

1 **Microbes are potential key players in the evolution of life histories and aging in**

2 ***Caenorhabditis elegans***

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23 **Abstract**

24 Microbes can have profound effects on host fitness and health and the appearance of late-  
25 onset diseases. Host-microbe interactions thus represent a major environmental context for  
26 healthy aging of the host and might also mediate trade-offs between life-history traits in the  
27 evolution of host senescence. Here we have used the nematode *Caenorhabditis elegans* to  
28 examine whether host-microbe interactions might modulate the evolution of host life-history  
29 and aging. We first characterized the effects of two non-pathogenic *Escherichia coli* strains, a  
30 pathogenic *E. coli* strain and a pathogenic *Serratia marcescens* strain on the reproductive  
31 schedule and survival of an outbred *C. elegans* population, to be used in an experimental  
32 evolution study. Secondly, to investigate the dependency of these effects on host genotype,  
33 we assayed population growth rates and survival of five representative *C. elegans* inbred  
34 strains in response to these microbes. Our results show that host-microbe interactions have a  
35 substantial, host-genotype-dependent impact on reproductive effort and survival of the  
36 nematode host. Although pathogenic bacteria reduced host survival, as expected, they did not  
37 necessarily decrease host fertility or population growth rate. Given such microbe-specific  
38 genotypic differences in host life history, we predict that the evolution of reproductive  
39 schedules and senescence in this system might be critically contingent upon host-microbe  
40 interactions, a hypothesis which we will be testing using experimental evolution in future  
41 work.

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43 Key words: *C. elegans*; microbes; host-microbe interactions; life-history evolution; aging;  
44 trade-offs.

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## 48 *Introduction*

49 Microbes are thought to have major effects on the evolution and speciation of host  
50 populations due to their ubiquitous presence and ability to influence host physiology and  
51 health (Bordenstein et al. 2001; Zilber-Rosenberg and Rosenberg 2008; McFall-Ngai et al.  
52 2013). While microbes are best known for their pathogenic or mutualistic effects, they can  
53 also modulate how hosts perceive and respond to stressful conditions. This has been  
54 observed, for example, in contexts as diverse as viral infections (Martinez et al. 2014), the  
55 autoimmune response (Langan et al. 2019), drug therapy (Pryor et al. 2019), metabolic  
56 dysfunction (Ussar et al. 2016), exposure to high temperatures (Xie et al. 2013; Howells et al.  
57 2016), and chemical toxicity (Coryell et al. 2018). Microbes can therefore impact adaptation  
58 of host populations to conditions that are apparently unrelated to the host-microbe interaction  
59 itself (Martinez et al. 2016; Faria et al. 2016; Bates et al. 2021; Hoang et al. 2021). This  
60 ability of microbes to modulate host adaptation to various stressors might thus play a  
61 fundamental but still poorly understood role in shaping the evolution of host life history and  
62 aging.

63 The progressive loss of physiological function leading to a decline in fecundity and  
64 increased mortality, which defines aging, can be explained by the reduced efficacy of  
65 selection in purging mutations that have deleterious effects late in life (Fisher 1930; Haldane  
66 1941; Medawar 1946, 1952; Williams 1957; Hamilton 1966; Rose 1991; Kirkwood and  
67 Austad 2000; Flatt and Schmidt 2009; Flatt and Partridge 2018). A major mechanism  
68 underlying the evolution of aging is antagonistic pleiotropy, i.e., the existence of alleles with  
69 antagonistic effects upon early and late life-history traits which lead to genetic trade-offs  
70 between fitness components (Medawar 1946, 1952; Williams 1957; Stearns 1989; Rose 1991;  
71 Flatt and Promislow 2009; Flatt 2020). Under this model, aging evolves because strong  
72 selection for beneficial fitness effects early in life outweighs the deleterious effects of these

73 alleles late in life when selection is weak (e.g., Williams 1957). A large body of work in  
74 numerous organisms, for example the nematode worm *Caenorhabditis elegans* (Anderson et  
75 al. 2011), the fruit fly *Drosophila melanogaster* (reviewed in Flatt 2020), or the fish *Poecilia*  
76 *reticulata* (Reznick et al. 1990), has revealed antagonistic pleiotropy underlying trade-offs by  
77 showing correlated responses to selection in major fitness components such as developmental  
78 rate, early and late-fecundity, and lifespan.

79 Even when populations harbor genetic variation sustaining antagonistic pleiotropic effects,  
80 they may not always experience phenotypic trade-offs nor correlated responses of life-history  
81 traits to selection, as these also depend on environmental factors (Giesel et al. 1982; Stearns  
82 1989; Ackermann et al. 2001; Sgró and Hoffmann 2004; Gutteling et al. 2007; Swanson et al.  
83 2016). For this reason, microbes are likely to play a relevant but underappreciated role in the  
84 evolution of aging, especially given their known effects on life-history traits (Little et al.  
85 2002; Deckaester et al. 2003; Brummel et al. 2004; Vale and Little 2012; Leroy et al. 2012;  
86 Laughton et al. 2014; Parker et al. 2014; Diaz et al. 2015; Zurowski et al. 2020) and their  
87 evolution (Sorci and Colbert 1995; Gibson et al. 2015; Walters et al. 2020). Causal  
88 relationships between the composition of the intestinal microbiome and aging observed in  
89 humans (Claesson et al. 2011) and other organisms (Clark et al. 2015; Sonowal et al. 2017;  
90 Bárcena et al. 2019) are consistent with this notion.

91 Studies with the *C. elegans* model hold great promise for an improved understanding of  
92 the interplay between host-microbe interactions and the evolution of aging. For example, the  
93 worm system has been extensively used in the identification of the genetic pathways  
94 underpinning aging and longevity (Garsin et al. 2003; Kurz and Tan 2004; Antebi 2007;  
95 Evans et al. 2008; Leroy et al. 2012), many of which are shared with humans (Kurz and Tan  
96 2004). At the same time, *C. elegans* has also been a valuable tool for studying host-microbe  
97 interactions (Tan et al. 1999; Abbalay et al. 2000; Garsin et al. 2003; Schulenburg et al. 2004;

98 Coolon et al. 2009; Leroy et al. 2012; Diaz et al. 2015; Dirksen et al. 2016; Schulenburg and  
99 Félix 2017) and how such interactions regulate host development, reproduction, metabolism,  
100 immunity and lifespan (MacNeil et al. 2013; Pang and Curran 2014; Chan et al. 2019 ).  
101 Notably, links between immunity and aging have been established in *C. elegans* (Kurz and  
102 Tan 2004), for example in the context of lifespan expansion obtained with specific bacterial  
103 metabolites (Virk et al. 2012; Han et al. 2017) or by transferring worms from their regular  
104 food source (*Escherichia coli* OP50) to other bacteria such as *Bacillus subtilis* (Aballay et al.  
105 2000; Portal-Celhay et al. 2012; Donato et al. 2017). Moreover, long-lived *C. elegans*  
106 mutants have been found to be resistant to pathogenic bacteria such as *Enterococcus faecalis*  
107 and *Staphylococcus aureus* (Garsin et al. 2003).

108 The extent to which the evolution of life histories and senescence in the nematode host  
109 might be contingent upon specific host-microbe interactions remains poorly understood. To  
110 begin to address this question, we sought to examine how different pathogenic and non-  
111 pathogenic bacteria impact the reproductive schedule and survival of *C. elegans*. To this end,  
112 we first measured survival and fertility throughout the reproductive lifespan of a genetically  
113 diverse population of *C. elegans* in response to two non-pathogenic *E. coli* strains, a  
114 pathogenic *E. coli* strain and a pathogenic *Serratia marcescens* strain. Second, we sought to  
115 investigate whether microbial effects on host life history might depend on host genotype, i.e.,  
116 whether host life history might be affected by interactions between microbe strain, host  
117 genotype and environment (i.e., reproductive timing). To do so, we assayed the population  
118 growth rates of five isogenic *C. elegans* strains, derived from different wild isolates, at the  
119 beginning of the reproductive period and past the reproductive peak in response to the same  
120 four bacterial strains mentioned above. Our results demonstrate that host-microbe interactions  
121 can have profound, host-genotype-dependent effects on reproductive effort and survival in *C.*  
122 *elegans*. Based on these results we conjecture that bacterial symbionts can modulate the

123 outcome of host life-history evolution in response to selection for different reproductive  
124 schedules, a prediction which we aim to test with experimental evolution in future work.

125

126 **Methods**

127 **BACTERIAL STRAINS**

128 Bacterial strains used in our experiments included two commonly employed non-pathogenic  
129 *Escherichia coli* strains, OP50 (Brenner 1974) and HT115(DE3) (Timmons et al. 2001), and  
130 two pathogenic strains, *E. coli* IAI1 (Picard et al. 1999; Diard et al. 2007) and *Serratia*  
131 *marcescens* Db11 (Flyg et al. 1980; Kurz et al. 2003). *E. coli* HT115(DE3) had been used as  
132 food during the establishment of the *C. elegans* D00 population described below. The strains  
133 *E. coli* HT115(DE3), *E. coli* OP50, and *S. marcescens* Db11 were obtained from the  
134 Caenorhabditis Genetics Center (CGC), and the *E. coli* IAI1 strain was kindly provided by  
135 Ivan Matic.

136 **NEMATODE POPULATIONS**

137 To assay life-history responses to the above-mentioned microbe strains we used an outbred  
138 experimental *C. elegans* population (D00) and 5 wild isolates (N2, CB4852, CB4855,  
139 CB4856, PX174). The D00 population was first described by Theologidis et al. (2014); it is a  
140 genetically diverse dioecious population with males and females, established by introgression  
141 of the *fog-2(q71)* mutant allele (Schedl and Kimble 1988) into the genetic background of a  
142 previously laboratory-adapted androdioecious population consisting of males and  
143 hermaphrodites (Teotónio et al. 2012; Chelo and Teotónio 2013). Throughout laboratory  
144 adaptation, worms were provided with *E. coli* HT115(DE3) as a food source and evolved  
145 under discrete (non-overlapping) generations imposed by a 4-day life-cycle, herein referred to  
146 as “early reproduction”. The D00 population is characterized by obligate outcrossing; its  
147 standing genetic variation results from an initial mixture of 16 isogenic strains, which were

148 chosen to represent a significant proportion of the known genetic diversity in *C. elegans*  
149 (Rockman and Kruglyak 2009; Teotónio et al. 2012). Here we have analyzed 5 of these 16  
150 isolates (N2, CB4852, CB4855, CB4856, PX174).

151 **GROWTH CONDITIONS**

152 Bacteria were grown overnight in NGM-lite solid media at 37 °C from LB-grown cultures.  
153 Nematode maintenance followed previously described protocols (Stiernagle 1999; Chelo  
154 2014). On day one, L1 larvae were seeded on NGM-lite supplemented with ampicillin (100  
155 mg/ml), carrying a confluent lawn of *E. coli* HT115(DE3). 10<sup>3</sup> larvae were used per plate,  
156 and development proceeded at 20°C and 80% (RH) for 72 hours, until day four of the life-  
157 cycle. Plates were washed with M9 buffer and a KOH:sodium hypochlorite solution was  
158 added (“bleaching”) to kill adults and larvae but allowing unhatched embryos to survive.  
159 Eclosion of first-stage larvae (L1) occurred overnight in 4 ml of M9 buffer with 2.5 mg/ml of  
160 tetracycline under constant shaking.

161 **REPRODUCTIVE SCHEDULE AND SURVIVAL OF THE D00 POPULATION**

162 Daily offspring number and survival were monitored to study the effects of different bacteria  
163 on individuals of the D00 population. Frozen (-80 °C) stock populations were thawed and  
164 maintained for two generations prior to the assay. To set up the experiment, 10<sup>3</sup> L1  
165 individuals were seeded on NGM-lite plates carrying each of the four bacteria and incubated  
166 until the beginning of day 3 (48 hours later). From each plate, 30 female larvae were placed  
167 (one larva per well) onto 24-well plates with antibiotic-free NGM-lite and matching bacteria,  
168 which had been grown from a 5 µl inoculum. Adult males from the same population and  
169 conditions, but which had been developing for one extra day, were added to the wells (two  
170 males per well). Individuals were transferred to fresh medium every 12 hours until day 6, and  
171 every 48 hours after day 6, until all individuals were found dead or considered to be missing.  
172 During the first five days, males that had died (or were missing) were replaced to ensure

173 mating and fertilization. After removal of adults, plates were kept in the incubator for one day  
174 and then transferred to 4°C for a maximum of two days before counting L2-L3 larvae under  
175 the stereoscope with 10x-30x magnification. These data were used to determine total fertility  
176 (lifetime reproductive success, LRS), variation in fertility through time and the age at first  
177 reproduction (AFR). Survival was scored based on daily observations during the entire period  
178 of the experiment. Monitoring of missing or dead females occurred at the time of transfer,  
179 and individuals were considered dead in the absence of movement or response when being  
180 gently touched with a platinum wire.

181 **GROWTH RATE OF THE D00 POPULATION AND OF INDIVIDUAL GENOTYPES**

182 Population growth rate in response to each of the four bacterial strains was measured at two  
183 different times: at 72 hours after L1 seed (transition from day 3 to day 4), i.e., within hours of  
184 reaching sexual maturity (“early reproduction”; Anderson et al. 2011) and at 114 hours post-  
185 seed (day 5; referred to as “delayed reproduction”). Frozen populations were thawed and  
186 maintained for two generations under standard maintenance conditions, plus one generation  
187 in presence of each bacterial strain for acclimatization. In the fourth generation, L1 larvae  
188 were seeded on NGM-lite plates ( $10^3$ /plate) with a lawn of each bacterial strain and allowed  
189 to develop for 72 or 114 hours. Following our standard maintenance protocol, cultures were  
190 bleached and the number of the live L1s was estimated the following day. Each estimate was  
191 obtained by pooling individuals from three plates. The D00 population and each of the five  
192 isolates (N2, CB4852, CB4855, CB4856, PX174) were assayed in independent experimental  
193 blocks. In the assays with the isolates, each block included the N2 strain feeding on *E. coli*  
194 HT115(DE3) as a common reference, the four different bacteria and the two time points. For  
195 each bacterial strain and each time point, we used five technical replicates for D00 and N2  
196 and four technical replicates for each of the other four isolates.

197

198 **SURVIVAL OF INDIVIDUAL GENOTYPES**

199 The effect of the four bacterial strains on survival was assayed for each of the five *C. elegans*  
200 isolates (CB4852, CB4855, CB4856, PX174). After thaw and growth for two generations  
201 under standard maintenance conditions, L1 larvae were seeded on NGM-lite media ( $10^3$   
202 individuals/plate) with a lawn of each of the four bacteria. 48 hours later (day 3), L4  
203 hermaphrodites were placed on 24-well NGM-lite plates (five individuals per well), with the  
204 corresponding bacteria, as described above for the survival assay of the D00 population. Each  
205 of the four non-N2 isolates was assayed in a different experimental block, which also  
206 included included N2 as a common reference. Four plates were used per block, and every  
207 plate included all four bacterial strains. Both the N2 and one of the non-N2 isolates were used  
208 in every plate, with N2 individuals occupying one fourth of the total number of wells. This  
209 experimental design enabled the estimation of plate effects within a block. In total, 480  
210 individuals were assayed in each block, with 120 being N2 individuals and 360 individuals  
211 from one of the other isogenic strains.

212 **DATA ANALYSIS**

213 Statistical analyses were performed in *R* (*R* Core Team 2019). Supplementary files with  
214 analyses and *R* code can be found at *FigShare* (see 10.6084/m9.figshare.15022566 for  
215 Supplementary Figures; and 10.6084/m9.figshare.15022599 for Supplementary Data and  
216 analysis scripts).

217 For fertility data, observations of 12h intervals were collapsed into daily measures until  
218 day 6 and into a single bin beyond that time. Thus, fertility reported for day 3 refers to  
219 embryos laid between 48 h and 72 h post-L1 seed, between 72 h to 96 h for day 4, between  
220 96 h to 120 h for day 5, between 120 h and 144 h for day 6, and 144 h onwards to “day 7”.  
221 Model fitting and model comparisons were performed with generalized linear models with  
222 appropriate error distributions (see below), and analysis of deviance was used to test for

223 significance. Parameter estimates were retrieved and tested with *emmeans* and *pairs* function  
224 (Lenth 2018). For pairwise comparisons, we used Tukey's post-hoc tests and report adjusted  
225 *p*-values. The reproductive schedule of the D00 population was modeled with a negative  
226 binomial distribution using the *R* function *glm.nb* in the MASS package. The following  
227 model was used: *Fertility* ~ *Bacteria* \* *Time*, where *Fertility* refers to the number of larvae  
228 observed per individual worm during a 24 h period, *Bacteria* represents the four bacterial  
229 strains tested, and *Time* is a categorical variable with 5 levels representing the day since the  
230 experimental set-up. Post-hoc comparisons were performed between fertility means within  
231 each day. Total fertility was modeled with a Poisson distribution using the *glm* function, as  
232 follows: *LRS* ~ *Bacteria*, *family* = "poisson"(link = "log"), where *LRS* is the total number of  
233 observed larvae. A Gaussian fit was used to analyze AFR with the following code: *AFR* ~  
234 *Bacteria*, *family* = "gaussian", where *AFR* (age at first reproduction) refers to the time  
235 between L1 seed and the time at which offspring was first observed.

236 Cox regression (proportional hazards analysis; Cox 1972) was used to test for differences  
237 in survivorship, with *E. coli* HT115(DE3) defining the baseline risk. The following model  
238 was implemented with the functions *Surv* and *coxph* in the *survival* package in *R* (Therneau  
239 2015): *Surv(S.time, S.event)* ~ *Bacteria*, with *S.time* being the time at which an individual was  
240 found dead or missing (*S.event*), assuming right-censored data. Kaplan-Meier estimation  
241 (Kaplan and Meier 1958) was used to estimate survival curves and mean lifespan.

242 Analysis of population growth rate was carried out using the natural logarithm (ln) of the  
243 observed rates. Whenever L1 larvae could not be detected, which would lead to growth rate  
244 estimates of zero (two samples; see Supplementary Table 2), values were replaced assuming  
245 that one L1 had been observed. To standardize the different blocks with *C. elegans* isolates,  
246 the growth rates of *C. elegans* N2 with *E. coli* HT115(DE3) were first estimated in each  
247 block and at each time point with a random-effects model using a block-specific baseline.

248 The following model was implemented in *R*:  $\log(\text{GrowthRate}) \sim \text{Time} * \text{Bacteria} *$   
249 *Celegans*,  $\text{offset} = \text{Block\_offset}$ , where *GrowthRate* is the observed L1 growth rate in  
250 consecutive generations, *Time* is the number of hours since L1 seed, *Bacteria* represents the  
251 bacterial strains, *Celegans* represents the 5 different isolates, and *Block\_offset* is the value of  
252 the block effects obtained with N2 and *E. coli* HT115(DE3).

253 Survival analysis of the *C. elegans* isolates was performed with Cox regression, using  
254 mixed-effect models with the *coxme* function in *R* (Thernau 2020) in order to include plate  
255 effects. The following model was used:  $\text{Surv}(\text{S.time}, \text{S.event}) \sim \text{Bacteria} * \text{Celegans} + (1 |$   
256 *Plate*) + *Block\_offset* (see above). Mean lifespan values based on Kaplan-Meier estimation  
257 were corrected by the values obtained for each block with N2 (see Supplementary Fig. 3).

258

259 **Results**

260 **DIFFERENT BACTERIA HAVE SPECIFIC EFFECTS ON THE LIFE-HISTORY**

261 **SCHEDULE OF A GENETICALLY DIVERSE *C. ELEGANS* POPULATION**

262 Reproduction and survival of the *C. elegans* D00 population were affected by the different  
263 bacteria in unique ways (Fig. 1). The presence of different bacteria had a significant effect on  
264 *C. elegans* survival ( $p$ -value  $< 0.0001$ , see Fig. 1A and Supplementary Fig. 1), with higher  
265 mortality risks observed in the presence of the pathogenic strains *E. coli* IAI1 ( $p = 0.0001$  for  
266 IAI1 vs. HT115;  $p < 0.0001$  for IAI1 vs. OP50) and *S. marcescens* Db11 ( $p = 0.02$  for Db11  
267 vs. HT115;  $p = 0.005$  for Db11 vs. OP50). Interestingly, no consistently detrimental (i.e.,  
268 pathogenic) effects were observed for fertility, even though fertility did vary with the  
269 different bacteria (Fig. 1B-C). Significant differences among bacterial strains were found for  
270 lifetime fertility ( $p < 0.0001$ , Fig. 1B), with the highest brood size being observed with *E. coli*  
271 HT115(DE3) ( $371 \pm 4$ , mean  $\pm$  standard error), followed by *E. coli* OP50 ( $185 \pm 2$ ), *E. coli*

272 IA1 ( $177 \pm 2$ ) and *S. marcescens* Db11 which resulted in a markedly reduced lifetime fertility  
273 ( $61 \pm 1$ ). These differences were also reflected in the reproductive schedule (Fig. 1C), as  
274 revealed by a significant time by bacteria interaction (likelihood ratio test, LRT = 42.5, df =  
275 12,  $p < 0.001$ ). Although fertility was always maximized at day 4, the relative contribution of  
276 offspring produced before and after this peak day was dependent on the bacterial strains. For  
277 instance, with *E. coli* HT115(DE3) the higher mean estimates of fertility observed throughout  
278 the entire reproductive lifespan of the host only become statistically significant after day 5  
279 (pairwise comparisons, adjusted  $p$ -values  $< 0.05$ ). In contrast, the initially diminished fertility  
280 of *S. marcescens* Db11 was no longer different from the majority of values observed with the  
281 three *E. coli* strains from day 4 onwards (Fig. 1C). Interestingly, comparing the start of  
282 offspring production of *S. marcescens* Db11 with the ones from all *E. coli* indicates a delay in  
283 reproduction, suggesting a possible interference with *C. elegans* development (Fig. 1D).

#### 284 **BACTERIA-HOST GENOTYPE INTERACTIONS AFFECT *C. ELEGANS***

#### 285 **POPULATION DYNAMICS**

286 The differential effects of the bacterial strains on the fertility dynamics of the host shown in  
287 Fig. 1C above suggest that the outcome of selection for early reproduction (reproduction at  
288 72 h) versus delayed reproduction (114 h), and hence the evolution of lifespan, might depend  
289 critically on microbe-host genotype interactions. Indeed, we found that the population growth  
290 rate (i.e., a fitness proxy) of the D00 population at those two ages was dependent on the  
291 bacterial strains (Fig. 2A): this was revealed by a significant time by bacteria interaction  
292 (LRT = 2.58, df = 3,  $p < 0.001$ ), with different slopes for *E. coli* IAI1 (CI = -0.028 to -0.014  
293 per hour), *E. coli* HT115(DE3) (CI = -0.011 to 0.002 per hour), *E. coli* OP50 (CI = -0.009 to  
294 0.004 per hour), and *S. marcescens* Db11 (CI = 0.006 to 0.0120 per hour). The main effects  
295 of time (LRT = 0.25, df = 1,  $p = 0.03$ ) and bacterial strain (LRT = 1.89, df = 3,  $p < 0.0001$ )  
296 were also significant.

297 Importantly, the time-bacteria interaction effects on growth rates also varied among the  
298 five *C. elegans* genotypes, as revealed by the pervasive crossing of lines in Fig. 2B and  
299 confirmed by a significant three-way interaction term (LRT = 17.71, df = 12,  $p < 0.0001$ )  
300 (main effects and all two-way interactions were also significant, not shown). Individual plots  
301 by bacterial strain (Supplementary Fig. 2) indicate that these effects were comparable to those  
302 obtained for the D00 population. This can be seen, for example, with *E. coli* IAI1 which  
303 imposed the largest average decrease in growth rate with time (CI = -0.097 to -0.084 per  
304 hour), or with *S. marcescens* Db11 which caused a shallower slope (CI = -0.015 to -0.003 per  
305 hour). In this latter case, it is noteworthy that the usual reduction of growth rate with time  
306 was reversed for the CB4855 genotype in presence of *S. marcescens* Db11 (Supplementary  
307 Fig. 2D).

308 **BACTERIA-HOST GENOTYPE INTERACTIONS MODULATE *C. ELEGANS***

309 **LIFESPAN**

310 The different bacteria also affected the adult survival of the *C. elegans* isolates (Fig. 2C and  
311 Supplementary Fig. 3,  $\chi^2 = 629.6$ , df = 3,  $p < 0.001$ ), with a significant bacteria-host genotype  
312 interaction on lifespan ( $\chi^2 = 72.9$ , df = 12,  $p < 0.0001$ ). Interestingly, for one of these isolates  
313 we also observed a departure from the overall deleterious effect of the pathogenic bacteria on  
314 lifespan: for the PX174 genotype, lifespan in presence of *S. marcescens* ( $8.7 \pm 0.2$  days) was  
315 clearly not lower than in presence of *E. coli* HT115(DE3) ( $8.1 \pm 0.4$  days).

316

317 **Discussion**

318 Aging represents one of the most compelling examples in evolutionary biology of how trait  
319 optimization can have detrimental side effects leading to physiological dysfunction. Here,  
320 using the nematode model *C. elegans*, we have confirmed that microorganisms can shape the

321 host environment in which life-history traits are expressed by showing that distinct bacterial  
322 strains have specific effects on host survival, lifetime fertility and reproductive schedule (Fig.  
323 1). Importantly, we also observed that different strains affect the nematode's reproductive  
324 dynamics and survival in a host-genotype-specific manner, suggesting that such bacteria-host  
325 interactions might affect the evolution of aging in the host.

326 Given the diverse type of interactions that bacteria can establish with *C. elegans* (Diard et  
327 al. 2007; Abalay et al. 2009; Coolon et al. 2009; Baeriswyl et al. 2010; Diaz et al. 2015;  
328 Dirksen et al. 2016; Stuhr and Curran 2020), the observed differences in life-history  
329 responses to specific bacterial strains (Fig. 1) were not entirely surprising. Nevertheless, the  
330 observed bacterial effects did not follow simple expectations based on our knowledge of  
331 strain pathogenicity. For instance, while the pathogenic bacteria *E. coli* IAI1 and *S.*  
332 *marcescens* Db11 had clear detrimental effects on host survival, they did not affect fertility;  
333 similarly, age at first reproduction was unaffected in presence of *E. coli* IAI1. Another  
334 illustration of this are *E. coli* IAI1 and *E. coli* OP50, which imposed the highest and lowest  
335 death rates (Fig. 1A), respectively, but which had similar effects on lifetime fertility (Fig.  
336 1B), in agreement with previous findings (Baeriswyl et al. 2010). Moreover, the recent  
337 evolutionary history of the D00 population might also explain some of the observed patterns:  
338 *E. coli* HT115(DE3), the bacterium used as the food source during the previous 140  
339 generations of laboratory adaptation (Teotónio et al. 2012; Chelo and Teotónio 2013;  
340 Theologidis et al. 2014), led to the highest lifetime fertility observed.

341 The apparent decoupling of the effects of pathogenic bacteria on *C. elegans* reproduction  
342 and survival (also see Diaz et al. 2015) are consistent with the existence of a degree of  
343 specificity in how bacteria interact with their nematode host's physiology, such that  
344 development, metabolism or immunity can be affected independently (Coolon et al. 2009;  
345 MacNeil et al. 2013, Maynard and Weinkove 2020). This may also explain why *C. elegans*

346 survival can differ considerably even between bacteria that are generally regarded as benign  
347 (Brooks et al. 2009; Baeriwisyl et al. 2010; Reinke et al. 2010; Pang and Curran 2014).

348 Our observation that the time-dependence of host population growth and survival differs  
349 among different host isolates in a bacterial-dependent way (Fig. 2) strongly suggests that the  
350 evolution of life-history traits might be subject to microbial modulation. In this context,  
351 bacteria can be regarded as alternative environments where the fitnesses (here given by  
352 population growth rate) of the different genotypes are ranked in an environment-specific  
353 manner.

354 Our results are consistent with several lines of independent evidence showing that in *C.*  
355 *elegans* and/or in other organisms: (i) microbes can influence the expression of life-history  
356 traits (Coolon et al. 2009; Storelli et al. 2011; Diaz et al. 2015); (ii) genotype-by-age effects  
357 are common (Leips et al. 2006; Viñuela et al. 2010); and (iii) genetic correlations between  
358 life-history traits can be subject to modulation by environmental (external) factors (Giesel et  
359 al. 1982; Stearns 1989, 1992; Gutteling et al. 2007; Swanson et al. 2016).

360 A demonstration of a major causal role of microbes as environmental determinants of the  
361 evolution of aging could be obtained with an experimental evolution experiment based on  
362 selection for delayed reproduction (Figure 3), an approach we are currently taking with *C.*  
363 *elegans*. In such an experiment, different bacterial strains might change the genetic  
364 correlations between life-history traits expressed early and late in life. Such an approach  
365 might be able to reveal to what extent the expression of genetic life-history trade-offs and the  
366 evolution of aging, subject to such trade-offs, are constrained by microbial effects.

367

## 368 **AUTHOR CONTRIBUTIONS**

369 M.M., T.F. and I.M.C. conceptualized and designed the project. J.S and I.M.C. collected and  
370 analyzed the data. J.S., M.M., T.F. and I.M.C. interpreted the data and wrote the manuscript.

371

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383

384 **FIGURE LEGENDS**

385

386 **Figure 1.** Different bacteria have specific effects on the reproductive output and survival of  
387 the genetically diverse *C. elegans* D00 host population. **(A)** shows hazard rates imposed by  
388 the benign *E. coli* HT115(DE3) and *E. coli* OP50 strains or the pathogenic *E. coli* IAI1 and *S.*  
389 *marcescens* Db11 bacteria (means plus standard errors SE). **(B)** shows lifetime reproductive  
390 success; and **(C)** shows the reproductive schedule. **(D)** display results for age at first  
391 reproduction, given in hours and days after L1 seed (for comparison with other panels in the  
392 figure). Shown are means plus SE. Letters above bars indicate group assignment based on  
393 post-hoc tests (adjusted  $p$ -value  $< 0.05$ , see Methods), which in **(C)** were performed within  
394 each time period.

395

396 **Figure 2.** Genotype-by-environment (bacteria) interactions affect *C. elegans* population  
397 growth and survival. In **(A)**, population growth rates of the genetic variable D00 population,  
398 measured at the early (72 h) and delayed reproduction period (114 h), reveal bacteria-specific  
399 effects on the temporal dynamics of reproductive output . Similarly, in **(B)**, differences  
400 between population growth at both times are bacterial dependent (color code as in **(A)**), but  
401 specific for each of the five *C. elegans* genotypes (significant three-way interaction, *p*-value  
402 < 0.001). **(C)** shows that mean lifespan depends on the interaction between *C. elegans*  
403 genotype and bacterial type. Letters above symbols show group assignment from significant  
404 post-hoc tests (*p*-value < 0.05) obtained with data for each bacterium independently. Mean  
405 estimates and SE are shown in **(A)** and **(B)**; predicted values are shown in **(C)**. Note the  
406 logarithmic scale of the *y* axis in **(A)** and **(B)**.

407

408 **Figure 3.** Experimental evolution scenarios. Using experimental evolution, different  
409 scenarios of life-history evolution may be obtained by comparison with the effects of delayed  
410 reproduction in the ancestral population. The figure depicts different hypothetical  
411 evolutionary outcomes. The white line shows hypothetical phenotypic values of *C. elegans*  
412 populations feeding on a reference bacterial strain (control); black lines show measurements  
413 taken while feeding on another strain (pathogenic or benign). In **(A)**, host-microbial  
414 interactions (HM) reveal the interplay between reproductive timing and bacterial type for the  
415 genetically diverse ancestral population. These values are displayed as dashed lines in other  
416 plots for comparison. Comparing the ancestral population with derived populations  
417 maintained under control conditions **(B)** or selected for delayed fecundity **(C)** in the presence  
418 of different bacteria should allow understanding the role of *C. elegans*–bacteria interactions  
419 in the evolution of life-history and aging phenotypes **(C)**, by first accounting for adaptation to  
420 the different bacteria **(B** vs. **C**).

421

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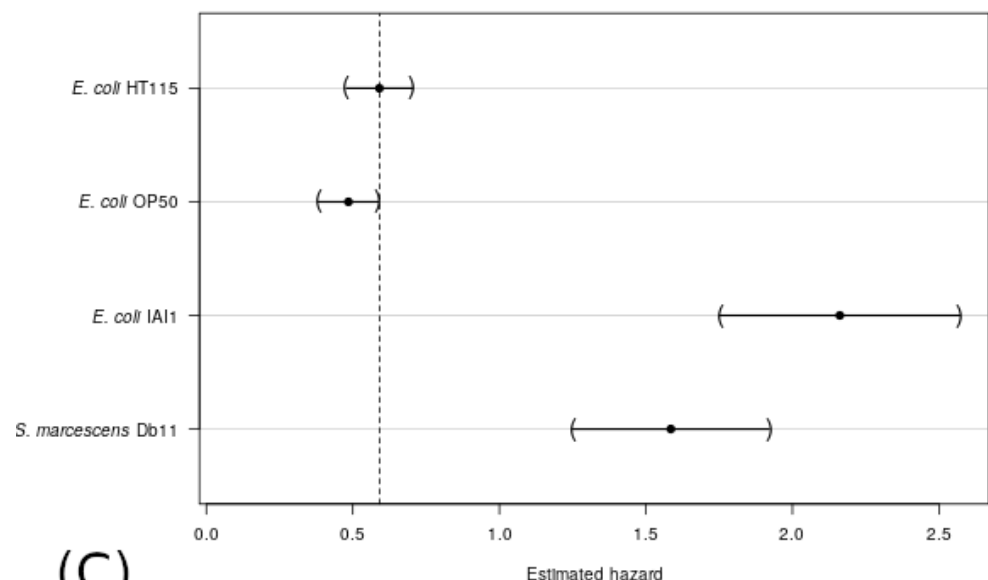
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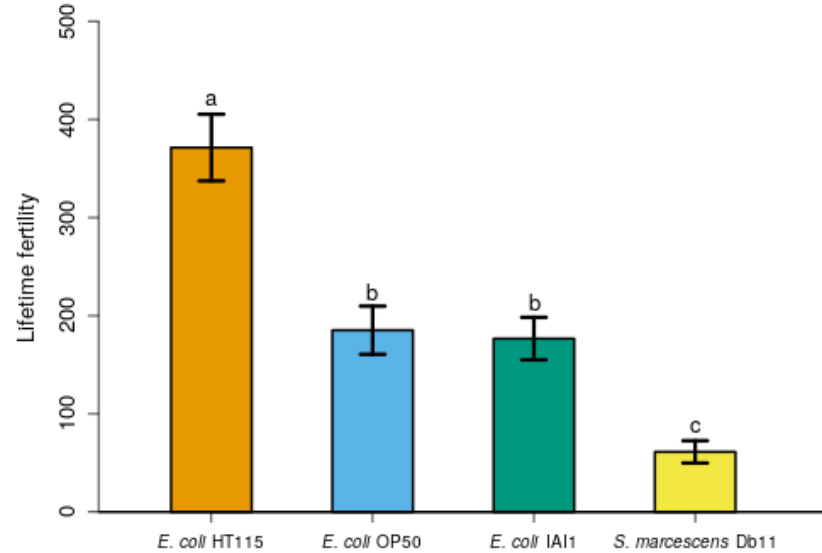
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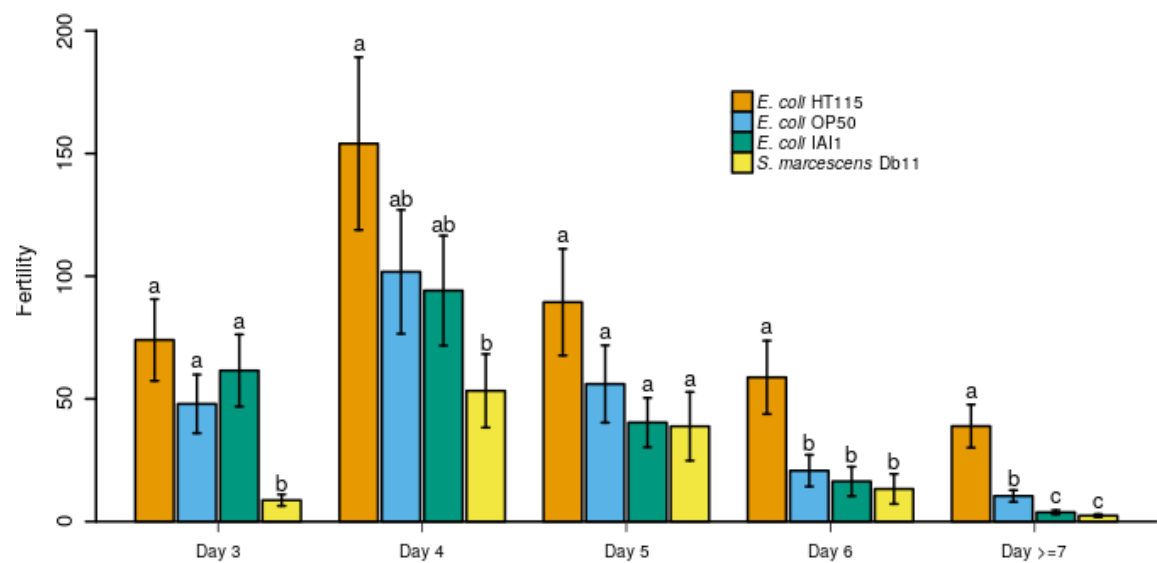
(A)



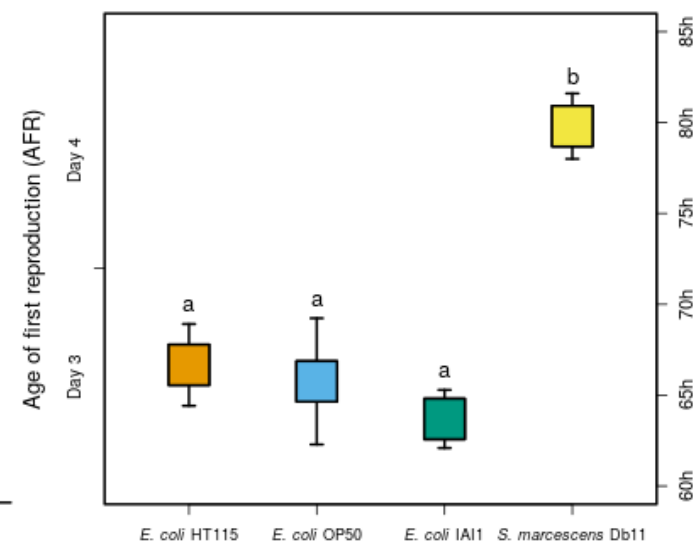
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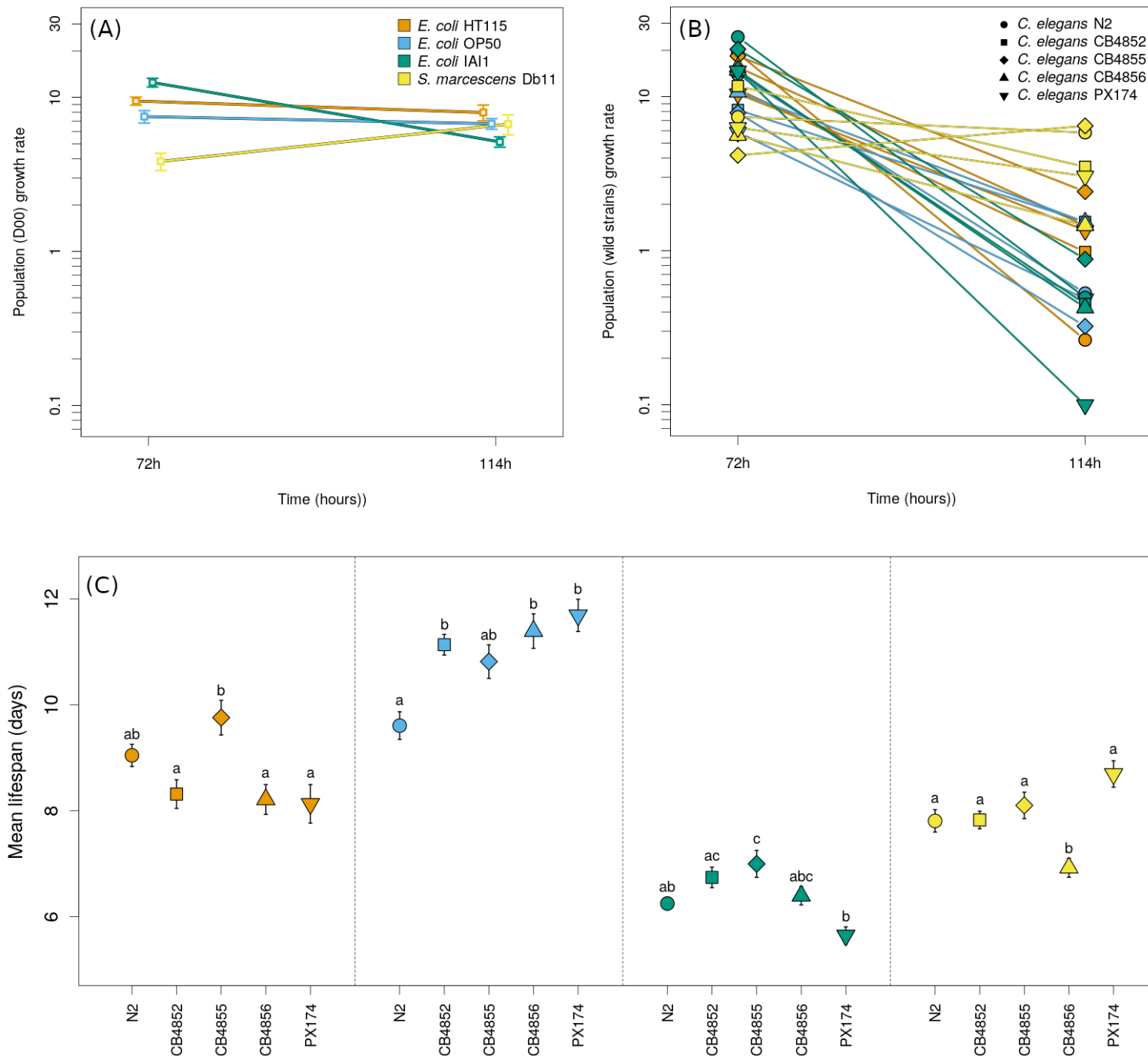


(C)

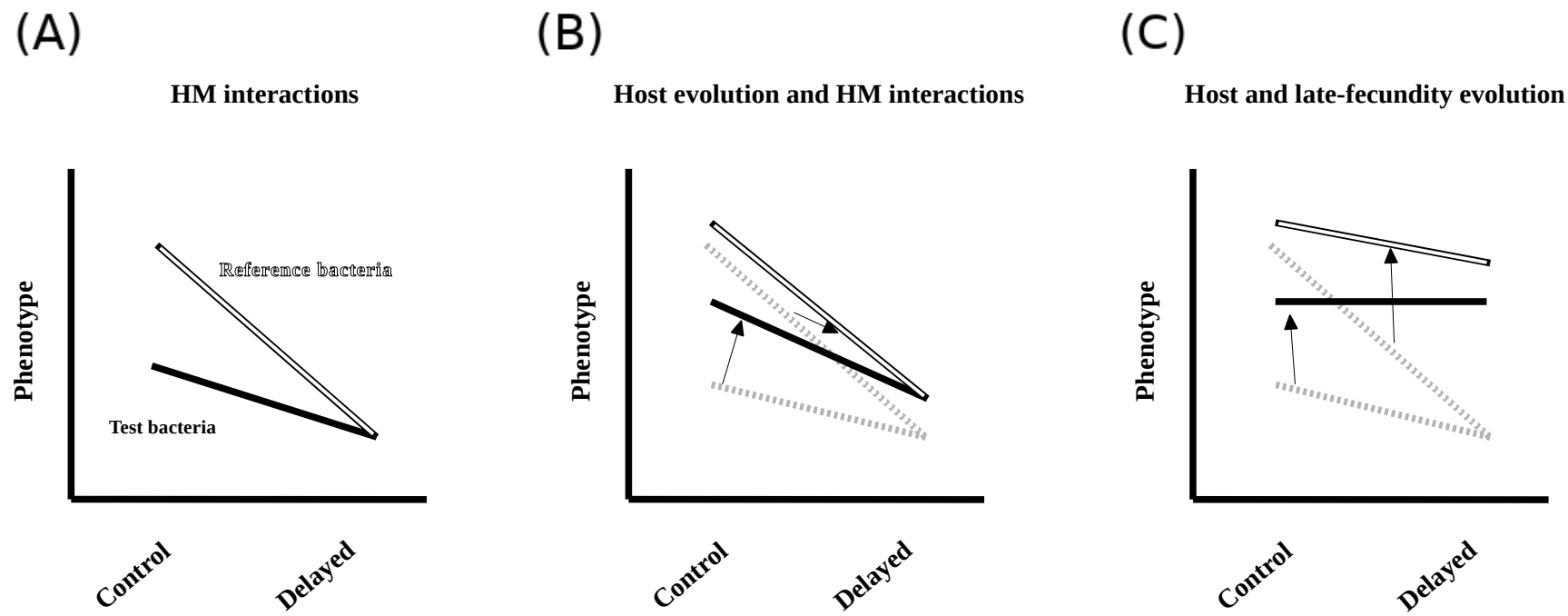


(D)

**Figure 1.**



**Figure 2.**



**Figure 3.**