

1 **Evolution of *Drosophila* resistance against different pathogens and infection routes**
2 **entails no detectable maintenance costs**

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22 **Abstract**

23 Pathogens exert a strong selective pressure on hosts, entailing host adaptation to
24 infection. This adaptation often affects negatively other fitness-related traits. Such
25 trade-offs may underlie the maintenance of genetic diversity for pathogen resistance.
26 Trade-offs can be tested with experimental evolution of host populations adapting to
27 parasites, using two approaches: (a) measuring changes in immunocompetence in
28 relaxed-selection lines and (b) comparing life-history traits of evolved and control lines
29 in pathogen-free environments. Here, we used both approaches to examine trade-offs
30 in *D. melanogaster* populations evolving for over 30 generations under infection with
31 *Drosophila C Virus* or the bacterium *Pseudomonas entomophila*, the latter through
32 different routes. We find that resistance is maintained after up to 30 generations of
33 relaxed selection. Moreover, no differences in several classical life-history traits
34 between control and evolved populations were found in pathogen-free environments,
35 even under stresses such as desiccation, nutrient limitation and high densities. Hence,
36 we did not detect any maintenance costs associated with evolved resistance to
37 pathogens. We hypothesize that extremely high selection pressures commonly used
38 lead to the disproportionate expression of costs relative to their actual occurrence in
39 natural systems. Still, the maintenance of genetic variation for pathogen resistance calls
40 for an explanation.

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45 **Introduction**

46 Several studies have shown that resistance to pathogens evolves rapidly in host
47 populations (Boots and Began 1993; Kraaijeveld and Godfray 1997; Lohse et al. 2006;
48 Zbinden et al. 2008; Martins et al. 2013). This indicates that standing genetic variation
49 (SGV) for host resistance to parasites is maintained in most systems. However, parasites
50 are ubiquitous and they pose a strong fitness cost upon hosts. Hence, high resistance
51 should be fixed in host populations. In other words, the seemingly paradoxical
52 occurrence of SGV for traits involved in fighting pathogenic infections calls for an
53 explanation. Such maintenance is often attributed to the occurrence of a trade-off
54 between resistance to pathogens and other fitness-related traits (for a review see
55 McKean and Lazzaro 2011).

56 Experimental evolution allows for robust tests of the occurrence of evolutionary-
57 relevant genetic trade-offs. Indeed, with this methodology, the ancestral state is known,
58 hence comparisons between control and evolved lines allows identifying traits modified
59 by a specific selection pressure as well as correlated responses to selection. Moreover,
60 the method avoids spurious correlations due to individuals (or their parents) having
61 been in different conditions, or subject to different recent evolutionary histories
62 (Kawecki et al. 2012; Magalhães and Matos 2012).

63 Trade-offs between immunity and fitness-related traits in experimentally-
64 evolving lines are tested using two main approaches. The first consists in creating lines
65 of relaxed selection (Lenski 1988; Ye et al. 2009; Meyer et al. 2010; Duncan et al. 2011).
66 These lines derive from populations evolving in the presence of the pathogen and are
67 then placed for several generations in pathogen-free conditions. The occurrence of a
68 trade-off is inferred if individuals from these lines show a lower performance when

69 exposed to pathogens, as compared to the pathogen-resistant ancestral population they
70 were derived from. In short, a costly defense is expected to be rapidly lost in the absence
71 of the pathogen it targets. This logic is appealing but may not be universal. Indeed,
72 reverting to the ancestral state may be prevented by the loss of genetic variation
73 allowing for such a reversion, although this possibility is seldom tested (but see Teotónio
74 and Rose 2000). Alternatively, resistance may be costly but evolution in a pathogen-free
75 environment selects for mutations that compensate such cost. This is widely shown in
76 antibiotic-resistant bacteria (reviewed in MacLean et al. 2010) but has never been tested
77 in multicellular sexual species, possibly because it relies upon the appearance of novel
78 mutations, which require large populations and a high number of generations.

79 Another possible approach to test such costs is by measuring the performance
80 of individuals from lines selected for pathogen resistance when placed in a pathogen-
81 free environment (Boots and Began 1993; Kraaijeveld and Godfray 1997; Lohse et al.
82 2006; Schwarzenbach and Ward 2006; Luong and Polak 2007; Cotter et al. 2008; Zbinden
83 et al. 2008; Vijendravarma et al. 2009; Koskella et al. 2012; cf. review in Duncan et al.
84 2011). Under such an approach, several life-history traits, thought to correlate with
85 fitness, can be measured. Moreover, these tests can be done in several environments.

86 Irrespective of the method used, all studies addressing the consequences of the
87 evolution of pathogen resistance have found a cost for this trait, with two exceptions.
88 First, using both methods described above, adaptation of the cabbage looper to a virus
89 was found to be free of cost (2002). Second, Meyer and colleagues (2010) found no cost
90 in *E. coli* resistance to phage T6 (but a cost in resistance to other phages). Therefore,
91 such costs seem to be the rule, with few exceptions. This ubiquity of costs to immunity

92 lends support to the hypothesis that such costs underlie the maintenance of SGV for
93 host resistance (Antonovics and Thrall 1994).

94 Experimental evolution using *Drosophila* as a model host has repeatedly shown
95 that the evolution of resistance to pathogens is costly (Kraaijeveld and Godfray 1997;
96 Fellowes et al. 1998; Luong and Polak 2007; Vijendravarma et al. 2009; Ye et al. 2009).

97 In our previous work, we have performed experimental evolution of an outbred
98 population of *Drosophila melanogaster* adapting to infection with different pathogens,
99 *Drosophila C virus* (DCV) or the gram-negative bacterium *P. entomophila*, the latter
100 being administrated via either an oral or a systemic route (Martins et al. 2013, 2014).

101 We found that these populations increased resistance against these challenges within
102 few generations, thereby demonstrating the presence of ample SGV for this trait. Here,
103 we took advantage of this resource to test whether *Drosophila* resistance to such
104 immune challenges entailed a cost. We did this using the two approaches mentioned
105 above: 1/ we created relaxed-selection lines, i.e., lines in which selection for pathogen
106 resistance was relaxed, and tested for its maintenance over several generations; and 2/
107 we compared the values of several life-history traits in control and evolved lines in
108 several pathogen-free environments, including the ancestral environment.

109

110 **Materials & Methods**

111 **Pathogen stocks and cultures**

112 *P. entomophila* (a generous gift of B. Lemaitre) was grown in LB inoculated with a single
113 bacterial colony, taken from glycerol stocks kept at -80 °C and streaked in fresh Petri
114 dishes. Bacteria were prepared from an overnight culture grown at 30 °C, centrifuged

115 and adjusted to the desired OD using fresh LB. Virus aliquots were grown and titrated as
116 described elsewhere (Teixeira et al. 2008), kept at -80 °C and thawed prior to infection.

117

118 **Experimental evolution lines**

119 From a highly outbred population of *D. melanogaster* (Martins et al. 2013), we derived
120 20 lines corresponding to 3 distinct immune challenges and 2 matched controls with 4
121 replicate lines each: a) oral infection with *P. entomophila* (BactOral), b) systemic
122 infection by pricking flies with *P. entomophila* (BactSys), c) systemic infection by pricking
123 flies with DCV (VirSys), d) one control under standard conditions (Control), and e) blank
124 injected controls (ControlSys). At each generation, 600 flies were exposed to each
125 challenge, and the survivors used to form the next generation. We selected an initial
126 concentration of pathogens that killed approximately 66% of the fly population. At each
127 generation, survival to infection was monitored by following the survival of 100-120
128 adults challenged with the same pathogen they were exposed to during selection every
129 day until at least the 10th day post-infection. Flies were maintained under constant
130 temperature (25 °C), humidity (60–70%) and light-darkness cycle (12:12), and fed with
131 standard cornmeal-agar medium. Detailed protocols for the selection experiment can
132 be found in our previously published work (Martins et al. 2013, 2014). We hereafter
133 refer to lines continuously exposed to the parasites as ‘Selection lines’, to distinguish
134 them from ‘Relaxed-Selection lines’, see below.

135 **Relaxed-Selection lines (and test to their immunocompetence)**

136 We first established that a plateau of resistance was reached in each selection regime.

137 This was estimated to occur whenever no difference in the response to pathogen

138 infection was found in five consecutive generations, which took place at different
139 periods for each selection regime. BactOral reached this plateau from generation 9
140 onwards, VirsSys from generation 21 onwards and BactSys from generation 25 onwards
141 (Martins et al. 2013, 2014). We then derived Relaxed-Selection lines, one per each
142 Selection line (i.e., 4 per Selection Regime, cf. Fig 1A). To do this, 600 individuals of each
143 population of a given Selection Regime were placed in new population cages.
144 Reproduction took place at the same days as the matching Selection lines, and in the
145 subsequent generations, the Relaxed-Selection population sizes (600 individuals)
146 mirrored those of the Control lines. Survival of Relaxed-Selection following exposure to
147 the parasites/route of infection matching to the corresponding Selection lines was
148 monitored daily until at least the 10th day post-infection at each generation, in parallel
149 with the Selection and Control lines.

150 **Fitness costs in parasite-free environments**

151 Fitness-related traits in parasite-free environments were compared between individuals
152 from Selection and Control lines. To avoid possible artefacts due to maternal effects,
153 flies used in these tests were the progeny of flies that spent at least one generation in a
154 common environment without pathogens, *i.e.*, in the standard environment of the base
155 population. These assays were performed at generations 23 or 24 for reproductive
156 output, development time and resistance to desiccation and starvation. Nutritional
157 restriction and competition assays were done more than 30 generations after the end
158 of the selection experiment (between generations 64 and 75 for all lines), hence evolved
159 lines had been under a Relaxed-Selection regime for 30 generations. Therefore, a test
160 for the maintenance of immunocompetence was performed on those lines at that

161 moment, to ensure that differences between control and evolved lines were still
162 present. This test was done as described in the last section.

163 Reproductive output

164 Reproductive output assays were designed to mimic the procedure followed during
165 experimental evolution. Fifteen male-female pairs from each Selection and Control lines
166 were transferred to fresh food vials 8-10 days post-eclosion and let to lay eggs for 48h.
167 Reproductive output was assayed as the number of adults emerging from pupae 12 days
168 after oviposition.

169 Development time

170 To determine the mean fly development time, 10 replicate groups of 5 uninfected
171 females (10-11 days old) were let to lay eggs for 1 hour in standard food vials. Egg never
172 exceeded 52 per vial (mean density 17). The assay conditions mimic the experimental
173 evolution procedure. The number of emerging adults was counted every 3 hours after
174 the 9th day post-oviposition.

175 Resistance to starvation and desiccation

176 For the desiccation assay, 100 individuals (males and females) from each population
177 were placed in groups of 10 in empty vials, and mortality was scored every 3 hours. For
178 the starvation assay, 100 individuals (males and females) from each population were
179 placed in groups of 10 in empty vials, with water supplied *ad libitum* by moisturizing the
180 vial plugs.

181 Nutritional restriction

182 For each assay, 200 eggs from each population were placed in 10 groups of 20 eggs, both
183 in standard food vials and nutritionally-restricted food (standard food diluted 1:8 with
184 water maintaining the agar concentration). Viability in both conditions was estimated as

185 the number of adults emerging from pupae. To determine the mean fly development
186 time, the number of emerging adults was counted every 12 hours after the 9th and 14th
187 day post-oviposition for standard and restricted food, respectively.

188 Larval competitive ability

189 Finally, we tested whether populations that had evolved increased immunocompetence
190 against each pathogen had lower larval competitive ability compared to control lines.
191 To this aim, we competed first instar larvae of the evolved populations (and their
192 controls) against the same outbred control population carrying an introgressed white
193 mutation. Pharates were weighted and classified as males or females, red eyes or white
194 eyes.

195 **Statistical analyses**

196 Relaxed selection

197 To compare survival across generations in the different Selection and Relaxed-Selection
198 lines, the proportion of individuals surviving at day 10 after infection in each vial was
199 first estimated using the Kaplan-Meier method. Subsequently, a generalized linear
200 mixed model (GLMM) was fitted to the data, assuming a binomial distribution and an
201 underlying logit link function. The proportion of survivors, weighted by the number of
202 individuals in each vial as dependent variable was fitted in a model with sex, generation
203 and regime (Control, Selection or Relaxed-Selection) as fixed factors. Line nested within
204 Selection Regime and sex at each generation was considered a random factor.

205 Subsequently, we tested for differences in survival between lines, both overall
206 and across generations. When differences in survival between Selected and Relaxed
207 selection lines were found, we then tested for changes in the mean difference between
208 Control and Selection or Relaxed-Selection lines, between the first and subsequent

209 generations after the derivation of the Relaxed-Selection lines. In addition, we also
210 tested if there was a linear trend for change (increase or decrease) across generations
211 in the mean survival of the different lines, by considering Generation an ordered factor.

212 Moreover, we tested for differences in the slope of the mean survival across
213 generations, by fitting a logistic regression mixed model with generation as a continuous
214 variable, assuming a binomial distribution and an underlying logit link function. The
215 proportion of survivors, weighted by the number of individuals in each vial as dependent
216 variable was fitted to a model with sex and regime (Control, Selection or Relaxed-
217 Selection) as fixed factors and generation of relaxed selection as a continuous covariate.

218 To compare survival among Control, Selection, and Relaxed-Selection lines in the
219 last generation of selection, we used a Cox's proportional hazards mixed effect model
220 for each treatment, with survival time of individual flies as the dependent variable,
221 Selection Regime and sex as fixed factors and replicate vial nested within line as a
222 random factor.

223 In the tests for maintenance of immunocompetence, done at generations 60-75
224 we used a GLMM identical to that used for the relaxed selection analysis, comparing
225 survival after infection between Control and Relaxed Selection lines.

226 Life-history traits in parasite-free environments

227 To compare reproductive output in the Control and Selection lines in the absence of
228 infection, we used a linear mixed model (LMM), with the number of hatching eggs within
229 48h by a single female as dependent variable, Selection Regime and Generation as fixed
230 factors and Replicate vial nested within line and generation as a random factor.

231 To compare development time among lines, we fitted a LMM with days to
232 eclosion of individual flies as dependent variable, Selection Regime as fixed factor and
233 replicate vial nested within line as a random factor.

234 To compare survival under starvation and desiccation conditions, we used a Cox's
235 proportional hazards mixed effect model for each treatment (starvation or desiccation),
236 with survival time of individual flies as the dependent variable, Selection Regime and sex
237 as fixed factors and replicate vial nested within line as a random variable. We also
238 compared differences in the mean time to death (TTD) between selection regimes. For
239 this, TTD was calculated for each vial, using the Kaplan-Meier method, and was fitted as
240 a dependent variable in a GLMM with sex and Selection Regime as fixed factors and line
241 nested within each Selection Regime and sex as random factor.

242 To compare viability in nutrient limiting conditions, we used a GLMM with the
243 number of eclosing vs non-eclosing individuals as a binomial variable, Selection Regime
244 and food type (Regular vs. Nutrient limited) and their interaction as fixed factors, and
245 test vials nested into line as random factors, with an underlying logit link function.
246 Development time was compared as above, including food type as an additional fixed
247 factor and removing egg density as covariate. Least-square estimates of viability and
248 development time were then compared between Selection Regimes, independently for
249 each food type.

250 To test for differences in larval competitive ability, the variable weight was log-
251 transformed to comply with normality. To confirm that a higher density implied a cost
252 in larval weight, we compared the weight in each density using a generalised mixed
253 model with competition level (either 15 or 30 flies from each line), selection regime and
254 sex, and their interactions, as fixed factors and replicate as random factor. Following a

255 significant effect of the density (cf. results) we then performed the analysis at the
256 highest density, to address potential costs in flies derived from the selection lines. To
257 this aim, we compared the weight of individuals from each selection regime to that of
258 tester individuals from the same assay using a glm with selection regime (either BactSys,
259 BactOral; ContSys, VyrSys or Tester populations), sex and their interaction as factors.

260 All statistical analyses were done in R (version 3.1.2). Linear mixed models were
261 fitted using the *lmer* function and generalized linear mixed models with the *glmer*
262 function, both in the “lme4” package in R. The effects of the fixed factors and of the
263 hierarchical interaction terms were compared using Type II Wald χ^2 tests (*Anova*
264 function in the “car” package). Contrasts of least-square means estimates and of
265 regression coefficients were done on the most parsimonious model, i.e. in models
266 including only significant ($P < 0.05$) factors and interactions, using the *lsmeans* and
267 *lstrends* function in the “lsmeans” package. Survival data was compared using the *coxme*
268 function in “coxme” package. Hierarchically nested models were compared using
269 likelihood ratio tests. The sex-averaged hazard ratios were then compared, using the
270 *glht* function in the “multcomp” package in R. The reported p -values for tests involving
271 multiple comparisons were adjusted using a sequential Bonferroni correction.

272

273 **Results**

274 **Maintenance of resistance under relaxed selection**

275 For all pathogen challenges, significant differences in survival were found among
276 Control, Selection, and Relaxed-Selection lines (Figure 1B and Table S1). This effect was
277 mainly caused by the difference between Control and either Selection or Relaxed-
278 Selection lines (Figure 1B). To get a more detailed description of mortality dynamics

279 upon infection of the different selection lines, we also measured survival over 10 days
280 after infection in flies from the last generation of selection (Figure 1C and Table S5).

281 Differences between both Selection and Relaxed-Selection lines to Controls were
282 always significant in the BactSys, BactOral and VirSys lines (Figure 1B and 1C), either
283 globally ($|z| > 23.5$, $P < 0.001$, $|z| > 29.3$, $P < 0.001$ and $|z| > 37.2$, $P < 0.001$, respectively),
284 at each generation ($|z| > 7.31$, $P < 0.001$, $|z| > 5.7$, $P < 0.001$ and $|z| > 9.46$, $P < 0.001$,
285 respectively, for all comparisons), or when comparing mortality dynamics in the last
286 generation of selection ($|z| > 5.58$, $P < 0.001$, $|z| > 10.06$, $P < 0.001$ and $|z| > 6.30$, $P <$
287 0.001 , respectively, for all comparisons). Excluding in the third generation of relaxed
288 selection, where the Relaxed-Selection lines showed significantly lower mortality the
289 Selection lines ($|z| = -2.87$, $P = 0.029$), we did not observe significant differences
290 between these lines at different generations ($|z| < 1.38$, $P > 0.999$, for all comparisons),
291 nor in the mortality dynamics in the last generation of Selection ($|z| = 0.83$, $P = 0.405$).
292 In the VirSys vs. VirSys-Relaxed comparisons, no differences were found when
293 comparing survival at each generation ($|z| < 2.49$, $P > 0.4$, for all comparisons), nor when
294 comparing the mortality dynamics in the last generation ($|z| = 0.38$, $p P = 0.704$; Tables
295 S2 and S6). We also did not find a significant difference in the linear slope of survival
296 across generations between the different selection regimes (GLMM, Generation X
297 Selection Regime effect, $\chi^2_2 < 3.79$, $P > 0.150$), despite a significant Generation effect
298 (Generation effect, $\chi^2_1 > 18.67$, $P < 0.001$), indicating no differences between the regimes
299 in the overall trend in survival across generations (Tables S3 and S4).

300 In contrast, there was a significant difference, between the BactOral lines and
301 their matched-Relaxed Selection lines ($|z| = 5.8$, $P < 0.001$), in 4 generations across the
302 experiment, including in the last generation of selection ($|z| = 3.63$, $P < 0.001$) (Table S2

303 and S4). This difference cannot be attributed to either an increased relative mortality in
304 the Relaxed-Selection lines (comparison between Control and Relaxed-Selection lines
305 remained constant across generations, $|z| < 1.74$, $P > 0.9$) or a decrease relative mortality
306 in the Selection lines (comparison between Control and Selection lines remained
307 constant, $|z| < 2.76$, $P > 0.53$).

308 To explore the reason for this difference, we tested changes in absolute survival
309 across generations, separately for the Selection, Control and Relaxed Selection Lines. In
310 this analysis, whereas in the Selection lines survival increased significantly ($|z| = 3.74$, P
311 < 0.001), this trait did not change significantly in Relaxed Selection and Control lines over
312 11 generations ($|z| = 1.44$, $P = 0.450$ and $|z| = 1.29$, $P = 0.595$, respectively). In agreement
313 with this finding, we also did not find a significant difference in the linear slope of
314 survival across generations among selection regimes (GLMM, Generation X Selection
315 Regime effect, $\chi^2_2 = 2.91$, $P = 0.233$), again indicating no differences among regimes in
316 changes in survival across generations (Tables S3 and S4). Therefore, we attribute the
317 small but significant differences between Selection and Relaxed-Selection lines (less
318 than 7% in the last generation of selection) to a marginal increase in survival in the
319 former (approximately 9%), where selection was continued, while there was no increase
320 (or decrease) in mortality in the latter.

321 At generations 60 and 70, at the moment we tested for larval competitive
322 ability, relaxed-selection lines were still significantly more immunocompetent than
323 control lines (Ime, BactOral vs Control: $z = 3.04$ $P = 0.0002$, BactSys vs ContSys $z = 8.28$ P
324 < 0.0001 , VirSys vs ContSys $z = 9.48$ $P < 0.0001$).

325

326 **Costs of resistance in parasite-free environments**

327 We also tested for the occurrence of trade-offs by comparing several life-history traits
328 between Selection and Control lines. We started by measuring the reproductive output
329 (Figure 2A) and developmental time at generation 23 and 24 (Figure 2B) in these lines in
330 the absence of infection. We found no effect of Selection Regime in the reproductive
331 output ($\chi^2_{24} = 0.640$, $P > 0.959$).

332 For developmental time (Figure 2B), and despite a statistically significant Selection
333 Regime and Relaxed-Selection Regime by egg density interaction ($\chi^2_{24} = 12.20$, $P = 0.016$,
334 Table S7), no difference between any Selection line and their matched Controls was
335 detected ($|t_{22}| < 2.21$, $P > 0.114$, Table S8).

336 Next we measured desiccation resistance and starvation resistance in Control vs
337 Selection lines. These stressors that have putative ecological importance for *Drosophila*
338 (David et al. 1983). For both traits we failed to detect statistically significant differences
339 between selection regimes (Table S9, Selection regime effect, $\chi^2_{24} < 5.21$, $P > 0.266$; χ^2_{24}
340 < 9.3 , $P > 0.053$ for both starvation and desiccation assays, considering either the mean
341 time to death or the full mortality dynamics, respectively). This indicates an absence of
342 a correlated response between adaptation to infection and both stress-related traits
343 (Figure 3).

344 Moreover, because it has been often argued that costs are more easily revealed
345 in nutrient limited environments (McKean et al. 2008), we measured egg-to-adult
346 viability and developmental time under these conditions (Figure 4A,B). Since these tests
347 were done in lines that derived from the Selection lines in the end of the selection
348 experiment, but maintained in control conditions (without selection) for > 30
349 generations, these lines represent a second set of Relaxed-Selection lines.

350 Although we detected increased mortality and developmental time in individuals
351 raised on nutritionally-limited food relative to those raised on standard food (Food type
352 effect $\chi^2_2 > 141.3$, $P < 0.001$, for both traits), no differences were detected in either
353 viability or development time among Selection regimes (Selection regime effect, $\chi^2_4 <$
354 7.4 , $P > 0.11$, in both traits; Table S10). Since we observed a significant Regime by Food
355 interaction in the viability assay ($\chi^2_4 < 12.99$, $P < 0.05$, Table S10), we tested for
356 differences between Selection and their matched Control lines independently in the
357 different food types. The absence of differences in viability among selection regimes was
358 confirmed in both food types ($|z| < 1.94$. and $|z| < 2.14$ for comparisons in standard and
359 nutritionally-limited food, respectively, $P > 0.194$, Table S11).

360 Concerning differences in weight following larval development at high or low densities,
361 the final model retained sex, density, selection regime, the interaction between sex and
362 each of the other factors, and the triple interaction. Overall, adults were smaller at the
363 highest density relative to the lowest, indicating an effect of competition on this trait
364 (glm, effect of density: $F_{1,165} = 74.99$, $P < 0.0001$, Figure 4C). We then compared the
365 weight of flies from each selection regime to that of tester flies from the same assay, at
366 the highest density. No differences were found between tester flies and flies from
367 ContSys, BactOral or VirSys regimes ($F_{1,24} = 1.996$, $P = 0.158$; $F_{1,52} = 0.938$ $P = 0.333$; $F_{1,38}$
368 $= 2.311$ $P = 0.128$, for ContSys, BactOral and Virsys, respectively, Figure 4c). In contrast,
369 flies from the BactSys selection regime were on average bigger than tester flies ($F_{1,41} =$
370 5.916 , $P = 0.015$, Figure 4c). Although the interaction between sex and selection regime
371 was never significant ($F > 1.562$, $P > 0.211$), the factor sex was always significant ($F <$
372 8.22 , $P < 0.004$), as males were on average lighter than females.

373

374 **Discussion**

375 In this study, we used a large-scale experimental evolution study addressing host
376 adaptation to pathogen infection to test for the occurrence of trade-offs between
377 immunity and other traits. We used two complementary methodologies (relaxation of
378 selection and direct measurements of costs in selected lines), and tested 12 Selection
379 lines, distributed over 3 different selection regimes, encompassing two distinct parasites
380 (viruses and bacteria) and two infection routes (oral or systemic). Taken together our
381 observations support the absence of maintenance costs in *Drosophila* populations
382 evolved for higher immunocompetence against pathogens.

383 Using lines subject to relaxed selection allows testing the response as a whole.
384 That is, had we observed a decrease in immunocompetence in individuals stemming
385 from those lines, we would have concluded that a trade-off with some fitness-related
386 trait existed. Nonetheless, we would not attribute this trade-off to a particular trait. The
387 fact that none of the lines in this study has lost its immunocompetence suggests that
388 these trade-offs with fitness-traits are absent in ancestral environment conditions. Still,
389 this pattern could have also been explained by a loss of genetic variation in the selection
390 lines, such that relaxed-selection lines would be stuck in a maladaptive peak (Teotónio
391 and Rose 2000). However, two lines of evidence suggest that this is not the case: first,
392 whole genome sequencing revealed that genetic variation in a subset of these lines was
393 the same in Control and Selection lines, and that even loci under selection did not reach
394 fixation (Martins et al. 2014). Second, the performance of relaxed-selection lines in the
395 ancestral, pathogen-free environment, showed no difference to Control for the fitness
396 traits measured. Together, these results indicate that adaptation of our populations to

397 pathogen infection entails no maintenance costs in conditions pertaining to the
398 ancestral environment.

399 To further understand how our evolved populations respond in different pathogen-
400 free environments, we performed direct tests for the occurrence of trade-offs between
401 immunity and several life-history traits. The problem with this approach is that we may
402 miss the trait in which the cost is expressed. However, we tested a comprehensive set
403 of classical life-history traits, namely reproductive output, developmental time,
404 starvation resistance, desiccation resistance and larval competitive ability, to maximize
405 the possibility of detecting trade-offs. Moreover, we measured these traits in both males
406 and females, thereby discarding the possibility of sexual antagonism for such costs
407 (Vincent and Sharp 2014). This further reinforces the notion that, in the pathogen-free
408 environment, evolution for increased survival upon infection by *P. entomophila* or DCV,
409 has no observable costs.

410 Given that the large majority of studies using experimental evolution detected
411 trade-offs between immunity and life-history traits (reviewed in Duncan et al. 2011), the
412 absence of such a trade-off calls for an explanation. First, although we can state that
413 maintenance costs were not present and that we did not find trade-offs related to the
414 tested traits, some costs in other traits or environments might exist. Indeed, we did find
415 a (relatively minor) cost of BactSys lines in presence of viruses: they performed worse
416 than control lines (Martins et al. 2013). The reverse, however, was not found: no costs
417 were detected of VirSys lines in presence of other pathogens when testing the
418 performance of these lines in presence of other pathogens (Martins et al. 2014).
419 Moreover, apart from survival (Martins et al. 2013, 2014) and reproduction after
420 infection (Figure S1), we did not test for the occurrence of deployment costs, or of costs

421 in many other environments. Second, a cost may have occurred at a transient state then
422 be compensated for during evolution. Although we know much about compensatory
423 evolution in bacteria, we know little about its occurrence and dynamics in sexual
424 organisms, with some remarkable exceptions in extensively-studied systems (e.g., Labbé
425 et al. 2007). However, compensatory evolution is not likely in the system used here
426 because the performance of relaxed-selection lines does not decrease and recovers
427 across generations: it is always similar to that of evolved lines. This suggests that no
428 transient cost was compensated for.

429 We hypothesize that the probability of finding a cost hinges on the selection
430 pressure posed on the populations: a high selection pressure may sweep away most of
431 the genetic variation that would allow for adaptation to the challenge posed, leaving
432 only the most effective but most costly alleles. Indeed, the selection protocol we used
433 was such that 33% of the population survived in the first generations (this percentage
434 then increased due to adaptation). In the other studies of adaptation to pathogens, the
435 selection pressure, when reported, was much higher, ranging from 90-95% mortality
436 (Kraaijeveld and Godfray 1997; Fellowes et al. 1998; Ye et al. 2009). In contrast, in the
437 single study that has also reported no cost in multicellular organisms, the selection
438 procedure was such that 20-30% of the hosts (a cabbage looper) survived (Milks and
439 Myers 2000). This reasoning may also explain why some studies failed to find a trade-
440 off with immunity when selecting for other life-history traits (Sanders et al. 2005; Kolss
441 et al. 2006; Hangartner et al. 2013). In particular, the results reported in Sanders et al.
442 (2005) are surprising, as the relaxed-selection process (i.e., selection for immunity and
443 measuring consequences in life-history traits) did reveal a trade-off. The traits selected
444 in these experiments (larval competitive ability, learning and reproductive investment,

445 respectively) have a looser link to survival than resistance to pathogens. Hence, it may
446 well be that the selection pressure that populations were exposed to in these studies
447 was lower than that of studies selecting for increased immunocompetence, and this may
448 account for the absence of a trade-off. Clearly, this hypothesis calls for a direct test. For
449 example, one could set up selection lines evolving in presence of the same parasite but
450 at different doses, and test whether trade-offs appeared in the treatments with higher
451 selection pressures only. In any case, the lack of symmetry in the trade-off between
452 immunity and other life-history traits suggests that the trade-off is not a universal
453 genetic characteristic of the organisms under study, but a conditional property, which
454 may hinge upon the selection pressure posed.

455 Unfortunately, it is not possible to validate this hypothesis with studies that have
456 used other approaches to test the occurrence of a cost of immunity. A cost was found
457 in circa 50% of such studies (reviewed in Labbé et al. 2010). However, either the
458 evolutionary trajectories leading to host resistance are unknown or resistant clones
459 have been generated via artificial selection, which may lead to spurious correlations
460 among traits (Rose 1984). Hence, these data cannot be used to test whether the
461 strength of selection underlies the probability of finding a cost (see also the discussion
462 in Labbé et al. (2010) for other potential confounding factors in that data set).

463 Our hypothesis, however, is congruent with data concerning pesticide
464 resistance. Indeed, in one of the best-documented examples of allele replacement in
465 the wild, Labbé et al. (2009) have shown that pesticide resistance in the mosquito *Culex*
466 *pipiens* in Southern France first evolved via a highly-resistant but highly-costly allele.
467 When mosquito populations were established in the treated area (hence selection for
468 increased pesticide resistance was weaker), this allele was replaced by one conferring a

469 lower cost. Similarly, Lopes et al. (2008) found no cost for resistance to levamisole in
470 experimentally-evolving *C. elegans* lines in which a dose killing initially 25% of individuals
471 was used. This contrasts with most studies of natural populations, in which a cost for
472 pesticide resistance was found (Coustau et al. 2000).

473 Given the low prevalence of costs in this system, the question remains: what
474 maintains genetic diversity for resistance to pathogens in our system? One possibility is
475 that alleles conferring resistance have a large effect, such that susceptibilities differ
476 widely in the population. This has been shown to allow for the maintenance of
477 polymorphisms for resistance even when the cost is negligible (Antonovics and Thrall
478 1994). In line with this, we have found that the majority of the selection response for
479 increased resistance to DCV could be attributed to alleles of 3 genes in our populations,
480 all of which with a considerable effect upon host survival (Martins et al. 2014).
481 Moreover, we have shown that adaptation to all immune challenges occurred via
482 resistance, rather than tolerance. Models predict that the maintenance of genetic
483 variation for resistance is more likely than for tolerance mechanisms, although a cost is
484 still necessary (Roy and Kirchner 2000). Another possibility is that the maintenance of
485 genetic diversity in host populations in the field is due to coevolutionary dynamics. In
486 that case, diversity for pathogen resistance may be maintained for a wider range of
487 parameters than contemplated in models that consider host evolution alone (Sasaki
488 2000; Best et al. 2010). Coevolution in natural populations of *Drosophila* could have
489 maintained the standing genetic variation present in our populations at the onset of
490 experimental evolution.

491 Overall, this study suggests that the occurrence of maintenance costs for
492 immunity traits is not a universal feature of organisms, raising questions as to (a) under

493 which conditions such costs evolve and (b) what maintains genetic diversity for costless
494 immunity traits.

495

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504 **Figure legends**

505

506 **Figure 1 – Increased immunocompetence is maintained in relaxed-selection**

507 **populations.** (A) Diagram representing the different selection regimes used in this study.

508 Lines represented by solid branches were challenged with a pathogen at every

509 generation (Selection) or kept unchallenged (Control). From each Selection line, a line

510 was derived and maintained in the ancestral environment (dashed lines, Relaxed-

511 Selection). (B) Mean survival (\pm 95% CI) 10 days post-infection of individuals from

512 Control (circles), Selection (squares) and Relaxed-Selection (triangles) lines, across 10 to

513 15 generations (see Materials & Methods). (C) Dynamics of survival after infection at the

514 last generation of relaxed selection. Control lines die much faster than either of its

515 counterparts, Selection or Relaxed-Selection lines, which display comparable profiles.

516

517 **Figure 2 – Reproductive output and developmental time of individuals from Control**

518 **and Selection lines in the absence of pathogens.** (A) Mean (\pm 95% CI) reproductive

519 output 5-7 days after females reached adulthood, (B) Mean egg-to-adults

520 developmental time from egg to adult.

521

522 **Figure 3 – Starvation and desiccation resistance of individuals from Control and**

523 **Selection lines.** Mean time to death (\pm 95% CI) after (A) starvation or (B) desiccation of

524 males (dark grey bars) and females (light grey bars).

525

526 **Figure 4 – Survival and developmental time of individuals from Control and Selection**

527 **lines in nutrient-limiting conditions.** Mean (\pm 95% CI) (A) egg-to-adult viability and (B)

528 development time of individuals developing in standard (left subpanel) and nutrient-
529 limited (right subpanel) medium. (C) Mean ($\pm 95\%$ CI) weight difference between
530 individuals from the experimental lines and Tester mutants (outbred [w1118]), at high
531 larval competition conditions (30:30 larvae in XX ml of food); light grey bars: females;
532 dark grey bars: males.

533

534

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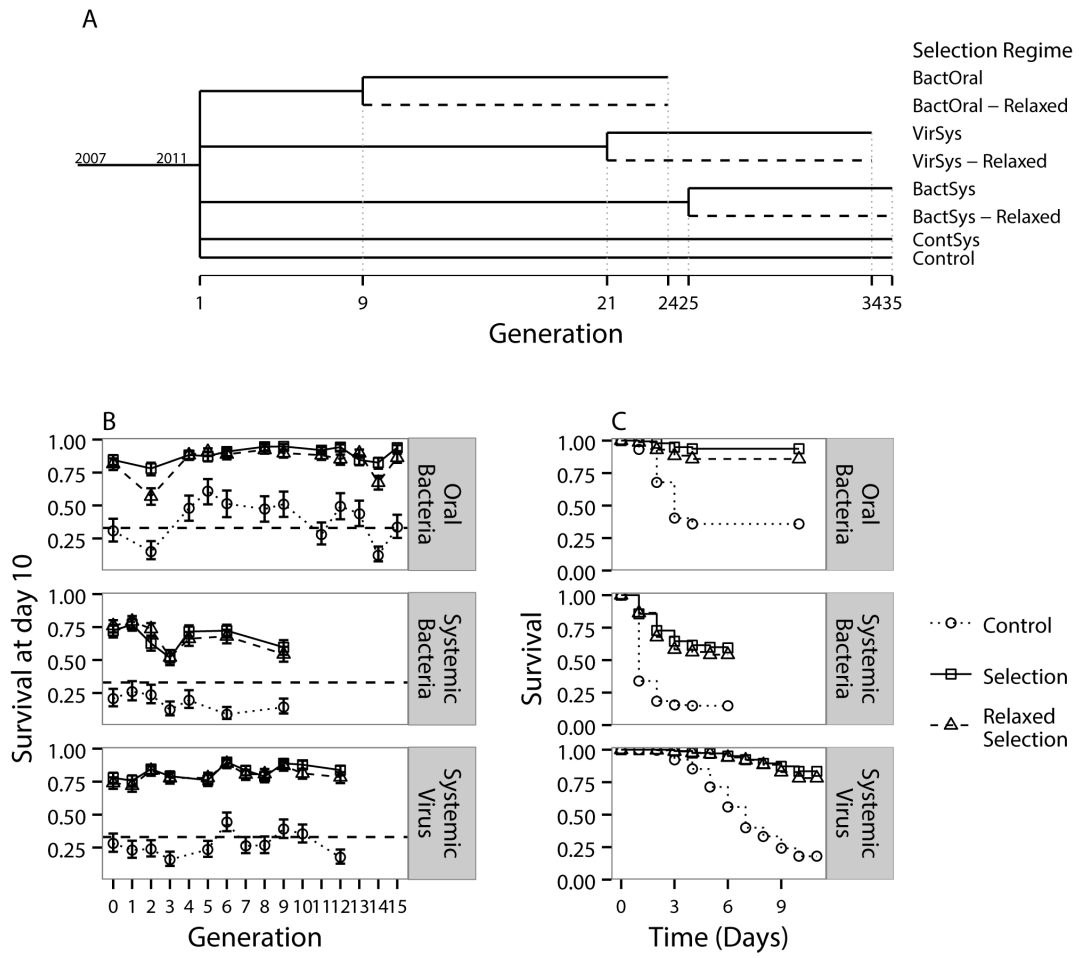
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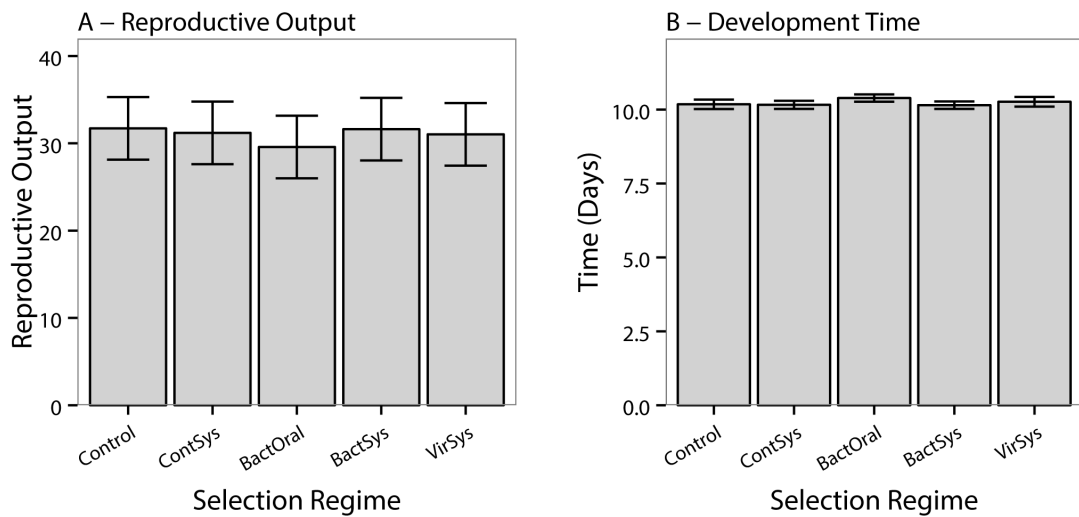
640 Figure 1



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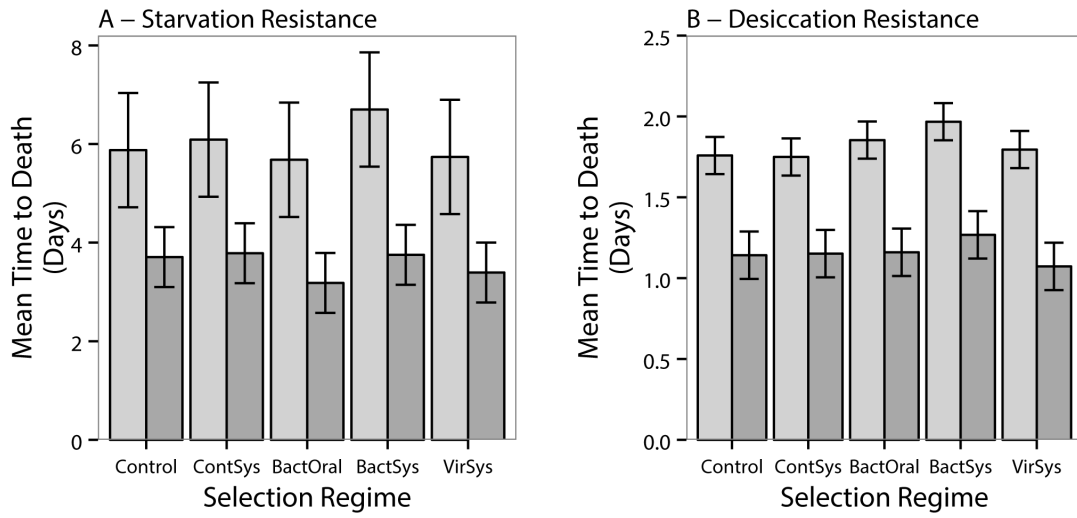
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643 Figure 2



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645 Figure 3



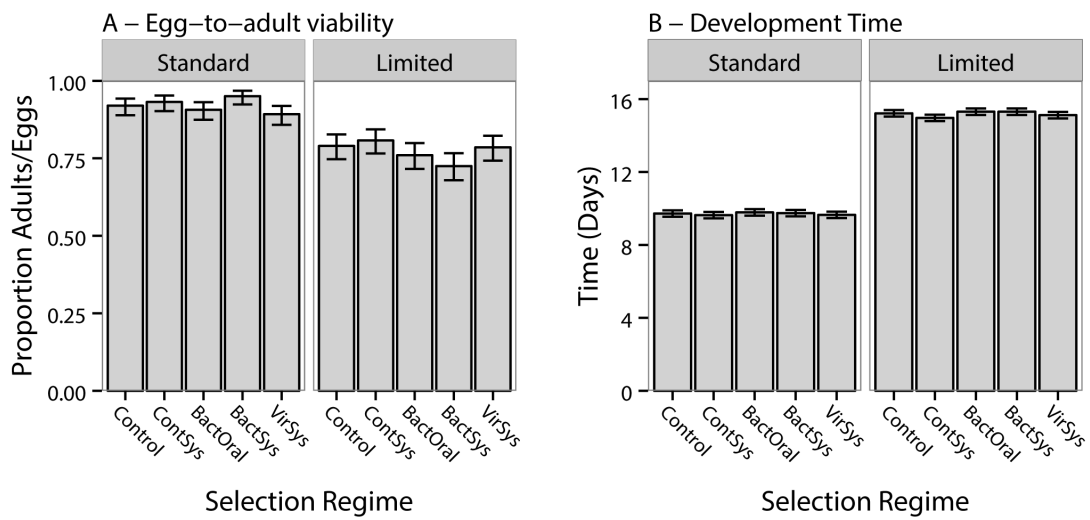
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650 Figure 4



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