

1 **Experimental evolution of virulence and associated traits in a *Drosophila melanogaster* –**
2 ***Wolbachia* symbiosis**

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12
13 **Abstract**

14 Evolutionary theory predicts that vertically transmitted symbionts are selected for low
15 virulence, as their fitness is directly correlated to that of their host. In contrast with this prediction,
16 the *Wolbachia* strain wMelPop drastically reduces its *Drosophila melanogaster* host lifespan at
17 high rearing temperatures. It is generally assumed that this feature is maintained because the
18 *D. melanogaster*–wMelPop symbiosis is usually not exposed to environmental conditions in which
19 the symbiont is virulent. To test this hypothesis, we submitted wMelPop-infected *D. melanogaster*
20 lines to 17 generations of experimental evolution at a high temperature, while enforcing late
21 reproduction **by artificial selection**. The fly survival was measured at different time points, as well

22 as two traits that have been proposed to be causally responsible for *w*MelPop virulence: its relative
23 density and the mean number of copies of octomom, an 8-genes region of the *Wolbachia* genome.
24 We hypothesised that these conditions (high temperature and late reproduction) would select for a
25 reduced *w*MelPop virulence, a reduced *w*MelPop density, and a reduced octomom copy number.
26 Our results indicate that density, octomom copy number and virulence are correlated to each other.
27 However, contrary to our expectations, we could not detect any reduction in virulence during the
28 course of evolution. We discuss the significance of our results with respect to the evolutionary
29 causes of *w*MelPop virulence.

30

31 **Introduction**

32 Symbionts live inside (endosymbionts) or on (ectosymbionts) bigger organisms referred to
33 as hosts. As a result, their evolutionary success crucially depends on their ability to colonize, or be
34 transmitted to, new hosts. Deleterious symbionts, also known as parasites, exploit their hosts in a
35 way that maximizes their transmission, but usually face a trade-off between instantaneous
36 transmission rate and opportunities for transmission. Indeed, the reproductive rate of a parasite is
37 likely to be positively correlated to both its instantaneous transmission rate and its virulence (the
38 reduction in fitness it incurs to its host). Increased virulence is in turn associated, notably through
39 increased host mortality, with a reduction in the opportunities for transmission (Anderson & May
40 1982, Ewald 1983, see Alizon et al. 2009 for a review and defence of this hypothesis).

41 Theoretical and experimental works suggest that the mode of transmission – which ranges
42 from fully vertical (from parents to offspring) to fully horizontal (contagion) – is an important
43 factor shaping parasite virulence. The fitness of vertically transmitted symbionts being tightly

44 correlated to that of their hosts, they are expected to be selected for low virulence (Ewald 1983).
45 This reasoning, just like the transmission/virulence trade-off hypothesis, rests solely on the
46 inter-host level of selection (*i.e.*, selection between symbionts infecting different hosts), ignoring
47 the potential effects of intra-host selection (*i.e.*, selection between symbionts infecting the same
48 host). Indeed, the most competitive symbionts within a host may replicate faster and be more
49 virulent than what would be optimal from the standpoint of inter-host selection (Alizon et al. 2013).
50 However, the causal link from vertical transmission to low virulence is well supported by the
51 experimental evolution of parasites following the enforcement of different transmission modes
52 (Bull et al. 1991, Turner et al. 1998, Messenger et al. 1999, Stewart et al. 2005). Accordingly, the
53 most widespread vertically-transmitted symbiont in insects, the intracellular bacterium *Wolbachia*,
54 is often found to be either avirulent (Hoffmann et al. 1994, Giordano et al. 1995, Bourtzis et al.
55 1996, Poinot & Mercot 1997, Hoffmann et al. 1998) or slightly virulent (Hoffmann et al. 1990,
56 Turelli & Hoffmann 1995, Clancy & Hoffmann 1997).

57 The most striking exception to this pattern is *wMelPop*, a *Wolbachia* strain hosted by
58 *Drosophila melanogaster* that drastically reduces its host lifespan at high rearing temperatures
59 (Min & Benzer 1997). This exceptional virulence raises the question of its causes, both proximal
60 and ultimate. Min & Benzer (1997) suggested that the proximal cause of the virulence of *wMelPop*
61 is its over-replication, which was further supported by subsequent studies (McGraw et al. 2002,
62 Strunov et al. 2013). Furthermore, the repetition of an eight-gene region (“octomom”), has recently
63 been proposed as the genomic basis of *wMelPop* high density and virulence (Chrostek & Teixeira
64 2015). Octomom includes genes encoding proteins potentially involved in DNA replication, repair,
65 recombination, transposition or transcription (Chrostek et al. 2003) and was found to be correlated
66 with *wMelPop* density and virulence (Chrostek & Teixeira 2015). However, both the link between

67 octomom copy number and density and the link between density and virulence have been called
68 into question. Rohrscheib et al. (2016) indeed found that differences in survival between flies
69 reared at 24°C and 29°C cannot be explained by differences in bacterial density. Similarly,
70 comparing flies of the same age but reared at different temperatures, or of different ages but reared
71 at the same temperature, they found no correlation between octomom copy number and either
72 density or virulence. However, to exclude the possibility that octomom copy number has an effect
73 on density, and density on virulence, the effect of these variables should be assessed independently
74 of temperature and age. The ultimate cause of *wMelPop* virulence has been assumed to be
75 non-adaptive: as this *Wolbachia* strain is not known to occur in nature but may have originated in
76 the lab, and as flies are typically reared at temperature lower than those at which *wMelPop* is
77 virulent and reproduce early in the lab, there is no selective pressure for reduced virulence
78 (Reynolds et al. 2003). It was indeed found that *wMelPop* reduces fly survival at 25°C, but not at
79 19°C (Reynolds et al. 2003). If this explanation is correct, raising *wMelPop*-infected flies at a high
80 temperature, while enforcing late reproduction, should select for: (i) a reduced *wMelPop* virulence,
81 (ii) a reduced *wMelPop* density (under the assumption that *wMelPop* virulence at high temperature
82 is due to its high density), and (iii) a reduced number of octomom copies (under the assumption
83 that *wMelPop* virulence at high temperature is due to its high octomom copy number).

84 In the present study, we used experimental evolution in conjunction with artificial selection
85 over 17 generations to test these hypotheses, raising *wMelPop* infected flies at 29°C, and enforcing
86 reproduction eight days after emergence. In addition, in one experimental condition, we fed the
87 flies with paraquat (1,1'-dimethyl-4,4'-bipyridinium dichloride), a pro-oxidant compound and
88 herbicide that was previously shown to increase the survival of *wMelPop*-infected flies (while
89 having no effect on the survival of uninfected flies) and to reduce the symbiotic density (Monnin

90 et al. 2016). We hypothesised that this treatment would reduce or cancel the selective pressure for
91 reduced virulence, density, and octomom copy number, resulting in higher values for these
92 parameters, compared to the control condition. The results of these experiments were interpreted
93 in light of our assessment of the relationship between octomom copy number and the age of the
94 host. Despite not being known to occur in nature, the *D. melanogaster-wMelPop* association can
95 help to understand the real-life dynamics of host-*Wolbachia* associations, especially in their early
96 stages. Indeed, one can expect that the low virulence typically observed in *Wolbachia* symbioses
97 is the outcome of a relatively long process of coevolution between the symbiotic partners. Newly
98 formed associations, by contrast, might be more likely to exhibit levels of virulence similar to what
99 is observed in *D. melanogaster-wMelPop* associations. Furthermore, our results contribute to the
100 ongoing debate concerning the relationships between *wMelPop* octomom copy number, density,
101 and virulence (Rohrscheib et al. 2016, Chrostek & Teixeira 2017, Rohrscheib et al. 2017).

102

103 **Material & methods**

104 Biological material

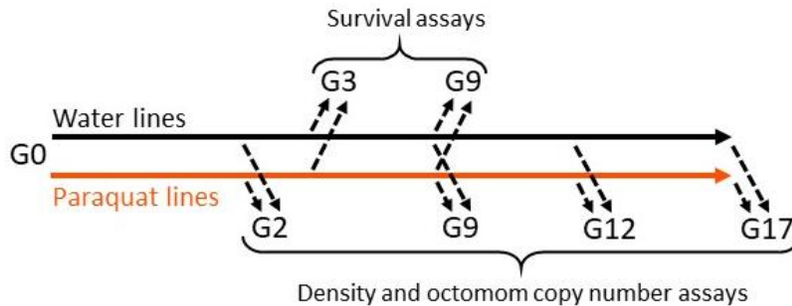
105 We used 17 isofemale lines of *wMelPop*-infected *D. melanogaster*^{w1118}. These lines,
106 originated from flies provided by Scott O'Neill (Monash University, Australia), were founded
107 approximately six months before the start of the experiment, and then maintained under controlled
108 rearing conditions at 18°C (12 LD cycle), a temperature at which *wMelPop* is expected to be
109 avirulent (Reynolds et al. (2003) found no effect of the infection on mortality rate at 19°C and no
110 virulence of *wMelPop* was ever detected at a lower temperature). The flies were maintained on
111 standard drosophila medium (David & Clavel 1965).

112

113 Experimental evolution procedures

114 The experiment was conducted at 29°C (12 LD cycle), a temperature at which *w*MelPop is
115 known to be virulent (Min & Benzer 1997, Strunov et al. 2013, Monnin et al. 2016). Emerging
116 flies from the 17 isofemale lines were mixed to constitute the founding generation G0 and were
117 allowed to lay eggs. Twenty experimental lines were then constituted, each originating from 100
118 randomly-picked eggs that were deposited in vials containing 1.5 g of *Drosophila* medium. The
119 experimental lines thus constituted were split into 10 “paraquat lines”, in which larvae developed
120 on drosophila medium supplemented with 150 µL of a 10 mM paraquat (1,1'dimethyl-4-
121 4'bipyridinium dichloride) solution (as described in Monnin et al. 2016), and 10 “water lines”, in
122 which the paraquat was replaced by 150 µL of water. The G1 flies emerging from each vial were
123 then allowed to lay their eggs on unsupplemented medium for eight days (the medium was not
124 supplemented with paraquat or water, as this would have been a confounding factor for the
125 subsequent measurements). The medium was changed regularly, including at the end of day seven,
126 and the eggs already laid on it were therefore removed. For each experimental line, a subset
127 (n=100) of the eggs laid on the eighth day following emergence was deposited in new vial similar
128 to those in which the parents developed – *i.e.*, paraquat-supplemented medium for the paraquat
129 lines and water-supplemented medium for the water lines. The protocol was repeated for the
130 emerging G2 flies, and again until G17. The evolution of *w*MelPop virulence, density and mean
131 octomom copy number was measured as follow: in each line, eggs laid seven days after emergence
132 were collected at G2, G3, G9, G12 and G17 and allowed to develop on unsupplemented medium.
133 Then, emerging females were used either to assess their survival, or their *Wolbachia* density and
134 mean octomom copy number (fig. 1). These traits were not measured at every single generation, as

135 we do not expect the changes from one generation to the next to be so great as to justify measuring
136 all traits at each generation.



137
138 **Figure 1.** Summary diagram of the experimental evolution experiment, showing the generations (G0-G17) at which
139 the different measurements were performed.

140
141 Though we chose – based on preliminary observations – the eight days period to enforce
142 late reproduction while reducing the risk of line extinction, several lines nevertheless went extinct
143 before the end of the experiment, as a result of lack of egg laying. Three control lines and two
144 paraquat lines went extinct between G3 and G9, four control lines and one paraquat line between
145 G9 and G12, and one control line and two paraquat lines between G12 and G17.

146
147 Survival assays

148 Fly survival was measured, as a proxy for *wMelPop* virulence, at G3 and G9. At emergence,
149 15 females from each experimental line were dispatched into three vials containing
150 sugar-supplemented agar (10% sugar). The surviving flies were then counted every day. The line
151 extinction rate – the ratio of lines that went extinct over the total number of lines – was calculated
152 for both the water and the paraquat lines, both during the 8 generations prior to G9 and during the

153 9 subsequent generations. It serves as another proxy for *wMelPop* virulence. We used Fisher's
154 exact tests to assess differences in extinction rates.

155

156 DNA extraction

157 The *wMelPop* density and mean octomom copy number were measured at G2, G9, G12 and
158 G17. For each generation and line, four of the emerging females were transferred on
159 sugar-supplemented agar (10% sugar), to be collected four days later and then kept at -80°C.
160 Because of technical issues during DNA extraction or qPCR, the final number of replicates varies
161 from 2 to 4 (mean number > 3.8). The DNA extractions were performed using the 96 wells Biobasic
162 EZ-10 BS4372 kit. The entire flies were mechanically crushed for 30 s at 25 Hz using a 5-mm
163 stainless steel bead in a TissueLyser (Qiagen). Three hundred µL of ACL solution and 20 µL of
164 16 g.L⁻¹ proteinase K were then added to the samples. Following an overnight incubation at 56°C,
165 300 µL of AB solution was added. The samples were then transferred on the plate from the kit, and
166 the purification was performed following the instructions of the manufacturer. The DNA was eluted
167 in 100 µL of elution buffer and stored at -20°C.

168

169 Quantitative PCR

170 Both relative *wMelPop* density and mean octomom copy number were measured in each
171 DNA extract by quantitative PCR, conforming to the Minimum Information for Publication of
172 Quantitative real-time PCR Experiments (MIQE) guidelines (Bustin et al. 2009). The relative
173 *wMelPop* density was quantified by amplifying two monocopy genes: one *Wolbachia* gene
174 (*wd0505*) and one *D. melanogaster* gene (*rp49*). The mean octomom copy number was quantified

175 by amplifying two *Wolbachia* genes: one that is part of the octomom region (*wd0513*), and one that
176 is not (*wd0505*). The primers used to amplify *rp49* were as follow: *rp49dd*-F: 5'-CTG-CCC-ACC-
177 GGA-TTC-AAG-3', *rp49dd*-R: 5'-CGA-TCT-CGC-CGC-AGT-AAA-C-3'. The primers used to
178 amplify the *wd0505* and *wd0513* genes were published in Chrostek et al. (2013). The primers were
179 synthesized by Eurogentec (Seraing, Belgium). qPCR mixes consisted of 5 μ L of Sso Advanced
180 SYBR Green Supermix (Bio-Rad, Hercules, California, USA), 1 μ L of water, 0.5 μ L of each primer
181 (10 mM), and 3 μ L of DNA samples diluted at 1/10. The reaction conditions for amplification were
182 95°C for 3 min, followed by 40 cycles of 95°C for 10 s, 59°C for 10 s and 68°C for 15 s. A melting
183 curve was recorded from 70°C to 95°C to ensure the specific amplification of the transcript.
184 Amplification and detection of DNA by quantitative PCR were performed with CFX96 instrument
185 (Bio-Rad). Standard curves were plotted using seven dilutions (10^2 to 10^7 copies) of a previously
186 amplified PCR product that had been purified using the kit Nucleospin extract II (Machery-Nagel).
187 Primer sets exhibited mean efficiencies of $93.2\% \pm 1.2$ for *rp49dd*, $91.0\% \pm 0.7$ for *wd0505*, and
188 $92.7\% \pm 0.6$ for *wd0513*. Two technical replicates per biological replicate were used for the
189 determination of DNA starting quantities. As the deviations between these duplicates were below
190 0.5 cycles, the mean C_p values were calculated. As PCR efficiency were close to 100%, we
191 calculated the relative w MelPop density and the mean octomom copy number using the following
192 formulas: $2^{C_{p_rp49}-C_{p_wd0505}}$ and $2^{C_{p_wd0505}-C_{p_wd0513}}$, respectively.

193

194 Relationship between mean octomom copy number and the age of the host

195 To evaluate the possibility that intra-host selection is involved in the evolution of w MelPop
196 octomom copy number, we assessed the relationship between the mean octomom copy number and
197 the age of the host. To do so, we performed qPCR as described above on samples obtained in the

198 course of a previously published experiment (Monnin et al. 2016). Briefly, wMelPop-infected
199 *D. melanogaster*^{w¹¹¹⁸} were raised until emergence on standard drosophila medium supplemented
200 with water or paraquat (as described above). Emerging females were either immediately collected
201 and kept at -80°C (1 fly per treatment) or transferred on sugar-supplemented agar (10% sugar) to
202 be collected at days 7, 14 and 21 of adulthood (four flies per day and per treatment). DNA
203 extractions were performed as described above.

204

205 Statistical analyses

206 We used the R software (version 3.3.2) for all analyses (R Core Team 2016).

207 We tested for evolution in survival between G3 and G9 using a mixed generalized linear
208 model with a gamma distribution (in the absence of censored data, a survival model is not
209 necessary, and the gamma distribution is suitable for continuous positive data). Individual
210 longevity was used as the response variable. Generation (G3 or G9) and treatment (water or
211 paraquat) were treated in the model as fixed factors, while the line and vial factors were included
212 as random effects.

213 The evolution of density and mean octomom copy number was tested using mixed linear
214 models, with the generation included as a fixed quantitative variable, the treatment as a fixed
215 qualitative variable, and the line factor included as a random effect. Normality and
216 homoscedasticity were checked graphically for both models.

217 The relationship between *Wolbachia* density and survival at G9 was tested by fitting a linear
218 model with median longevity as the response variable, mean density as a quantitative explanatory
219 variable, and the treatment (water or paraquat) as an explanatory factor.

220 The relationship between density and mean octomom copy number was tested by fitting a
221 mixed linear model with density as the response variable, mean octomom copy number as a fixed
222 quantitative explanatory variable, and generation, line and treatment as random effects.

223 The relationship between age and mean octomom copy number was tested by fitting a linear
224 model with mean octomom copy number as the response variable, age as a quantitative explanatory
225 variable and treatment as a qualitative explanatory variable.

226

227 **Results & discussion**

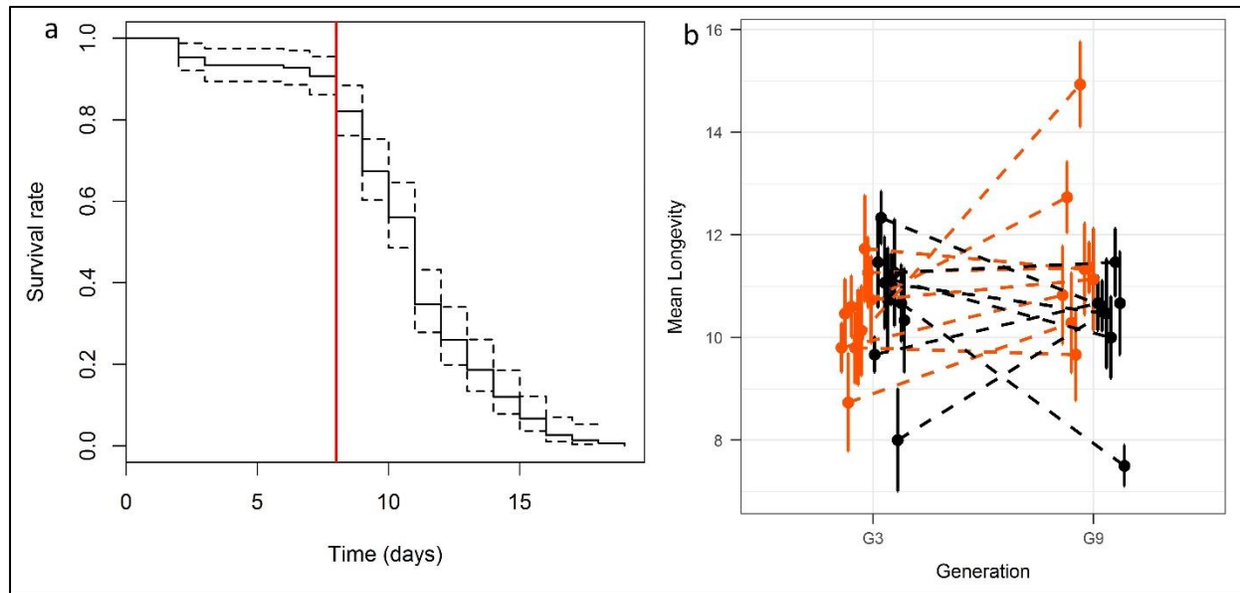
228 Test of the predictions

229 We followed the evolution of *wMelPop* virulence, density, and octomom copy number at a
230 high temperature while enforcing late reproduction. Our first prediction was that during the
231 experimental evolution, *wMelPop* virulence would have decreased in the **water** lines, and less so,
232 or not at all, in the paraquat lines. To test this, we compared fly survival at G3 and G9. **We can be**
233 **confident that fly mortality is induced by *wMelPop*, as the strong reduction of host longevity**
234 **induced by *wMelPop* has been established previously. A study performed in our lab (Monnin et al.**
235 **2016), using the same *D. melanogaster* line and the same survival protocol, found a large mean**
236 **difference in longevity (4.4 ± 0.66 days) between *wMelPop*-infected and uninfected individuals.**

237 Survival data at G3 confirmed that mortality began before the eighth day of the adult life of
238 the flies (fig. 2a), suggesting that the late reproduction we enforced should have induced a selective
239 pressure for decreased *wMelPop* virulence, at least in the **water** lines. In the paraquat lines, we
240 expected a lesser reduction, or no reduction at all, in virulence.

241 Neither the interaction between generation and treatment ($\chi^2_1=3.23$, $p=0.07$) nor the
242 treatment effect ($\chi^2_1=0.38$, $p=0.54$) was statistically significant, suggesting that the paraquat
243 treatment did not modify the evolution of survival, as compared to the water condition.
244 Furthermore, there was no effect of the generation factor, indicating that survival did not increase
245 between G3 and G9 ($\chi^2_1=0.98$, $p=0.32$; fig. 2b; median longevity at G3 in the control lines: 11 days;
246 at G9 in the control lines: 10.5 days; at G3 in the paraquat lines: 10 days; at G9 in the paraquat
247 lines: 11 days). As the survival was last assessed at G9, we cannot directly test whether it evolved
248 between G9 and G17. Given that an increased survival would presumably lead to a decreased
249 probability of extinction, we compared the rates of line extinction per generation before and after
250 G9. In the water condition, it was 0.038 before G9 (3 out of 10 lines went extinct at some point
251 during these eight generations), and 0.079 after G9 (5 out of 7 lines went extinct at some point
252 during these nine generations), showing that no increase in extinction rate occurred during the
253 experimental evolution, even after G9. Likewise, in the paraquat condition, the rate of line
254 extinction per generation was 0.025 before G9 (2 out of 10 lines went extinct at some point during
255 these eight generations), and 0.042 after G9 (3 out of 8 lines went extinct at some point during
256 these nine generations). Differences in extinction rate between the early and late generations were
257 not significant (Fisher's exact tests, $p=0.15$ in the water lines, $p=0.61$ in the paraquat lines, $p=0.16$
258 overall).

259 Taken together, these survival and extinction rates suggest the contrary to our prediction,
260 the virulence of *wMelPop* did not decrease during the experiment and was not differentially
261 affected by the treatment (paraquat or water).

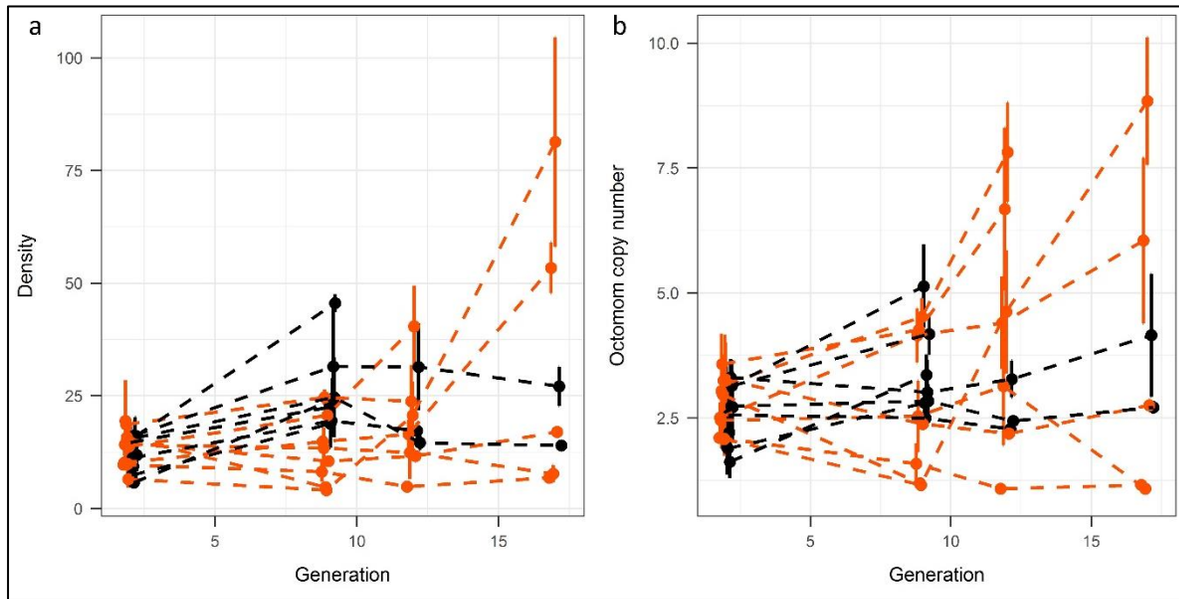


262
 263 **Figure 2.** Evolution of fly survival. (a) Survival of female flies in the **water** condition at G3. Dotted lines represent the
 264 95 % confidence interval. The red vertical line shows the age (8 days) at which reproduction was enforced during the
 265 experimental evolution. **The corresponding figure for the paraquat condition is provided as supplementary material** (b)
 266 Mean longevity (\pm standard error) in all lines in the paraquat (orange) and **water** (black) conditions, at generations 3
 267 and 9.

268

269 The discrepancy between the first prediction and the results invalidates the next two
 270 predictions, as the expected reduction in *Wolbachia* density was supposed to causally explain the
 271 expected reduction in virulence, and the expected reduction in mean octomom copy number was
 272 supposed to causally explain the expected reduction in density. Accordingly, we observed neither
 273 a decrease in density nor in mean octomom copy number during the experimental evolution. On
 274 the contrary, both density ($\chi^2_1=27.84$, $p<0.001$; fig. 3a) and mean octomom copy number
 275 ($\chi^2_1=16.95$, $p<0.001$; fig. 3b) increased (mean density in the **water** lines at G2: 11.73; G9: 26.33;
 276 G12: 21.05; G17: 20.50; in the paraquat lines at G2: 13.05; G9: 12.74; G12: 18.59; G17: 33.23;
 277 mean octomom copy number in the **water** lines at G2: 2.50; G9: 3.45; G12: 2.66; G17: 3.43; in the
 278 paraquat lines at G2: 2.84; G9: 2.67; G12: 4.27; G17: 3.97). Neither the interactions between
 279 generation and treatment (density: $\chi^2_1=0.52$, $p=0.47$; octomom copy number: $\chi^2_1=1.21$, $p=0.27$) nor

280 the treatment effects (density: $\chi^2_1=0.44$, $p=0.51$; octomom copy number: $\chi^2_1=0.44$, $p=0.51$) were
281 statistically significant.



282
283 **Figure 3.** Evolution of *Wolbachia* density and octomom copy number. (a) Mean *Wolbachia* relative density (\pm standard
284 error) in all lines in the paraquat (orange) and water (black) conditions, at generations 2, 9, 12 and 17. (b) Mean
285 octomom copy number (\pm standard error) in all lines in the paraquat (orange) and control (black) conditions, at
286 generations 2, 9, 12 and 17.

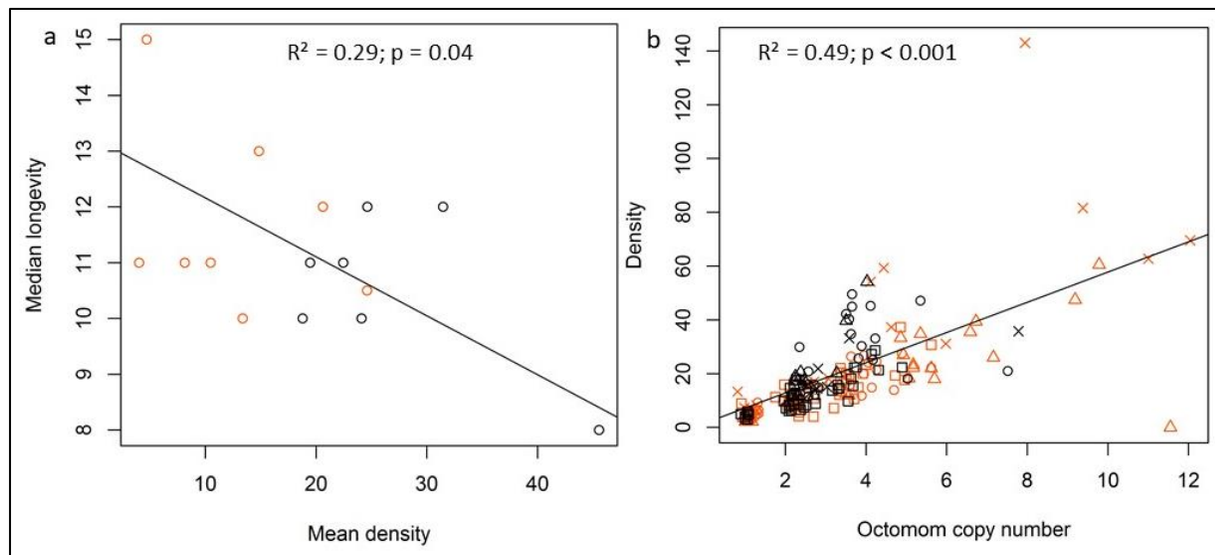
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288 Potential explanations for the falsification of the predictions

289 The falsification of our predictions is especially puzzling, as our results show that both
290 density and octomom copy number evolved, suggesting that it is not by lack of time or heritable
291 variability that the *D. melanogaster*-wMelPop interaction did not evolve in the expected ways.

292 As some lines went extinct before the end of the experiment, our statistical analyses may
293 have been biased if those extinctions were not random. However, it is unlikely to explain why our
294 predictions were not vindicated. Indeed, although a reduction in virulence could have been
295 concealed by the preferential extinction of lines exhibiting low virulence, this pattern is the

296 opposite of what we would expect. Furthermore, the observed increase in relative *Wolbachia*
297 density and mean octomom copy number is not likely to be due to a mere increase in genetic
298 similarity: at G2, no measured fly had a relative density superior to 40, contrary to 21% of the G17
299 flies (the maximum being 143); similarly, no flies at G2 had a mean octomom copy number over
300 6, contrary to 18% of the flies measured at G17 (the maximum being 12). Finally, we did not vary
301 the experimental temperature – a factor recently shown to impact on the evolution of
302 *D. melanogaster-Wolbachia* associations by Mazzucco et al. (2020) – and used only one host
303 genetic background (*D. melanogaster*<sup>w¹¹¹⁸) and one *Wolbachia* strain (*w*MelPop). We cannot
304 exclude the possibility that different conditions would have led to different coevolutionary
305 outcomes.</sup>

306 Among our starting assumptions was that the virulence of *w*MelPop is due to its high
307 density, which in turn is due to its high octomom copy number. It is therefore noteworthy that both
308 assumptions were recently called into question (Rohrscheib et al. 2016; see also Chroskek &
309 Teixeira 2017, Rohrscheib et al. 2017). If, as claimed by Rohrscheib et al. (2016), the virulence of
310 *w*MelPop is independent of its density and octomom copy number, our second and third predictions
311 do not hold. Although this would not explain why virulence did not decrease during the
312 experimental evolution, it would remove our puzzlement regarding the increase in *Wolbachia*
313 density and octomom copy number. However, our results do not support Rohrscheib et al. (2016)
314 findings. First, *w*MelPop density and fly survival were negatively related at G9, the only generation
315 at which both survival and density data were collected ($F_{1,18}=5.25$, $p=0.04$; fig. 4a). Second,
316 *w*MelPop density and mean octomom copy number were found to be positively related ($\chi^2_1=116.31$,
317 $p<0.001$; fig. 4b).



318

319 **Figure 4.** (a) Correlation between *Wolbachia* density and fly survival. Each circle represents a line in the paraquat
 320 (orange) or water (black) condition, at G9. The black line represents the regression of median longevity on mean
 321 density (data for both conditions are pooled, as the treatment effect was found not to be statistically significant).
 322 Statistics from Pearson's correlation test are indicated. (b) Correlation between *Wolbachia* density and octomom copy
 323 number. Each symbol represents an individual fly, from the paraquat (orange) or water (black) condition, at G2
 324 (squares), G9 (circles), G12 (triangles), or G17 (crosses). The black line represents the regression of density on
 325 octomom copy number. Statistics from Pearson's correlation test, with all conditions pooled together, are indicated.

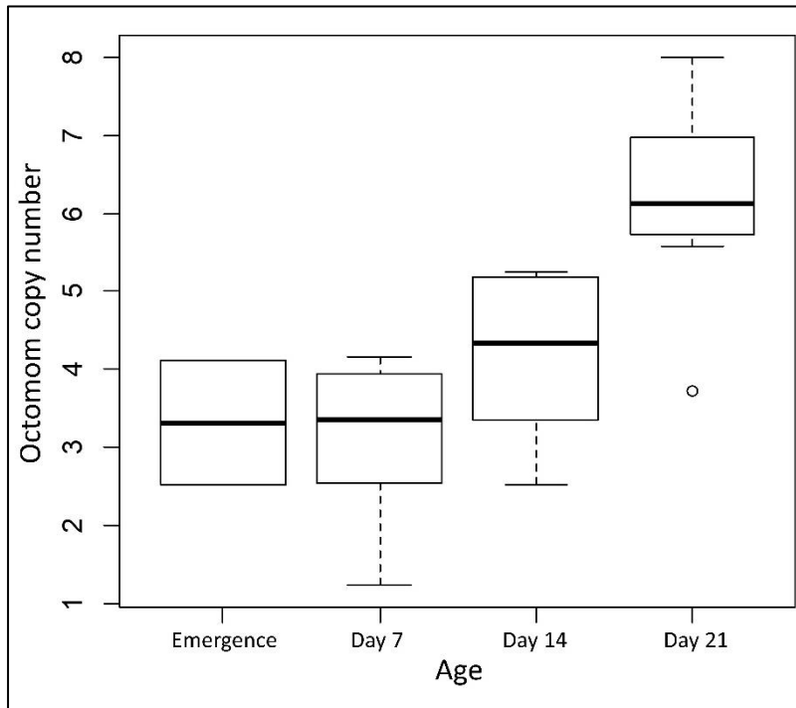
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327 It is therefore more likely that our assessment of the evolutionary forces at play, rather than
 328 our assumptions regarding the proximal causes of *wMelPop* virulence, was mistaken. The increase,
 329 over time, in both density and octomom copy number, appears to be consistent with two
 330 non-mutually exclusive hypotheses. First, this pattern could be explained by the relaxation of an
 331 unknown selective pressure maintaining a low octomom copy number in flies kept at relatively low
 332 temperatures without enforcement of late reproduction, perhaps amplified by a mutational bias
 333 favouring a high octomom copy number. As each new generation of experimental evolution was
 334 started with only 100 eggs, genetic drift could have been increased by the experimental
 335 evolution/selection protocol, possibly limiting the efficiency of natural selection. However, it
 336 remains difficult to see why the selective pressure in favour of a low octomom copy number should
 337 be weaker in the experimental evolution setting than in the maintenance conditions, as our current

338 knowledge of *wMelPop* suggests that the opposite is true: in the absence of *wMelPop*-induced
339 mortality in the maintenance conditions, the octomom copy number is not expected to be under
340 selection.

341 A second explanation for the observed pattern is that it results from directional selection in
342 favour of high density, high octomom copy number variants, more or less constrained, in each line,
343 by the amount of available heritable variation. Other things being equal, flies harbouring low
344 density, low octomom copy number should be longer-lived and therefore have more opportunities
345 to transmit their symbionts to their offspring than their counterparts infected by high density, high
346 octomom copy number variants. As a result, inter-host selection is expected to favour *wMelpop*
347 variants that do not reach densities so high that they compromise the fitness of their host. However,
348 intra-host selection is also likely to affect the evolution of virulence (Alizon et al. 2013). Indeed,
349 one can expect high density, high octomom copy number variants to replicate more rapidly than
350 (and therefore outnumber) their low density, low octomom copy number counterparts, which will
351 make them more likely to be transmitted to their host offspring. Crucially, the competitive
352 advantage of rapidly-replicating variants will increase with host age, as it is positively correlated
353 with the number of *wMelPop* generations. (At 29°C, the doubling time of *wMelPop* was estimated
354 by Duarte et al. (preprint) to vary from 0.88 to 1.38 days.) ‘Late reproduction’ may therefore allow
355 intra-host selection to be more powerful than inter-host selection. This is especially likely given
356 that survival was still high at eight days (fig. 2a), suggesting that inter-host selection may have
357 been relatively weak. Chrostek & Teixeira (2018) provide evidence for such intra-host selection
358 by showing that the mean octomom copy number increases as flies age and reach a plateau by day
359 eight of adulthood. Our own results confirm the increase ($F_{1,24}=25.64$, $p<0.001$), but no plateau
360 was observed (fig. 5). These results show that intra-host variability can exist, and therefore allow

361 for intra-host selection, despite the bottleneck experienced by vertically transmitted symbionts
362 (estimated to range from 850 to 8000 cells in aphid-*Buchnera* symbioses (Mira & Moran 2002)).



363
364 **Figure 5.** Relationship between the age of the fly and the octomom copy number of *wMelPop*. Data pooled from two
365 treatments (paraquat and water), as neither the interaction between age and treatment ($F_{1,22}=2.79$, $p=0.11$) nor the
366 treatment effect ($F_{1,23}=0.14$, $p=0.72$) were significant.

367
368 More research is needed to determine the evolutionary consequences of this intra-host
369 dynamic. Rapid evolution of octomom copy number in *wMelPop* suggests that intra-host selection
370 could play an important role in the evolution of vertically transmitted symbiont and may contribute
371 to explain the persistence of relatively virulent *Wolbachia* strains in nature.

372 Taken together, our results show that *Wolbachia* traits associated with virulence can evolve
373 rapidly following a switch in selection regime. These evolutionary changes challenge common
374 assumptions on the evolution of virulence in vertically transmitted symbionts, as traits positively
375 correlated with virulence were found to evolve. This could indicate that the intra-host level of

376 selection plays a significant, yet underexplored, role in shaping associations with vertically
377 transmitted symbionts.

378

379 **Data availability**

380 Raw data, scripts and additional figure are available online:

381 <https://doi.org/10.5281/zenodo.4065517>

382

383 **Conflict of interest disclosure**

384 The authors of this article declare that they have no financial conflict of interest with the content
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386

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393

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