth high-mortality hypothesis: Direct experimental support in a leafmining fly.
   Ecological Entomology 40:221–228.
- <sup>371</sup> Visakorpi K, Riutta T, Martínez-Bauer AE, Salminen J-P & Gripenberg S (2019) Insect community structure <sup>372</sup> covaries with host plant chemistry but is not affected by prior herbivory. Ecology 100:e02739.
- Wan H, Haye T, Kenis M, Nacambo S, Xu H, Zhang F & Li H (2014) Biology and natural enemies of *Cydalima perspectalis* in Asia: Is there biological control potential in Europe? Journal of Applied Entomology 138:715–722.
- Weeraddana CDS & Evenden ML (2019) Herbivore-induced plants do not affect oviposition but do affect fitness of subsequent herbivores on canola. Entomologia Experimentalis et Applicata 167:341–349.
- <sup>378</sup> Wise MJ & Weinberg AM (2002) Prior flea beetle herbivory affects oviposition preference and larval <sup>379</sup> performance of a potato beetle on their shared host plant. Ecological Entomology 27:115–122.
- Wratten SD, Edwards PJ & Winder L (1988) Insect herbivory in relation to dynamic changes in host plant quality. Biological Journal of the Linnean Society 35:339–350.
- <sup>382</sup> Zuur AF (2009) Mixed effects models and extensions in ecology with R. Springer, New York; London.

## **383** 9 Appendix

#### <sup>384</sup> 9.1 Supplementary figure

<sup>385</sup> Figure S1 - Semivariogram of the number of egg clutches as a function of distance among box trees.





<sup>387</sup> 9.2 Supplementary table containing raw data

Table S1 - Raw data used in the present manuscript: x and y are the position of each box tree in the green house; Treatment is the prior herbivory treatment; Clutch.number is the total number of egg clutches counted on a given box tree; N.L3 is the number of retrieved L3 larvae, L3.mean is the mean weight of a L3 larvae (g); N.chrysalids is the number of retroved chrysalids; Chrysalid.mean is the mean weight of a chrysalid.

х	У	Treatment	Clutch.number	N.L3	L3.mean	N.chrysalids	Chrysalid.mean
1	1	Attacked	22	5	0.0221740	0	NaN
2	1	Attacked	8	5	0.0183980	0	NaN
3	1	Attacked	12	5	0.0187360	1	0.1020100
4	1	Non attacked	7	5	0.0142200	6	0.1612240
5	1	Attacked	6	5	0.0146140	0	NaN
6	1	Attacked	10	5	0.0165620	2	0.1101750
7	1	Non attacked	19	5	0.0140740	6	0.1295600
8	1	Attacked	20	5	0.0110140	1	0.0961900
9	1	Attacked	4	5	0.0132300	1	0.1438500
10	1	Attacked	25	5	0.0057520	1	0.0891900
11	1	Non attacked	14	5	0.0252560	4	0.0800775
1	2	Attacked	12	5	0.0226500	2	0.2062350
2	2	Attacked	10	5	0.0162200	1	0.0524100
3	2	Attacked	9	5	0.0200760	0	NaN
4	2	Attacked	11	5	0.0211200	5	0.1465000
5	2	Non attacked	9	5	0.0160880	2	0.1455000
6	2	Non attacked	11	4	0.0332225	11	0.1486660
7	2	Attacked	5	5	0.0112560	0	NaN
8	2	Non attacked	4	5	0.0192900	10	0.1718880
- 9	2	Attacked	4	5	0.0161760	1	0 1338800
10	2	Attacked	4	5	0.010173680	1	0.1706800
11	2	Non attacked	18	5	0.0175000	1	0.1142100
1	2	Attacked	10	5	0.0104100	1	0.1142100
$\frac{1}{2}$	<u>ु</u> २	Non attacked	14	5	0.0103000	4	0.1030525
$\frac{2}{2}$	- J - 2	Non attacked	5	5	0.0193180	8	0.1342200
	່ <u>ບ</u>	Attacked	5	5	0.0277000	10	0.1413480
-4-5	ა ე	Attacked	5	5	0.0139420	<u> </u>	0.1360100
-0-6-	ა ე	Non attacked	0	5	0.0121100	1	0.1319100
	<u>ა</u>	Non attacked	0	0 E	0.0353020	9	0.1404700
	<u>ა</u>	Non attacked	1	5	0.0197740	11	0.1815320
8	<u>ა</u>	Non attacked	2	5	0.0334440	9	0.2104500
10	<u>ა</u>	Non attacked	2	5	0.0144920	2	0.1980650
10	3	Non attacked	8	5	0.0188620	7	0.1529100
11	3	Attacked	11	5	0.0101960	1	0.0836300
	4	Attacked	2	5	0.0111600	2	0.1224050
2	4	Non attacked	21	5	0.0127880	2	0.1022250
3	4	Attacked	11	5	0.0157420	0	NaN
4	4	Attacked	11	5	0.0158140	4	0.1557575
5	4	Non attacked	2	5	0.0171240	1	0.1575800
6	4	Attacked	2	5	0.0238660	2	0.1728600
7	4	Non attacked	11	5	0.0151460	0	NaN
8	4	Attacked	5	5	0.0187260	2	0.1527050
9	4	Non attacked	5	5	0.0256880	5	0.1158560
10	4	Non attacked	5	5	0.0171440	3	0.1927033
11	4	Attacked	8	5	0.0181900	0	NaN
1	5	Non attacked	4	5	0.0185720	3	0.1905400
2	5	Non attacked	3	5	0.0363840	20	0.1862420
3	5	Non attacked	4	5	0.0224060	8	0.1504320
4	5	Non attacked	4	5	0.0291180	9	0.1339320
5	5	Attacked	1	5	0.0201820	1	0.1914500
6	5	Non attacked	2	5	0.0313100	2	0.1725900
7	5	Attacked	1	5	0.0148320	3	0.1752800
8	5	Attacked	2	5	0.0176160	1	0.0853400
9	5	Non attacked	15	5	0.0130960	1	0.1001600
10	5	Attacked	10	134	0.0171925	1	0.1453700
11	5	Non attacked	7	5	0.0128840	2	0.1279200
1	6	Attacked	4	5	0.0158160	2	0.1774000
2	6	Non attacked	4	5	0.0225620	11	0.1544600
	-	37	i .				

```
library(tidyverse)
library(knitr)
library(kableExtra)
library(nlme)
library(car)
library(citr)
library(here)
wd = here()
my.ggplot <- function(){</pre>
  theme_bw() + theme(legend.key = element_blank())
}
knitr::opts_chunk$set(echo= F, warning= F, message = F, results = "hide",
                      fig.width=4, fig.height=4, dpi= 100, fig.pos = "H",
                      fig.path = paste0(wd, '/Figures/'))
d = read.csv(paste0(wd, '/Data/BTM_data.csv'), header = T)
str(d)
d =
  d %>% mutate(L3.mean = Weight.L3 / N.L3) %>%
  mutate(Treatment = factor(Treatment, levels = c("Non attacked", "Attacked")))
knitr::include_graphics(paste0(wd, '/Figures/Figure 1.png'))
d %>%
  ggplot(aes(x, y, shape = Treatment, fill = log1p(Clutch.number))) +
  # geom_rect(aes(xmin = 1.5, ymin = 1.5, xmax = 10.5, ymax = 10.5), fill = 'grey90') +
  geom_point(size = 6) +
  scale_shape_manual(values = c(21, 24)) +
  my.ggplot() +
  labs(x = "", y = "") +
  scale_fill_gradientn(colours = c('white', 'grey', 'black'), values = c(0,0.6,1), name = 'No. egg clut
d %>%
  select(Treatment, Clutch.number, N.L3, N.chysalids) %>%
  gather(Resp, Val, 2:4) %>%
  group_by(Treatment, Resp) %>%
  summarise(N = sum(Val), M = round(mean(Val),2), SD = round(sd(Val),2)) %>%
  mutate(Val = paste(M, ' (', SD, ')', ', n = ', N, sep = '')) %>%
  select(- N, - M, - SD) %>%
  mutate(Resp = factor(Resp, levels = c('Clutch.number', 'N.L3', 'N.chysalids'), labels = c('Egg clutch
  spread(Treatment, Val) %>%
  kable(col.names = c('Response variable', 'Control', 'Herbivory treatment'),
        caption = "Repartition of egg clutches, larvae and chrysalis across box trees with or without p
d.center = droplevels(d[d$x > 1 & d$x < 11 & d$y > 1 & d$y < 11,])
m0 = gls(log1p(Clutch.number) ~ Treatment, data = d, na.action = "na.omit")
m1a = gls(log1p(Clutch.number) ~ Treatment, correlation = corExp(form = ~x + y, nugget = TRUE), data = 
m1b = gls(log1p(Clutch.number) ~ Treatment, correlation = corGaus(form = ~x + y, nugget = TRUE), data =
mic = gls(log1p(Clutch.number) ~ Treatment, correlation = corSpher(form = ~x + y, nugget = TRUE), data =
```

```
m1d = gls(log1p(Clutch.number) ~ Treatment, correlation = corLin(form = ~x + y, nugget = TRUE), data = .
m1e = gls(log1p(Clutch.number) ~ Treatment, correlation = corRatio(form = ~x + y, nugget = TRUE), data =
Table_AIC = AIC(m0, m1a, m1b, m1c, m1d, m1e)
Table_AIC$AIC = round(Table_AIC$AIC,1)
Table_AIC$Delta = Table_AIC$AIC - min(Table_AIC$AIC)
# Table_AIC %>% kable() %>% kableExtra::kable_styling()
# plot_resid = function(model) {
#
   df = data.frame(Residuals = residuals(model),
#
                   Fitted = fitted(model))
#
#
   A = df %>% ggplot(aes(Fitted, Residuals)) +
    my.ggplot() +
#
#
    qeom_point() +
#
     geom_smooth(method = "lm", se = F) +
#
     geom_hline(yintercept = 0)
#
#
   B = df %>% qqplot(aes(Residuals)) + my.qqplot() + qeom_histogram()
#
#
    cowplot::plot_grid(A, B)
# }
#
# plot_resid(m1e)
# Alternative approach fitting *x* and y*y position in the greenhouse together with treatment and `N.a
#
# m = MASS::glm.nb(Clutch.number ~Treatment * N.attacked, data = d.center)
# Anova(m)
# #
# # var.resid = Variogram(residuals(m), dist(d.center %>% select(x, y)))
# # var.resid %>%
    qqplot(aes(dist, varioq)) +
# #
# # my.ggplot() +
# # # geom_point(size = 3) +
# # geom_smooth(se = F) +
     labs(x = "Distance (m)", y = "Semivariogram")
# #
plot.resid = function(m){
  df = data.frame(f = fitted(m), r = residuals(m))
  A = df \%
    ggplot(aes(f, r)) + my.ggplot() + geom_point() + geom_smooth(method = "lm", se = F) + geom_hline(yi)
  B = df \%
    ggplot(aes(r)) + my.ggplot() + geom_histogram()
  cowplot::plot_grid(A,B)
}
m_larvae = lm(log(L3.mean) ~ Clutch.number * Treatment , d)
# plot.resid(m_larvae)
d =
  d %>%
  mutate(Chrysalid.mean = Weight.chrysalids / N.weighted.chysalids)
```

```
m_chrys = lm(log(Chrysalid.mean) ~ Clutch.number * Treatment , d)
# plot.resid(m2)
Table_AIC %>%
  mutate(Model = paste('Model', 1:6),
         'Correlation structure' = c('None', 'Exponential', 'Gaussian', 'Spherical', 'Linear', 'Rationa
  select(Model, `Correlation structure`, everything()) %>%
  kable(caption = "Summary of AIC of GLS models testing the effect of prior herbivory on the number of
Anova(m1e)
Fig_3A =
  d %>%
  ggplot(aes(Treatment, Clutch.number)) +
  my.ggplot() +
  geom_point(position = position_jitter(0.1), colour = "grey80") +
  stat_summary(size = 0.6) +
  labs(x = "Prior herbivory", y = "Number of egg clutches") +
  annotate(geom = 'text', x = 1:2, y = 26, label = paste("n =", c(61, 60)))
res_larvae = Anova(m_larvae)
n = expand.grid(Treatment = levels(d$Treatment), Clutch.number = seq(0, 25))
p = predict(update(m_larvae, ~.- Clutch.number:Treatment), newdata = n)
nFit = 1000 * exp(p)
Fig 3B =
  d %>%
  ggplot(aes(Clutch.number, 1000 * L3.mean, colour = Treatment)) +
  my.ggplot() +
  geom_point(size = 2) +
  scale_colour_manual(values = c("grey25", "grey70")) +
  labs(x = "Number of egg clutches",
       y = expression("Mean larval weight" %+-% "SE (mg)")) +
  geom_line(data = n, aes(Clutch.number, Fit), size = 1.5) +
  theme(legend.position = c(0.8, 0.85))
r = with(d, cor.test(Chrysalid.mean, L3.mean))
cowplot::plot_grid(
  Fig_3A + labs(title = 'Eggs'),
  Fig_3B + labs(title = 'Larvae'),
 ncol = 2.
 labels = c('A', 'B')
)
X = c("Number of egg clutches", "Herbivory", "Eggs x Herbivory")
Fval = function(x) {round(x[,3], 2)}
Pval = function(x) {ifelse(x[,4] < 0.001, '< 0.001', round(x[,4], 3))}</pre>
df = rep("1, 117", 3)
b = round(summary(m_larvae)$coefficients[-1,1], 2)
b_se = round(summary(m_larvae)$coefficients[-1,2], 2)
Estimate = paste(b, ' (', b_se, ')', sep = '')
data.frame(#Response = c('Larvae', '', ''),
           Predictor = X,
           df = df,
           Estimate = Estimate,
           `F-value` = Fval(res_larvae)[-4],
```

```
`P-value` = Pval(res_larvae)[-4]) %>%
kable(caption = "Summary of models testing the effect of prior herbivory and initial egg clutch densi
col.names = c("Predictor", "df", "Estimate (SE)", "F-value", "P-value"),escape = T, digit = 2) "
m0 = gls(log1p(Clutch.number) ~ x + y, data = d, na.action = "na.omit")
variog0 <- Variogram(m0, form = ~x + y, resType = "pearson", nugget = T)
variog0 %>%
ggplot(aes(0.4*dist, variog)) +
my.ggplot() +
geom_point(size = 3) +
geom_smooth(se = F) +
labs(x = "Distance (m)", y = "Semivariogram") +
xlim(0, 4)
d %>%
rename(N.chrysalids = N.chysalids) %>%
select(x, y, Treatment, Clutch.number, N.L3, L3.mean, N.chrysalids, Chrysalid.mean) %>% kable() %>% kable() %>% kable() %>%
```