

1 | Sex makes them sleepy: ~~change in~~ host reproductive status induces diapause in  
2 | ~~parasitoids~~ parasitoid population experiencing harsh winters  
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6 | **Tougeron K.<sup>1,2</sup>, Brodeur J.<sup>2</sup>, van Baaren J.<sup>1</sup>, Renault D.<sup>1,3</sup> & Le Lann C.<sup>1</sup>**

7 | <sup>1</sup> Univ Rennes, CNRS, ECOBIO (Ecosystèmes, biodiversité, évolution) - UMR 6553, 263 Avenue du  
8 | Général Leclerc, F-35000 Rennes, France.

9 | <sup>2</sup> Institut de Recherche en Biologie Végétale, Département de Sciences Biologiques, Université de  
10 | Montréal, 4101 rue Sherbrooke Est, Montréal, QC, Canada, H1X 2B2.

11 | <sup>3</sup> Institut Universitaire de France, 1 rue Descartes, 75231 Paris Cedex 05, France

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14 | **Corresponding author:** [tougeron.kevin@gmail.com](mailto:tougeron.kevin@gmail.com)

15 | Current address: The University of Wisconsin – La Crosse, Department of Biology, La Crosse,  
16 | Wisconsin, United States of America, 1725 State street, 54601

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21 **Abstract**

22       When organisms coevolve, any change in one species can induce phenotypic changes in  
23 traits and ecology of the other species. The role such interactions play in ecosystems is central,  
24 but their mechanistic bases remain underexplored. Upper trophic level species have to  
25 synchronize their life-cycle to both abiotic conditions and to lower trophic level species'  
26 phenology and phenotypic variations. We tested the effect of host seasonal strategy on  
27 parasitoid diapause induction by using a holocyclic clone of the pea aphid *Acyrtosiphon pisum*  
28 producing ~~two morphs with either~~ asexual ~~(and sexual morphs that are~~ viviparous females) ~~or~~  
29 ~~sexual~~ ~~(i.e. laying embryos) and~~ oviparous females) ~~reproduction (laying eggs), respectively,~~  
30 the latter being only present at the end of the growing season. *Aphidius ervi* parasitoids from  
31 ~~populations of~~ contrasted climatic origin (harsh vs. mild winter areas) were allowed to parasitize  
32 each morph ~~in a split-brood design~~ and developing parasitoids were next reared under either  
33 fall-like or summer-like temperature-photoperiod conditions. We next examined aspects of the  
34 host physiological state by comparing the relative proportion of forty-seven metabolites and  
35 lipid reserves in both morphs: ~~produced under the same conditions~~. We found that oviparous  
36 morphs are cues *per se* for diapause induction; parasitoids entered diapause at higher levels  
37 when developing in oviparous hosts ( $19.4 \pm 3.0\%$ ) than in viviparous ones ( $3.6 \pm 1.3\%$ ), under  
38 summer-like conditions (i.e., when oviparous aphids appear in the fields). This pattern was only  
39 observed in parasitoids from the harsh winter area since low diapause levels were observed in  
40 the other population ~~dependent~~, suggesting local adaptations to overwintering  
41 cues. Metabolomics analyses show parasitoids' response to be mainly influenced by the host's  
42 physiology, with higher proportion of polyols and sugars, and more fat reserves being found in  
43 oviparous morphs. Host quality thus varies across the seasons: ~~and represents one of the~~  
44 multiple environmental parameters affecting parasitoid diapause. Our results underline strong  
45 coevolutionary processes between hosts and parasitoids in their area of origin, likely leading to  
46 phenological synchronization, and we point out ~~their~~ the importance of such bottom-up effects  
47 for trait-expression, and for the provision of ecosystem ~~services~~ services such as biological  
48 control in the context of climate change.

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50 **Key-words**

51       Coevolution; Phenotypic plasticity; Phenology; Host-parasite synchronization;  
52 Environmental cue; Metabolomics

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## Introduction

Interacting individuals from two biological entities can adjust their phenotypes in response to cues from each other, even when these cues vary across time (Agrawal 2001). Beneficial or antagonistic interactions, from mutualism to parasitism, predation and competition may lead to adaptive phenotypic responses. When interactions persist over generations, coevolution can occur and species adapt to the interacting species' life history traits, phenology and ecology (Agrawal 2001, Ellers et al. 2012). Interaction norms (Thompson 1988) arise from ecological responses of interacting organisms in varying environments, as any phenotypic change occurring in one "partner" species can cascade to the other species' phenotype (Fordyce 2006, Hughes 2012). Cues produced by one interacting species may indirectly inform the other species of environmental changes. For example, plant senescence in fall can inform herbivorous insects of upcoming detrimental winter conditions and induces phenotypic changes (e.g. diapause induction) or migration behaviour (Archetti et al. 2009).

Parasitoids are excellent models to study phenotypic expression in interacting species because they are strongly influenced during immature stages by changes in nutritional and physiological quality of their host (Godfray 1994). Diapause is an important ecological process in insects allowing them to survive recurrent unfavorable environmental conditions (Tauber et al. 1986). For parasitoids, diapause also contributes to maintain synchronization with their host's seasonal reproductive-cycle; it is induced before suitable hosts vanish from the environment (Lalonde 2004). As in most insects, diapause in parasitoids is mainly induced by abiotic cues perceived either by the generation that will enter diapause, or by the maternal generation (Tauber et al. 1986). A few studies also reported that diapause in parasitoids can be triggered by the onset of host diapause (Polgár and Hardie 2000, Gerling et al. 2009), or through intraspecific competition for hosts (Tougeron et al. 2017a). However, whether the phenotype of a non-diapausing host can influence parasitoid diapause remains poorly studied.

Aphids are hosts for Aphidiinae parasitoids and can have very complex cycles showing seasonal alternation between morphs with asexual and sexual reproduction (~~Dixon 1985~~)(Dixon 1985). Asexual females reproduce parthenogenetically and lay live offspring (i.e. viviparity) whereas ~~sexual~~sexually reproducing females produce eggs (i.e. oviparity) after mating with males. Sexual aphid morphs are present at higher proportions in harsh than in mild winter climates (~~Dedryver et al. 2001~~)(Dedryver et al. 2001), and they represent the last hosts available for aphid parasitoids before winter as they produce overwintering eggs in fall (~~Leather 1992~~)(Leather 1992). Consequently, sexual morphs have been suggested to promote diapause in parasitoids, indicating a host physiological effect (~~Polgár et al. 1991, 1995, Christiansen-Weniger and Hardie 1997~~)(Polgár et al. 1991, 1995, Christiansen-Weniger and Hardie 1997). No mechanistic understanding of this phenomenon has been proposed and the effects of the host morph have not been detangled from confounding factors such as host genotype and geographic origin, host size, abiotic conditions, or the season at which hosts are sampled in the fields. Hosts and parasitoids ~~share common evolutionary history~~have coevolved over long periods of time, they respond to similar seasonal cues and the physiological syndrome associated with overwintering is highly conserved among insects (~~Tauber et al. 1986, Denlinger 2002~~)(Tauber et al. 1986, Denlinger 2002). As a result, the related physiological state of the host may represent a reliable signal of upcoming seasonal changes for parasitoids.

~~Hormones, fats, carbohydrates and other types of metabolites are involved in the control of overwintering and diapause expression in insects (Chippendale 1977, Christiansen-Weniger and~~

102 ~~Hardie 1999, Denlinger 2002, Sinclair and Marshall 2018). In aphid parasitoids, metabolomic~~  
103 ~~and proteomic profiles differ between diapausing and non-diapausing individuals, with higher~~  
104 ~~amounts of sugars, polyols and heat shock proteins being found in diapausing parasitoids~~  
105 ~~(Colinet et al. 2012). In aphids, morphs differ in morphology and physiology; oviparous females~~  
106 ~~accumulate reserves to produce energetically costly diapausing eggs (Le Trionnaire et al. 2008)~~  
107 ~~with cryoprotectant compounds such as mannitol and glycerol (Sömme 1969), whereas~~  
108 ~~viviparous females metabolize energetic resources rapidly to produce embryos. Aphids'~~  
109 ~~triglyceride reserves change quantitatively and qualitatively across the seasons with alternating~~  
110 ~~morphs (Greenway et al. 1974). Immature parasitoids are known to consume sugars and lipids~~  
111 ~~from their hosts (Jervis et al. 2008) and are therefore influenced by host reserves for their~~  
112 ~~growth and development.~~

113 ~~We questioned the extent to which oviparous and viviparous morphs of a single clone of the~~  
114 ~~pea aphid *Acyrtosiphon pisum* (Harris) (Hemiptera: Aphididae) influences winter diapause~~  
115 ~~expression in the parasitoid *Aphidius ervi* Haliday (Hymenoptera: Braconidae) under summer~~  
116 ~~and fall conditions. Under laboratory conditions Hormones, fats, carbohydrates and other types~~  
117 ~~of metabolites are involved in the regulation of overwintering and diapause expression in insects~~  
118 ~~(Chippendale 1977, Christiansen-Weniger and Hardie 1999, Denlinger 2002, Sinclair and~~  
119 ~~Marshall 2018). In aphid parasitoids, metabolomic and proteomic profiles differ between~~  
120 ~~diapausing and non-diapausing individuals, with higher amounts of sugars, polyols and heat~~  
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131 ~~expression in the parasitoid *Aphidius ervi* Haliday (Hymenoptera: Braconidae) under summer~~  
132 ~~and fall conditions. Under laboratory conditions and using a split-brood design, we compared~~  
133 ~~the response to two aphid morphs of two populations of parasitoids from mild (France) and~~  
134 ~~harsh (Canada) winter areas that differed in their level of diapause expression (Tougeron et al.~~  
135 ~~2018)(Tougeron et al. 2018). In *Aphidius* species, winter diapause is initiated at the prepupal~~  
136 ~~stage within the aphid mummy (i.e. dead aphid containing a developing parasitoid) following~~  
137 ~~stimuli perceived by the mother or early developmental stages (Brodeur and McNeil 1989,~~  
138 ~~Tougeron et al. 2017b)(Brodeur and McNeil 1989, Tougeron et al. 2017b). We hypothesized~~  
139 ~~that parasitoids of both populations developing in oviparous hosts enter diapause at higher~~  
140 ~~proportions than those developing in viviparous hosts, independently of photoperiod and~~  
141 ~~temperature. We predicted this pattern to originate from differences in aphids' physiological~~  
142 ~~contents. We thus performed physiological analyzes to measure lipid content and quantify aphid~~  
143 ~~morphs metabolites. We also hypothesized parasitoids from mild winter area to be less~~  
144 ~~responsive to diapause inducing cues from the host and the environment, because parasitoid~~  
145 ~~populations should be adapted to climatic conditions and relative occurrence of sexual hosts in~~  
146 ~~their respective area of origin.~~

147

148 **Material and Methods**

149 *Biological materials*

150 Two populations of the parasitoid *A. ervi* were collected in 2015 at the mummy stage in pea  
151 fields from two contrasted climatic origins: near Montréal, QC, Canada (45.584°N, 73.243°W;  
152 harsh winter area) and near Rennes, France (48.113°N, 1.674°W; mild winter area). ~~Parasitoids~~  
153 ~~were then reared under controlled conditions using a cyclically parthenogenetic clone (clone F2-~~  
154 ~~X9-47) of the pea aphid *A. pisum* provided by INRA Le Rheu, France, and known to produce~~  
155 ~~both oviparous and viviparous aphid morphs (Jaquiéry et al. 2014). One population per~~  
156 ~~geographic origins was used as high gene flow has been reported in *A. ervi* populations which~~  
157 ~~therefore present little genetic differentiation (Hufbauer et al. 2004). Even if gene flow was~~  
158 ~~weak, we would expect higher differences between Canadian and French populations than~~  
159 ~~among populations of a same location. Parasitoids were then reared under controlled conditions~~  
160 ~~using a cyclically parthenogenetic clone (clone F2-X9-47) of the pea aphid *A. pisum* provided~~  
161 ~~by INRA Le Rheu, France, and known to produce both oviparous and viviparous aphid morphs~~  
162 ~~(Jaquiéry et al. 2014). The symbiotic load of the aphid clone we used was not assessed, but~~  
163 ~~symbionts present in the grandparent generation from which our clone comes from had been~~  
164 ~~identified. Half of the grandparent was associated with *Serratia symbiotica*, the other half had~~  
165 ~~no secondary endosymbionts (J. Jaquiéry pers. comm.), it is thus likely that our clone was~~  
166 ~~inhabited by *S. symbiotica*. All insects were maintained on fava bean *Vicia faba* (Fabaceae) at~~  
167 20 °C, 70% relative humidity (RH) and 16:8 h Light:Dark (L:D) photoregime.

168  
169 *Production of sexual and asexual hosts*

170 Three aphid morphs were used in the experiments; oviparous females (O), viviparous  
171 females (V) and a control treatment for viviparous females (C), as detailed below.

172 Three parthenogenetic *A. pisum* adult females from the aphid culture were put on bean plants  
173 (N=15) and allowed to lay larvae during four days at 20 °C, 70% RH, 16:8 h ~~L:D~~L:D. Females  
174 were then removed and infested plants were put in a growing chamber at 17 °C, 70% RH, 12:12  
175 h (L:D), and under 36W, IRC 85, 6500 K day-light type fluorescent tubes to induce the  
176 production of sexual aphids ~~(Le Trionnaire et al. 2009)~~(Le Trionnaire et al. 2009). At each  
177 generation, plants were renewed, and less than five aphids were maintained per plant to prevent  
178 formation of ~~alate winged~~ individuals due to overcrowding ~~(Hardie 1980)~~(Hardie 1980). As  
179 embryos directly detect photoperiodic cue through the cuticle of the grand-mother ~~(Le~~  
180 ~~Trionnaire et al. 2008)~~(Le Trionnaire et al. 2008), the first sexual aphids: males (~20%) and  
181 oviparous females (30 to 60%) were formed, along with asexual aphids (20 to 50%): sexuparous  
182 (a particular type of parthenogenetic females producing sexual morphs) and viviparous aphids  
183 (parthenogenetic females producing only parthenogenetic morphs), after three generations under  
184 these conditions. As sexuparous and viviparous aphids cannot be distinguished  
185 morphologically, they were indistinctly considered as the “viviparous female” treatment.  
186 However, a control group of viviparous parthenogenetic females (C) was produced by rearing  
187 aphids under non-sexual-inductive conditions (20 °C, 70% RH, 16:8 h L:D). This treatment  
188 controls for potential stress effects of the sexual-inductive conditions on the aphid, and allows to  
189 solely measuring the response of viviparous aphids as sexuparous are not produced under this  
190 condition ~~(Dixon 1985)~~(Dixon 1985). Oviparous aphid morphs were differentiated from  
191 viviparous ones under a stereo microscope (x10) by observing the morphology of their legs:

192 oviparous female aphids have rhinaria on the tibia, and have a femur of the same width as the  
193 tibia, and viviparous females have a wider tibia than the femur without rhinaria (~~Lamb and~~  
194 ~~Pointing 1972, Hüllé et al. 2006~~)(Lamb and Pointing 1972, Hüllé et al. 2006). Aphid males  
195 were not included in our analyses since *A. ervi* does not parasitize them, probably because they  
196 are too small and have lower energetic reserves than female morphs (Tougeron et al.,  
197 unpublished data).

#### 198 199 *Diapause induction*

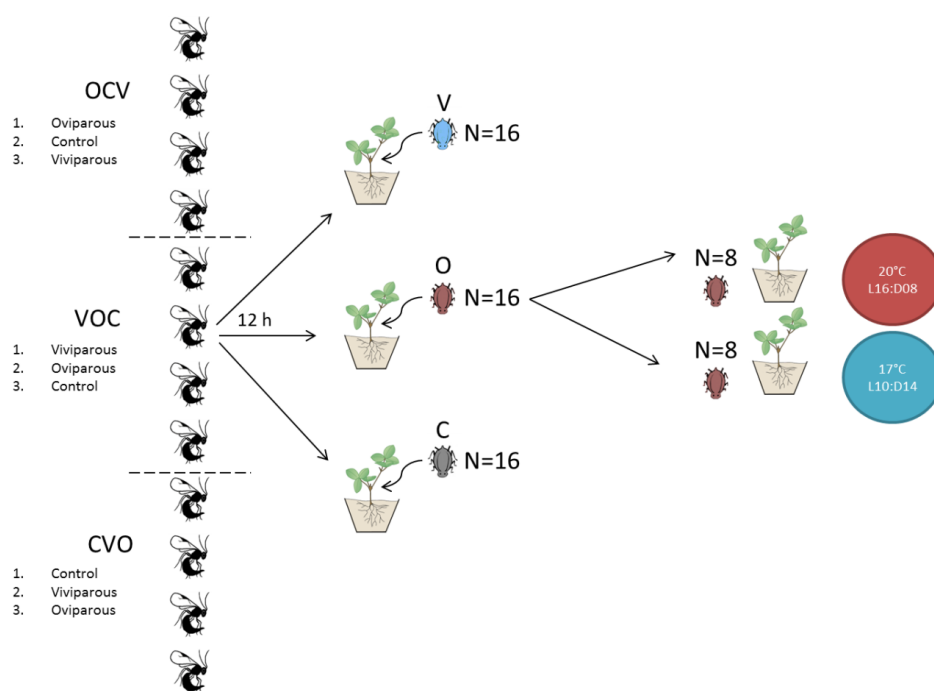
200 Aphid mummies from the colonies were isolated in a small gelatin capsule until parasitoid  
201 emergence. Newly emerged parasitoids were put in a 5 cm plastic tube for mating (5 females  
202 with 2 males) for 24 h, and were fed with a 70-% diluted honey solution. Maternal genotype,  
203 egg-laying order in different aphid morphs, in addition to parasitoids' age or host preference  
204 may affect diapause induction (~~Brodeur and McNeil 1989~~)(Brodeur and McNeil 1989). To  
205 consider these potential effects, twelve *A. ervi* females were individually allowed to parasitize  
206 16 adult aphids of the same age and size within the same cohort and of each of the three morph  
207 types (oviparous female, viviparous female, control viviparous females produced under non-  
208 sexual-inductive conditions, N=48 aphids offered for parasitism per female wasp) for 12 h over  
209 three consecutive days, by alternating the order of presentation of aphid morphs among females.  
210 Parasitoids rested at night, with an access to diluted honey. Aphids were introduced in a plastic  
211 tube (10 x 3 cm) and were given a few minutes to settle on a bean cut plant, after which a  
212 parasitoid was introduced into the tube. Four parasitoid females were first individually put in  
213 presence of oviparous aphids, then moved to a second tube with control viviparous aphids and  
214 next moved to a third tube containing viviparous aphids (OCV). Four other females were first  
215 offered viviparous aphids (VOC), and the last four females were first offered control viviparous  
216 aphids (CVO) (Fig. 1).

217 After each oviposition period, the 16 potentially parasitized aphids of each morph type were  
218 transferred by group of 8 on two bean plants. Plants were next enclosed into micro-perforated  
219 plastic bags and placed at either 20 °C, 16:8 h (L:D) (summer-like conditions not inducing  
220 diapause in *A. ervi*) or 17 °C, 10:14 h (L:D) (autumn-like conditions inducing diapause)  
221 (~~Tougeron et al. 2017b~~)(Tougeron et al. 2017b). When the plants began to wilt, aphids were  
222 transferred to another plant with a small paintbrush. Mummification was checked daily, and  
223 newly-formed mummies were placed individually into gelatin capsules, and remained under  
224 their respective temperature and photoperiod treatments until adult emergence. Mummies from  
225 which no parasitoid had emerged 15 days after mummification were dissected, and the content  
226 was recorded as dead parasitoids or diapausing individuals (golden-yellow prepupae, ~~Tougeron~~  
227 ~~et al. 2017b~~)(Tougeron et al. 2017b). This experiment was repeated twice per parasitoid  
228 population; diapause levels were thus calculated among the offspring of 24 females for each  
229 treatment. Patterns were consistent in each of the repeated experiments. Our split-brood family  
230 design also allowed comparing reaction norms (RN) of diapause levels in the offspring of each  
231 parasitoid female from each population, both within morphs at different abiotic conditions, and  
232 within abiotic conditions among morphs. We have excluded "control" morphs from the RN  
233 analysis as their effect on diapause induction did not differ from viviparous morphs.

234 The aphid morph (individual differences within a population due to developmental plasticity)  
235 and the aphid clone (differences in reproduction modes genetically determined between

236 populations) may both influence parasitoid diapause. To consider this aspect, we compared the  
 237 incidence of diapause when parasitoids developed in the cyclically parthenogenetic clone  
 238 (holocyclic, i.e., alternating between sexual and asexual morphs) described above and in an  
 239 obligate parthenogenetic clone, producing only viviparous females (anholocyclic clone F2-X9-  
 240 19; Jaquiéry et al. 2014)Jaquiéry et al. 2014). To achieve this goal, five *A. ervi* females were  
 241 individually allowed to sequentially parasitize 35 viviparous aphids of each clone during 12 h.  
 242 Parasitized hosts were next placed at 17 °C 10:14 h (L:D), and diapause induction was  
 243 measured as described above. We excluded any clone effect because diapause incidence was  
 244 similar for parasitoids developing in viviparous aphids of either the holocyclic ( $59.9 \pm 10.1\%$ ,  
 245  $n=132$  mummies) or the anholocyclic ( $66.0 \pm 7.7\%$ ,  $n=112$  mummies) clone (GLM,  $p=0.97$ ).  
 246 The cyclically parthenogenetic clone was thus used for the experiments.

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249

250 Figure 1: Experimental design for diapause induction in the parasitoid *Aphidius ervi*. Twelve parasitoid  
 251 females were individually allowed to parasitize 16 *Acyrtosiphon pisum* from each of the three host  
 252 morphs for 12 h: oviparous (O), viviparous (V) and viviparous control (C). First contact (parasitism  
 253 sequence) with an aphid was alternated between the three morphs (OCV, VOC, CVO). Following  
 254 parasitism, the aphid cohort was split in two and individuals were reared under a diapause-inductive  
 255 condition (17 °C 10:14 h L:D) or a non-diapause-inductive condition (20 °C 16:8 h L:D). This protocol  
 256 was repeated twice for parasitoid populations originating from mild or harsh winter.

257

258 *Metabolomic analyses and lipid reserves*

259 | ~~Non~~As sexual morphs could only be produced at 17°C, we compared non-parasitized  
 260 | apterous adult aphids of viviparous and oviparous females of the same age (between 24 and 48 h  
 261 | after imago molt), produced under the same conditions used for the diapause experiment (at 17  
 262 | °C, 12:12 h (L:D)). Samples were kept at -20 °C for metabolomic and lipid analyses.  
 263 | ~~Samples~~They were dried out at -60 °C for 2 days in a freeze-dryer and their dry mass measured  
 264 | using a Mettler-Toledo precision ~~balance~~scale (accurate to 0.001 mg). Viviparous aphids' dry  
 265 | mass ranged from 0.280 mg to 0.742 mg, and oviparous aphids' dry mass ranged from 0.358 mg  
 266 | to 0.739 mg.

267 | For metabolic analyses, 18 aphids of each morph (viviparous and oviparous females) were  
 268 | used. Nine replicates were analyzed for each morph condition, each consisting of a pool of two  
 269 | aphid females. The samples were put in 600 µL of ~~chloroform~~chloroform-methanol (1:2)  
 270 | solution and ~~homogenized~~homogenized using a tungsten-bead beating apparatus at 30 Hz for  
 271 | 1.5 min. Then, 400 µL of ultrapure water was added to each tube and samples were centrifuged  
 272 | at 4 °C, 4,000 g for 5 min. Finally, 90 µL of the upper aqueous phase containing metabolites  
 273 | were transferred to chromatographic vials. Injection order of the samples was randomized prior  
 274 | mass spectrometry detection. Metabolomic fingerprinting process was performed following the  
 275 | protocol of ~~Khodayari et al. (2013)~~Khodayari et al. (2013). Chromatograms were analyzed  
 276 | using XCalibur software (Thermo Fischer Scientific, Waltham, MA, USA). We accurately  
 277 | quantified 47 metabolites: 14 amino acids, 11 sugars / sugar phosphates, 8 organic acids, 7  
 278 | polyols, 4 other metabolites and 3 amines (Table 1). ~~Details of metabolite amounts measured~~  
 279 | ~~from each morph are provided in Figure S1.~~

280 | Lipid contents were measured using 52 oviparous females and 23 viviparous females. Each  
 281 | dry aphid was left for two weeks in a microtube containing 1 mL of chloroform-methanol  
 282 | solution (2:1) to extract lipids (~~Terblanche et al. 2004~~Terblanche et al. 2004). Aphids were  
 283 | then rinsed with the same solution, and placed back in the freeze-dryer for 24 h to eliminate the  
 284 | residues of the extracting solution and next weighted again to measure fat content (= fat mass  
 285 | (mg) / lean dry mass (mg), Colinet et al. 2007).

Code de champ modifié

286 |

287 | Table 1: Metabolites detected in each of the two morphs (viviparous and  
 288 | oviparous females) of the pea aphid, *Acyrtosiphon pisum*. Each metabolite has  
 289 | been found in each morph. Abbreviations used on Figure 43 are in brackets.

Amino acids	Organic acids
Alanine (Ala)	Citric acid (Cit_Ac)
Aspartic acid (Asp_Ac)	Galacturonic acid (Gal_Ac)
Citrulline (Citr)	Glyceric acid (Glyc_Ac)
Glutamic acid (Glu)	Lactic acid (Lact_Ac)
Glycine (Gly)	Malic acid (Mal_Ac)
Isoleucine (Ile)	Phosphoric acid (Phos_Ac)
Leucine (Leu)	Pipecolic acid (Pipe_Ac)
Lysine (Lys)	Quinic acid (Quin_Ac)
Ornithine (Orn)	
Proline (Pro)	<b>Sugars and sugar phosphates</b>
Serine (Ser)	Arabinose
Valine (Val)	Fructose
Threonine (Thr)	Fructose-6-phosphate (F6P)
Phenylalanine (Phe)	Galactose
	Glucose
<b>Polyols</b>	Glucose-6-phosphate (G6P)
Adonitol	Maltose
Arabitol	Mannose
Galacticol	Ribose
Glycerol	Saccharose



Inositol	Trehalose
Mannitol	<b>Other metabolites</b>
Xylitol	Gluconolactone (GNL)
<b>Amines</b>	Gamma aminobutyric acid (GABA)
Cadaverine (Cad)	Glycerol-3-phosphate (Gly3P)
Triethanolamine (TEA)	Dopamine (Dop)
Putrescine (Put)	

290

## 291 *Statistical analyses*

292 Generalized linear mixed-effects models (GLMM) with binomial distributions were fit to the  
 293 data using the *lme4* package. The response variable was the proportion of diapausing  
 294 parasitoids; the origin of the parasitoid population (Canada vs. France), the host morph (three  
 295 modalities, O, V, C), the temperature/photoperiod conditions (17°C 10:14h vs 20°C 16:8h), and  
 296 their interaction, were considered as fixed factors; the identity of each parasitoid female and the  
 297 egg-laying (parasitism) order were considered as random effect factors in the models. As  
 298 diapause incidence differed between parasitoid populations (GLMM,  $\chi^2=216$ ,  $df=1$ ,  $p<0.001$ ),  
 299 data from both populations were analyzed separately using similar GLMMs. Significance of  
 300 each term in the model was analyzed using the package *car*.

301 For metabolite data, concentrations of the compounds were first log-transformed. Then, a  
 302 Principal Component Analysis (PCA) was performed to detect which metabolites (expressed in  
 303  $nmol.mg^{-1}$ ) differed the most between host morphs. Log-transformed metabolite concentrations  
 304 were then summed up among each category (Table 1) and another PCA was performed using  
 305 metabolite groups as discriminatory factors. An ANOVA with FDR-adjusted p-values was next  
 306 performed to compare concentrations of each metabolite between morphs. Finally, an ANOVA  
 307 tested differences in fat content between oviparous and viviparous morphs. All statistical  
 308 analyses were carried out using the R software (~~R Core Team 2017~~)(R Core Team 2017).

309

## 310 **Results**

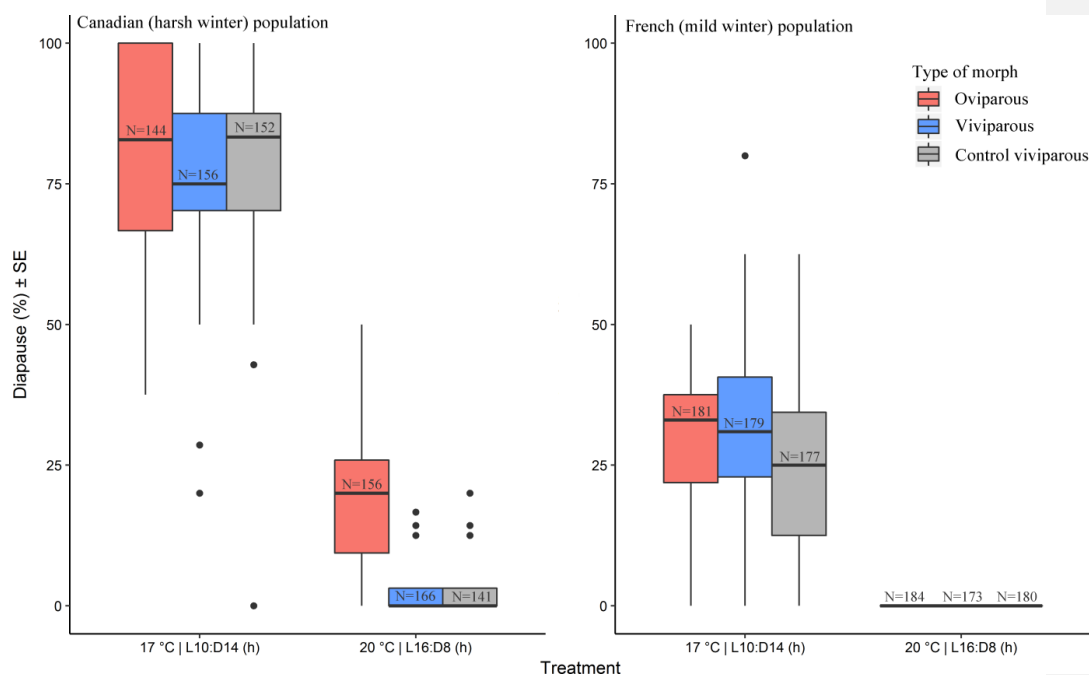
### 311 *Diapause incidence in the parasitoid A. ervi*

312 ~~Host~~In the Canadian (harsh winter area) population, diapause levels were affected by host  
 313 morph (GLMM,  $\chi^2=12.6$ ,  $df=2$ ,  $p<0.001$ ; Fig. 2) and abiotic conditions (GLMM,  $\chi^2=250.0$ ,  
 314  $df=1$ ,  $p<0.001$ ), with an interaction effect as host morphs influenced parasitoid diapause  
 315 incidence only ~~in the Canadian (harsh winter area) population~~ at 20 °C 16:8 h (L:D) (GLMM,  
 316  $\chi^2=16.9$ ,  $df=2$ ,  $p<0.001$ ; ~~Fig. 2~~). Diapause incidence was higher at 17 °C 10:14 h L:D than at 20  
 317 °C, 16:8 h L:D, for the Canadian population ( $76.9 \pm 2.5\%$  vs.  $9.0 \pm 1.5\%$ , respectively). At 20  
 318 °C, 16:8 h L:D, diapause incidence was higher when Canadian parasitoids developed in  
 319 oviparous aphids ( $19.4 \pm 3.0\%$  s.e.) than in viviparous aphids ( $3.6 \pm 1.3\%$ ),  $z=-4.3$ ,  
 320  $p<0.001$  or viviparous control aphids ( $3.8 \pm 1.4\%$ ). ~~The~~,  $z=-3.9$ ,  $p<0.001$ ).

321 In the French (mild winter area) population, the host morph did not have an effect on  
 322 diapause at 17 °C 10:14 h LD for both influence parasitoid populations (Fig. 2). diapause  
 323 (GLMM,  $\chi^2=1.84$ ,  $df=2$ ,  $p=0.39$ ), abiotic conditions did influence parasitoid diapause (GLMM,  
 324  $\chi^2=237.9$ ,  $df=1$ ,  $p<0.001$ ), but no interaction effect can be interpreted since no diapause was  
 325 expressed for the French population at 20 °C, 16:8 h L:D. Diapause incidence was higher at

326 17 °C 10:14 h L:D than at 20 °C, 16:8 h L:D, for the French population (27.9 ± 2.1% vs. 0%,  
 327 respectively). Random factors female identity and host exposition order had negligible effects  
 328 on total variance explained in both our models for both populations (variance ≤0.02).

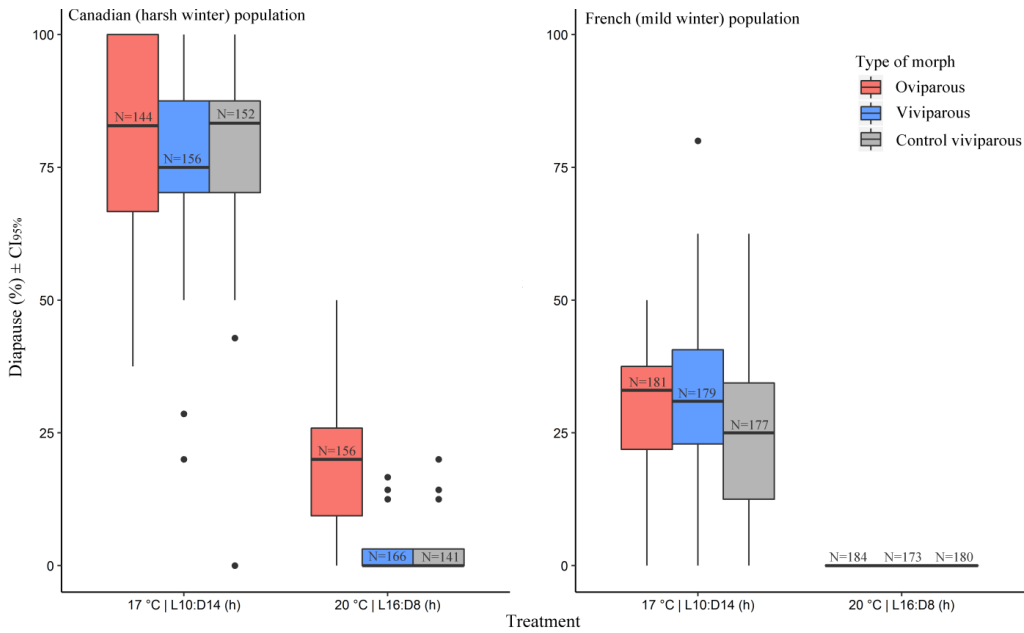
329 Diapause incidence was higher at 17 °C 10:14 h LD than at 20 °C, 16:8 h LD, for both the  
 330 Canadian (76.9 ± 2.5 % vs. 9.0 ± 1.5 %, GLMM,  $\chi^2=250$ ,  $df=1$ ,  $p<0.001$ ) and the French (27.9 ±  
 331 2.1 % vs. 0 %, GLMM,  $\chi^2=238$ ,  $df=1$ ,  $p<0.001$ ) populations. At 20 °C, 16:8 h LD, low levels of  
 332 diapause were observed for the Canadian population, except when parasitizing oviparous aphids  
 333 (Fig. 2), whereas no diapause was expressed for the French population.



334  
 335 Some female parasitoids produced offspring that had stronger responses to changes in host  
 336 morph or abiotic conditions than offspring of other females (Fig. 3). Data for each female are  
 337 made available as a supplementary material sheet. In some female's brood, there was no  
 338 variation in diapause plasticity in response to different biotic (morphs) or abiotic (photoperiod  
 339 and temperature) conditions (RN slope = 0). In the Canadian population at 17°C 10:14 h L:D,  
 340 reaction norm slopes (i.e., diapause level variations between conditions within a single brood)  
 341 ranged from -71% to 48%, for the diapause response to either oviparous or viviparous morphs  
 342 Fig. 3A). At 20°C 16:8 h L:D, these RN slopes ranged from -50% to 12% (Fig. 3B). In the  
 343 French population at 17°C 10:14 h L:D, RN slopes ranged from -29% to 38% for the diapause  
 344 response to either oviparous or viviparous morphs (Fig. 3C).

345 In the Canadian population, for parasitoids developing in viviparous morphs, RN slopes  
 346 ranged from -100% to -3% (Fig. 3D), and for parasitoids developing in oviparous morphs, RN  
 347 slopes ranged from -100% to -12% (Fig. 3E), for the diapause response to abiotic conditions  
 348 (17°C 10:14 h L:D vs. 20°C 16:8 h L:D). In the French population, for parasitoids developing in  
 349 viviparous morphs, RN slopes ranged from -80% to 0% (Fig. 3F), and for parasitoids  
 350 developing in oviparous morphs, RN slopes ranged from -50% to 0% (Fig. 3G) for the diapause  
 351 response to either abiotic conditions.

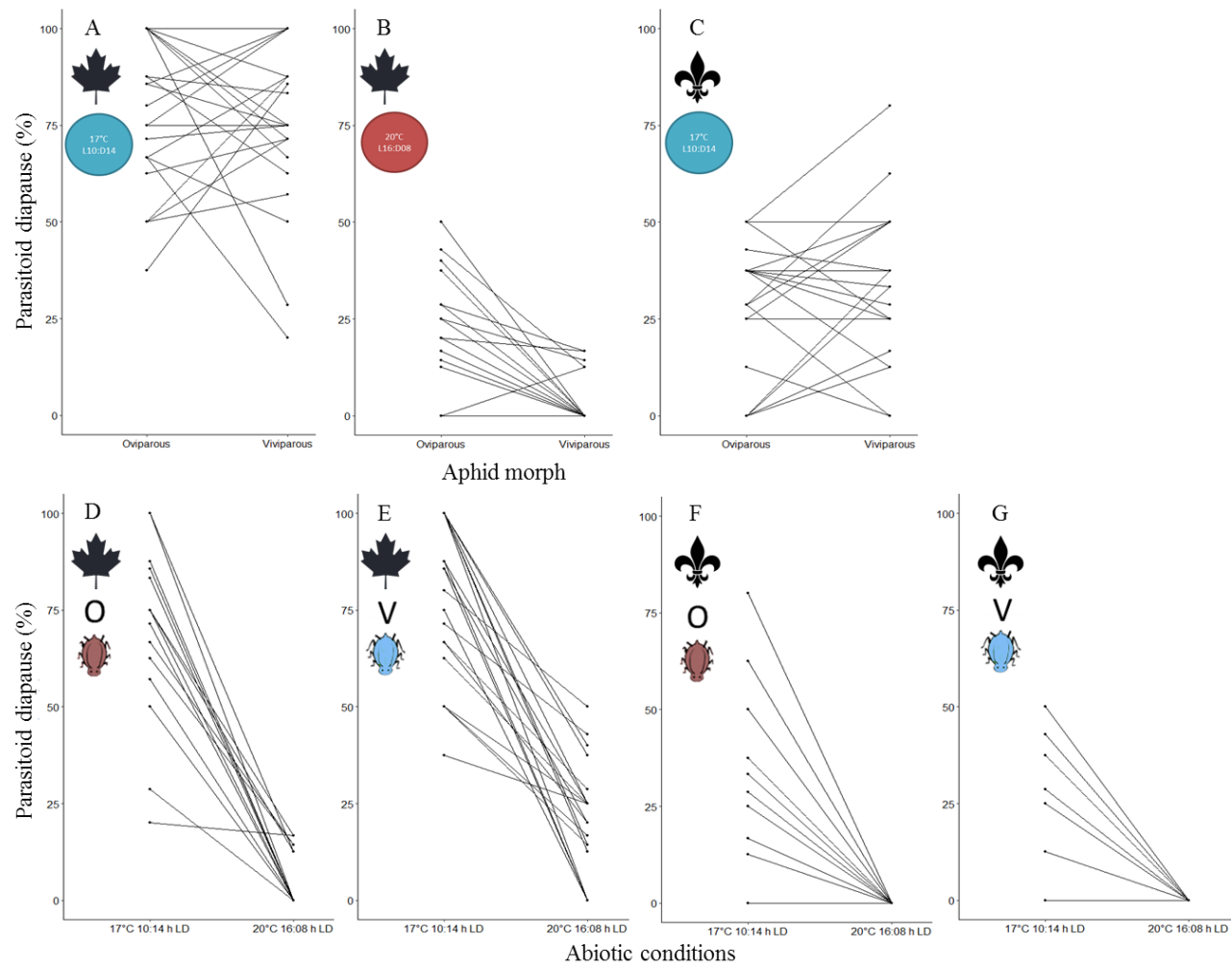
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

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354 Figure 2: Percent diapause incidence ( $\pm$  SE) CI<sub>95%</sub> in two *Aphidius ervi* populations. *Left*: Canadian population  
355 naturally experiencing harsh winter. *Right*: French population naturally experiencing mild winter. For both  
356 populations, three different morphs of the pea aphid *Acyrthosiphon pisum* (oviparous sexual females, viviparous  
357 parthenogenetic females produced under sexual-inductive conditions, and control viviparous females produced  
358 under non- sexual-inductive conditions) were used for parasitoid development, under two abiotic conditions  
359 (17- °C, 10:14 h L:D or 20°C, 16:8 h L:D). For each treatment, N represents the total number of parasitoid  
360 mummies used to calculate diapause incidence.

361



362

363 Figure 3: Reaction norms (RN) of diapause levels in the offspring of each parasitoid female from each parasitoid population (Canadian:  and French: ), both within  
 364 morphs at different abiotic conditions (top panel, **A & C**: 17°C 10:14 h L:D, **B**: 20°C 16:8 h L:D), and within abiotic conditions between morphs (bottom panel, **D & F**:

365 | oviparous morphs, E & G: viviparous morphs). RN for the French population at 20°C 16:8 h L:D are not displayed as no diapause was observed under these conditions. N=24  
366 | parasitoid female per condition. Note that some lines may be confounded.

367 *Metabolomic analyses and lipid reserves of aphid host morphs*

368 All measured compounds were found in both aphid morphs. The first and second principal  
 369 component (PC1 and PC2, respectively) of the PCA, accounted for 37.1-% and 26-% of the total  
 370 inertia, respectively (Fig. 3). Oviparous and viviparous female hosts were separated on PC1,  
 371 with oviparous females exhibiting significantly higher concentrations of trehalose, ribose,  
 372 arabinol, **GABA** gamma aminobutyric acid and mannose than viviparous ones (ANOVA, df=1,  
 373 p<0.05) (Fig. S1). Conversely, viviparous hosts had significantly higher concentrations of  
 374 alanine, gluconolactone, dopamine, putrescine, phenylalanine, glycerol, proline and quinic acid  
 375 than oviparous aphids (ANOVA, df=1, p<0.05) (details of metabolite amounts measured from  
 376 each morph are provided in Fig. S1). The second component of the PCA depicted the inter-  
 377 individual variation of metabolites within each of the two morphs (Fig. 34).

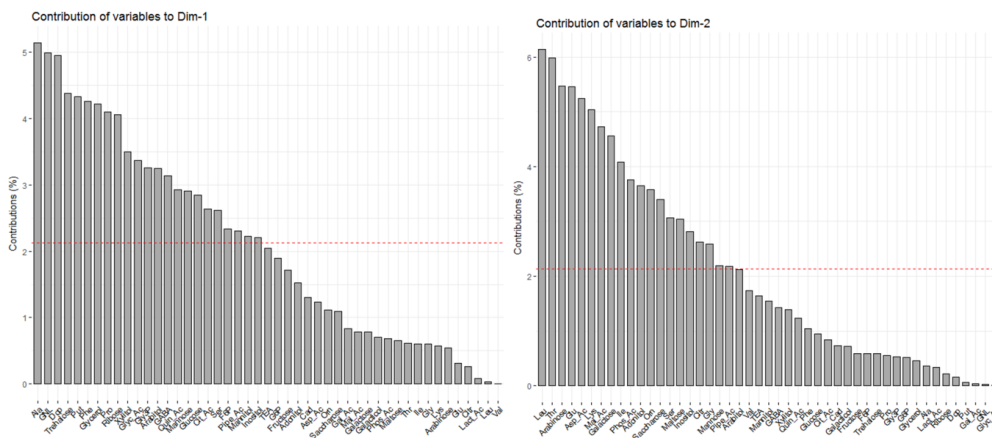
378 The analysis by metabolic family revealed that sugars / sugar phosphates (at the exception of  
 379 glucose) and polyols were measured in higher amounts in oviparous morphs, while amino acids,  
 380 amines and other metabolites were generally found in higher concentrations in viviparous hosts  
 381 (Fig. S2). Altogether, metabolic differences among oviparous and viviparous females revealed  
 382 that activities of the pathways involved in aminoacyl-tRNA biosynthesis and glutathione  
 383 metabolism were higher in viviparous females.

384 Oviparous hosts had a higher fat content ratio (mg fat/mg dry mass) than viviparous ones  
 385 ( $0.63 \pm 0.02$  and  $0.51 \pm 0.03$ , n= 52 and n= 23, respectively) (ANOVA, LR=8.0, df=1, p<0.005).  
 386 The fat mass represented  $37.8 \pm 0.8\%$  and  $33.3 \pm 1.3\%$  of the dry mass of oviparous and  
 387 viviparous morphs, respectively.

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388



389

390 Figure 34: Multivariate analysis (PCA) on the first two principal components (PC) representing links between  
 391 metabolic compounds (47 log-transformed variables, nmol.mg<sup>-1</sup>) and two aphid morphs (oviparous vs.  
 392 viviparous females) of *Acyrtosiphon pisum*. Enclosed figure in the upper panel shows a PCA of the six  
 393 metabolite categories. Confidence ellipses (95%) are constructed around each aphid group centroid (n=9  
 394 replicates by morph). Contributions of metabolite variables to PC1 and PC2 are provided in the two lower  
 395 panels. Abbreviations are listed in Table 1, supplementary figure S3. Abbreviations are Alanine (Ala), Aspartic  
 396 acid (Asp Ac), Cadaverine (Cad), Citric acid (Cit Ac), Citrulline (Citr), Dopamine (Dop), Fructose-6-phosphate  
 397 (F6P), Galacturonic acid (Gal Ac), Gamma aminobutyric acid (GABA), Gluconolactone (GNL), Glucose-6-  
 398 phosphate (G6P), Glutamic acid (Glu), Glyceric acid (Glyc Ac), Glycerol-3-phosphate (Gly3P), Glycine (Gly),  
 399 Isoleucine (Ile), Lactic acid (Lact Ac), Leucine (Leu), Lysine (Lys), Malic acid (Mal Ac), Ornithine (Orn),  
 400 Phenylalanine (Phe), Phosphoric acid (Phos Ac), Pipecolic acid (Pipe Ac), Proline (Pro), Putrescine (Put),  
 401 Quinic acid (Quin Ac), Serine (Ser), Threonine (Thr), Triethanolamine (TEA), Valine (Val).

402

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403 **Discussion**

404 Species interactions greatly contribute in shaping arthropods' seasonal ecological strategies,  
 405 because one species needs to synchronize or unsynchronize its life-cycle with interacting  
 406 partners or antagonists. However, biotic-induced diapause signals are poorly studied. A few  
 407 cases of predator-induced diapause have been documented in arthropods (Ślusarczyk 1995,  
 408 Kroon et al. 2008)(Ślusarczyk 1995, Kroon et al. 2008), such as in *Daphnia magna*  
 409 (Diplostraca: Daphniidae) in which the production of diapausing eggs is stimulated by predator  
 410 exudates and chemicals originating from injured conspecifics (Ślusarczyk 1999)(Ślusarczyk  
 411 1999). Reversely, low prey density was reported to influence summer diapause of the lady  
 412 beetle *Hippodamia undecimnotata* (Coleoptera: Coccinellidae) (Ipertti and Hodek 1974)(Ipertti  
 413 and Hodek 1974). Also, in herbivorous insects that require strong synchrony with their host  
 414 plant phenology, resuming activities after winter diapause is also influenced by the  
 415 physiological status of the plant (Leather et al. 1993)(Leather et al. 1993). Similarly, the host  
 416 plays a major role in parasitoid seasonal ecology. In addition to abiotic factors, such as  
 417 photoperiod and temperature, the host genotype, species, size, life-stage and abundance can  
 418 modulate parasitoid diapause (Tauber et al. 1986, Danks 1987)(Tauber et al. 1986, Danks 1987).

419 We report that parasitoids can use host oviparous morph as a cue for diapause induction,  
 420 with higher diapause incidence (up to 20%) expressed in *A. ervi* developing in oviparous  
 421 *A. pisum* females compared to viviparous conspecifics. This pattern is likely due to differences  
 422 in host physiology and metabolic contents. However, we have observed relatively high

423 intrapopulation variability within each female's offspring in response to the host morph, and in  
424 a lower extent in response to abiotic conditions, through the study of reaction norms.  
425 Polymorphism in the response of diapause induction cues (i.e., in plasticity) is known to be  
426 responsible for variability in diapause levels within populations experiencing different  
427 environmental conditions, but is still to be more deeply explored. As expected, parasitoids from  
428 the harsh winter environment expressed higher diapause levels than parasitoids from the mild  
429 winter environment. Of significance, only parasitoids from the harsh winter area and exposed to  
430 summer-like conditions relied on host morph as a cue for diapause induction.

431 Parasitoid populations of *A. ervi* from contrasted climatic environments (Canada and France)  
432 do not respond the same way to abiotic (photoperiod and temperature) and host cues. The  
433 French population of *Aphidius* spp. evolved under warming temperature conditions over the past  
434 decades, and this has allowed individuals of this species to remain active under mild winter  
435 conditions prevailing in this area, with none or small proportions of individuals entering  
436 diapause (Tougeron et al. 2017b). ~~In mild winter areas, non-diapausing parasitoids maintain~~  
437 ~~their populations by exploiting asexual anholocyclic aphid hosts during winter periods (Langer~~  
438 ~~and Hance 2000, Andrade et al. 2015, 2016) as sexual morphs are rare in these areas (Dedryver~~  
439 ~~et al. 2001). Diapause expression can be genetically lost or reduced in insects when they do not~~  
440 ~~experience the necessary environmental factors for its induction (e.g., Bradshaw and Holzapfel~~  
441 ~~2001, Garipey et al. 2015). Consequently, parasitoid populations from mild winter areas may~~  
442 ~~not have evolved a response to sexual hosts, or they may have lost the capacity to answer such a~~  
443 ~~cue to enter diapause(Tougeron et al. 2017b). In mild winter areas, non-diapausing parasitoids~~  
444 ~~maintain their populations by exploiting asexual anholocyclic aphid hosts during winter periods~~  
445 ~~(Langer and Hance 2000, Andrade et al. 2015, 2016) as sexual morphs are rare in these areas~~  
446 ~~(Dedryver et al. 2001). Diapause expression can be genetically lost or reduced in insects when~~  
447 ~~they do not experience the necessary environmental factors for its induction (e.g., Bradshaw and~~  
448 ~~Holzapfel 2001, Garipey et al. 2015). Consequently, parasitoid populations from mild winter~~  
449 ~~areas may not have evolved a response to sexual hosts, or they may have lost this capacity~~ under  
450 changing environments.

451 The opposite pattern is observed in Canadian populations, where all aphid parasitoids enter  
452 diapause during winter (Brodeur and McNeil 1994)(Brodeur and McNeil 1994). In these cold  
453 temperate regions, sexual morphs of aphids are produced at the end of the growing season, and  
454 represent the last hosts available for parasitoids before the onset of unfavorable winter  
455 conditions. In addition, parasitism of aphid sexual morphs on primary host plants allows  
456 parasitoids to overwinter nearby their hosts, thereby favoring host availability in spring for  
457 newly emerged parasitoids, and improving reproductive-cycles synchronization (Höller 1990,  
458 Christiansen-Weniger and Hardie 1997)(Höller 1990, Christiansen-Weniger and Hardie 1997).  
459 In regions with harsh winter climates, parasitoids have coevolved with the seasonal occurrence  
460 of host morphs and may use oviparous morphs as a convergent signal with temperature and  
461 photoperiod decrease in fall to enter diapause. Canadian Aphidiinae parasitoids begin to  
462 overwinter as early as mid-July, with all individuals being in diapause by early September  
463 (Brodeur and McNeil 1994, Tougeron et al. 2018)(Brodeur and McNeil 1994, Tougeron et al.  
464 2018). This seasonal pattern might be an adaptation to avoid early lethal frosts. Moreover, we  
465 showed that oviparous hosts only influenced diapause under summer-like conditions, suggesting  
466 that encountering this morph informs the parasitoids for upcoming deleterious conditions and  
467 modulates diapause expression. In natural settings, alternative host species can be present, and  
468 both anholocyclic and holocyclic aphid populations can coexist (Dedryver et al. 2001)(Dedryver



469 | [et al. 2001](#)), which may send confounding signals to parasitoids, and may explain why only a  
470 | fraction of the population responded to oviparous morphs. In Canada, oviparous morphs of the  
471 | pea aphid are present in the environment as soon as August (~~Lamb and Pointing 1972~~)([Lamb  
472 | and Pointing 1972](#)). In fall-like conditions, the morph effect was overridden by the  
473 | temperature/photoperiod effect, which remains the main signal for diapause induction.  
474 | Alternative diapause-inducing cues such as those associated with the host are usually viewed as  
475 | factors modulating diapause expression which is mainly triggered by temperature and  
476 | photoperiod (~~Tauber et al. 1986~~)([Tauber et al. 1986](#)). For example, in the polyphagous  
477 | herbivore *Choristoneura rosaceana* (Lepidoptera: Tortricidae), diapause is dependent upon  
478 | photoperiod and temperature, but under similar abiotic conditions, the proportion of larvae  
479 | entering diapause differs depending on the host-plant species (~~Hunter and McNeil  
480 | 1997~~)([Hunter and McNeil 1997](#)). Moreover, the effect of the host-plant was observed even  
481 | under photoperiod and temperature conditions known to induce low levels of diapause (~~Hunter  
482 | and McNeil 1997~~)([Hunter and McNeil 1997](#)). The relative importance of each environmental  
483 | cue at inducing diapause in insects remains to be evaluated for a significant number of species.

484 | Parasitoids' response to host morph could be partly shaped by maternal effects, as females  
485 | have the capacity to assess host quality through [a combination of physiological](#), morphological  
486 | ~~or behavioural and~~ chemical cues (~~van Baaren and Nénon 1996, Boivin et al. 2012~~)([van Baaren  
487 | and Nénon 1996, Boivin et al. 2012](#)). Developing immature parasitoids may also directly  
488 | respond to the quality and quantity of metabolites available from hosts, which could trigger the  
489 | onset of diapause. The overwintering metabolic and physiological syndrome is highly conserved  
490 | among insects (~~Tauber et al. 1986~~)([Tauber et al. 1986](#)), and both hosts and parasitoids may  
491 | respond to the same molecules involved in diapause initiation. As an example concurring to this  
492 | hypothesis, diapausing prepupae of the aphid parasitoid *Praon volucre* (Hymenoptera:  
493 | Braconidae) showed similar proportions of some sugars (e.g. trehalose, fructose) and polyols  
494 | (e.g. arabitol) (~~Colinet et al. 2012~~)([Colinet et al. 2012](#)) than non-parasitized oviparous morphs  
495 | of the pea aphid tested in our study. Our results suggest that high concentrations of some  
496 | polyols and sugar metabolites in the oviparous morphs, as well as accumulation of fat reserves  
497 | associated with the overwintering process, may either directly contribute to induce diapause in  
498 | parasitoids developing in such hosts or may trigger the internal physiological cascade  
499 | responsible for parasitoid diapause.

500 | [In the present work, oviparous \*A. pisum\* females have higher fat reserves than their  
501 | viviparous counterparts. This finding is consistent with the metabolic phenotypes of the hosts,  
502 | which revealed higher levels of sugar and sugar phosphate metabolites from the glycolytic  
503 | pathway in oviparous females, this pathway providing elementary bricks for fatty acid and  
504 | triacylglyceride \(TAG\) synthesis. Fatty acids serve as a main source of energy for physiological  
505 | or ecological processes, including flight, gametes production, egg maturation and hormones  
506 | synthesis \(Arrese and Soulages 2010\), and have been shown to represent up to 30% of aphids'  
507 | fresh mass \(Dillwith et al. 1993, Sayah 2008\). Interestingly, lipids can provide energy for  
508 | overwintering insects and sugars can be metabolized to produce sugar-based cryoprotectant  
509 | molecules \(Storey and Storey 1991, Hahn and Denlinger 2011, Sinclair and Marshall 2018\). In  
510 | oviparous females, the need for TAG may be higher than in viviparous ones, as eggs with yolk  
511 | \(vitellus\) are mostly composed of fat and proteins \(Brough and Dixon 1990\). Also, reserves  
512 | from the fat-body, including TAG and glycogen, play major roles in overwintering insects,  
513 | including diapause \(reviewed in Sinclair and Marshall 2018\) and could explain why oviparous  
514 | aphids have high fat content to prepare their eggs for successful overwintering. Diapause entails](#)

515 important energetic costs for insects (Ellers and Van Alphen 2002, Hahn and Denlinger 2011)  
516 and they may enter diapause only when a critical body-mass or amount of energetic reserves has  
517 been reached (Colinet et al. 2010); for parasitoids, developing in an oviparous host could  
518 contribute to reach this level.

519 Metabolites acting as compatible solutes greatly contribute to insect cold hardiness and  
520 overwintering survival (~~Storey and Storey 1991, Bale 2002, Hodkova and Hodek 2004~~)(~~Storey~~  
521 ~~and Storey 1991, Bale 2002, Hodkova and Hodek 2004~~). Metabolic analyses identified sugars  
522 and polyols in higher amounts in oviparous females containing eggs intended to overwinter.  
523 Overwintering eggs of the aphid *Hyaloperus pruni* (Homoptera: Aphididae) are characterized  
524 by high values of mannitol and trehalose (~~Sömme 1969~~)(~~Sömme 1969~~), as also observed in our  
525 *A. pisum* oviparous morphs. Glucose-6-phosphate and fructose were found at high  
526 concentrations in oviparous morphs of *A. pisum* and are precursors of sorbitol (~~Storey and~~  
527 ~~Storey 1991~~)(~~Storey and Storey 1991~~), a cryoprotective compound also observed in diapausing  
528 individuals of *P. volucre* parasitoids (~~Colinet et al. 2012~~)(~~Colinet et al. 2012~~). Fructose-6-  
529 phosphate is a precursor of mannitol, and both are cryoprotectant molecules (~~Storey and Storey~~  
530 ~~1991~~)(~~Storey and Storey 1991~~) highly concentrated in oviparous female hosts, and found in  
531 most of overwintering insects (~~Leather et al. 1993~~)(~~Leather et al. 1993~~). These metabolites may  
532 be responsible for diapause induction in parasitoids developing in oviparous morphs.  
533 ~~GABA~~Gamma aminobutyric acid was more concentrated in oviparous females and could also  
534 serve as an indirect seasonal cue for parasitoids because this neurotransmitter is known to be  
535 involved in insect perception of photoperiodic changes (~~Vieira et al. 2005~~)(~~Vieira et al. 2005~~).

536 Surprisingly, in viviparous females, we found high concentrations of glycerol a  
537 cryoprotective compound usually associated with the diapause syndrome (~~Hayward et al.~~  
538 ~~2005~~)(~~Hayward et al. 2005~~). As suggested by the high concentrations of glucose observed in  
539 these females, glycogen production through gluconeogenesis pathway could be used as main  
540 source of energy by these viviparous morphs (~~Dixon 1985~~)(~~Dixon 1985~~). In addition, observed  
541 physiological differences between host morphs are not necessarily linked to overwintering  
542 strategies. For example, viviparous aphids have high concentrations of proline, which is used as  
543 fuel for insect flight (~~Teulier et al. 2016~~)(~~Teulier et al. 2016~~). Viviparous aphids can rapidly  
544 produce ~~alate~~winged individuals for dispersal in case of overcrowding or degradation of host  
545 plant quality (~~Hardie 1980~~)(~~Hardie 1980~~).

546 To conclude, intra- and interspecific interactions are of primary importance for ecosystem  
547 functions, such as biological control, but still require deeper investigations in the context of  
548 diapause and seasonal strategies. Overwintering strategies are rapidly shifting in the context of  
549 climate change (Bradshaw and Holzapfel 2001, Bale and Hayward 2010) and may cause  
550 temporal mismatches between trophically interacting species (Tylianakis et al. 2008, Walther  
551 2010). Thus, potential bottom-up effects on diapause, such as reported in our study, should be  
552 given more attention and should be considered as a potential factor explaining the low levels of  
553 diapause expression in insects from mild winter areas, together with global warming (Jeffs and  
554 Lewis 2013, Andrade et al. 2016, Tougeron et al. 2017b). In addition, there was variation for  
555 plasticity in diapause induction among female genotypes, mostly in response to the parasitized  
556 morph but also to abiotic conditions, as determined by slopes of the reaction norms. This means  
557 that there is genetic polymorphism in diapause plasticity within populations, which may allow  
558 natural selection to act in the context of rapid environmental and climate changes (Sgrò et al.  
559 2016). Moreover, our results are of significance for the manipulation of insect diapause; e.g., in  
560 the context of mass rearing for the food industry, or for the biological control industry. More

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561 generally, a better appreciation of the processes governing phenology is needed to predict the  
562 consequences of such phenology changes on species interactions and synchrony across multiple  
563 trophic levels, community functioning and ecosystem services.

564

#### 565 **Authors' Contributions**

566 KT performed the diapause experiments, analyzed the data and wrote the manuscript. KT  
567 and DR performed the metabolomic experiments and analyzed the metabolomic data. All co-  
568 authors substantially contributed at designing protocols and revising the manuscript.

569

#### 570 **Acknowledgments**

571 We are grateful to ~~G~~both reviewers and both recommenders from PCI Ecology who made  
572 an excellent job in reviewing our manuscript and in providing strong advices on how to improve  
573 it. We thank G. Le Trionnaire at INRA Le Rheu for providing the aphid clones. We thank S.  
574 Llopis and J. Doyon for technical support and J. Jaquiéry for stimulating discussions. KT was  
575 funded by the Fyssen foundation, by the French Région Bretagne (ARED grant) and by the  
576 Canada Research Chair in Biological Control awarded to JB.

577

#### 578 **Data accessibility**

579 Metabolomics and diapause data have been made publicly available as a supplementary  
580 material attached to this publication.

581

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