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1	Variation in diet and microbial exposure shape the performance of the Asian tiger
2	mosquito, Aedes albopictus
3	
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# 16 ABSTRACT

17 Along their life cycle, mosquitoes colonize different ecological niches with various 18 microorganisms and diet sources that likely modulate their performance (i.e. a set of 19 mosquito fitness-related traits). However, which ecological parameters and how their 20 variations modulate mosquito performance is not completely understood. In this study, we 21 used Ae. albopictus surface-sterilized eggs re-associated or not to conventional bacterial 22 microbiota upon a range of diet concentrations and addressed the impact of microbial 23 inoculum and diet concentration variation on several mosquito performance traits. Results 24 showed that mosquito juvenile survival depends on the interaction between bacterial 25 inoculum load and diet concentration in the breeding water. Exposure to bacteria in rearing 26 water shorten larval development time although it impacted larvae survival in an inoculum 27 and diet concentration-dependent manner. Bacterial composition of larval rearing water 28 was mainly structured by the bacterial inoculum concentration, with some Operational 29 Taxonomic Units abundances correlating with larval traits. Ae. albopictus survival, 30 development and bacterial community patterns upon gradients of diet and bacterial 31 inoculum illustrated the complex impact of diet-microbiota interaction on mosquito 32 performance. These findings argue the importance of deciphering host-microbe-33 environment interactions and open promising perspectives to improve Ae. albopictus 34 control measures in the field.

35

## 36 **IMPORTANCE**

Microbiota is increasingly recognized as a driving force of metazoan biology, impacting diverse traits including nutrition, behaviour or reproduction. The microbial impact on host nutrition is among the most studied host-microbe interactions although it remains poorly understood in arthropod vectors like mosquitoes. Here, we manipulated mosquito microbiota using gnotobiology to decipher the impact of bacteria and diet on the Asian

- 42 tiger mosquito, *Ae. albopictus*. These results are key to understand the link between diet
- 43 and bacteria concentration on juvenile mosquitoes as well as carry-over effects in adults.
- 44 They unveil some specific aspects of mosquito-bacteria interactions while opening
- 45 interesting avenues for vector management of this vector of arboviruses.
- 46
- 47 KEYWORDS: Mosquito ecology ; symbiosis ; microbiota ; diet ; Aedes albopictus

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## 49 **INTRODUCTION**

50 From mammals to insects, host-microbiota interactions drive host nutrition, 51 behavior, reproduction or development thereby impacting host fitness (1, 2). However, the 52 ecological determinants governing these symbiotic interactions remain unknown in many 53 systems. During nutritional symbioses, microbial symbionts benefit to host by serving as 54 nutrient source, detoxifying diet, participating in nutrient digestion/intake or supplementing 55 diet with essential molecules (2, 3). Converging evidences show that diet is a driving force 56 of symbiosis, for instance by selecting for diet-adapted microbial communities (4, 5). 57 Advances in the field of symbiotic interactions benefited from the development of 58 gnotobiology (*i.e.* study of hosts with a controlled microbiota) as well as holidic diets (*i.e.* 59 synthetic nutrient source made of pure chemical components). In insects, individuals are 60 deprived from environmentally-acquired microorganisms by surface sterilization of eggs 61 that are put to hatch under sterile environment (6). Individuals can then be re-associated 62 to selected microorganisms to interrogate microbial influence on host traits, under varying 63 (a)biotic conditions such as diet. However, holidic diets and gnotobiotic models are 64 unevenly available among host species (8) although attempts are made to develop 65 gnotobiology and holidic diets in mosquitoes (9, 10).

Compelling evidences from the fruit fly Drosophila melanogaster obtained by 66 67 comparing axenic with gnotobiotic individuals support that bacteria are not only a source of 68 nutrients but establish a diet-dependent nutritional symbiosis to provide their host with essential factors when raised on scarce diets (11–13). However, in mosquitoes the impact 69 70 of the microbiota on mosquito performance under the influence of the diet in a still poorly 71 understood way. Mosquitoes (Diptera: Culicidae) are holometabolous insects with four 72 distinct life stages (egg, larvae, pupae and adult). Larval and pupal stages develop in 73 aquatic breeding sites before winged adults emerge. The breeding site provides larvae 74 with nutrients to complete their development (14-16) and shapes their microbiota (17-19).

75 Diet quantity and composition impact mosquito larval growth, development time and 76 survival (20–25) as well as adult immunity, vector competence and fitness (26–30). Natural 77 larval water habitats contain variable resource inputs that stimulate microbial growth, with 78 the decomposition of detritus by microorganisms releasing nutrients uptake by larvae (31– 79 33). In turn, microbiota also deeply impacts mosquito biology, from larval to adult stage 80 (18). Supplementation of sterile larval rearing water with living bacteria seemed mandatory 81 to reach adult stage (34, 35). However, optimal diet and rearing conditions can lead to 82 complete larval development in absence of environmental microorganisms, as 83 demonstrated in a proof-of-concept study in Ae. aegypti (36). Similarly to what was 84 reported in *D. melanogaster* (37, 38) axenic mosquito larvae present major physiological changes including larval developmental delay, reduced adult size or extended adult life 85 86 span that correlate with specific transcriptomic profiles compare to conventional larvae (36, 87 39, 40). More recently, study of axenic *Ae. aegypti* showed that bacteria in rearing water 88 mediate larval nutrient sensing and growth activation by provisioning larvae with essential 89 vitamins (41, 42). In Ae. aegypti, bacteria-mediated growth-promoting effect on larvae 90 depends on both the bacterial strain and diet type (rat chow or fish food) (43). Microbial 91 supplementation of water as unique nutrient source is not as efficient as synthetic rich diet 92 in promoting larval survival and developmental rate in conventionally reared (*i.e.* non 93 axenic) individuals, with major variations in larval performance being observed according 94 to the microbial strains used (44). This underlines that an interaction between microbiota 95 and diet during larval stage impacts performance in Ae. aegypti larvae, with potential 96 carry-over effects on adult traits (26, 45). Recent data in Ae. aegypti suggest that diet x 97 microbiota interaction occurs as diet concentration impacts bacterial abundance and 98 composition in rearing water and larvae (46). However, additional data are needed to 99 assess if diet and microbiota have similar effects on the biology of other mosquito species, 100 and if and how diet x microbiota interaction shapes mosquito performance.

101 The Asian tiger mosquito Aedes albopictus is an important vector of human 102 pathogens including dengue virus, chikungunya virus or Zika virus (47, 48). This mosquito 103 species thrives notably in urban and suburban environments where females lay eggs in a 104 broad range of breeding sites (51). Field-collected specimens harbor a core bacterial 105 microbiota although variations in bacterial diversity and community structure are observed 106 according to geographic origin, developmental stage or sex (52). Bacteria impact host 107 traits in Ae. albopictus including oviposition site selection (53), sugar feeding (54) or larval 108 development (55). Ae. albopictus juvenile development also depends on diet as larvae 109 present a developmental delay below a given diet concentration that microbiota, including 110 the native intracellular bacterium Wolbachia cannot counterbalance (56). Together, it 111 suggests that microbiota and diet impact Ae. albopictus performance. But to the best of 112 our knowledge, no work generated axenic Ae. albopictus nor study the concomitant impact 113 of diet and microbiota concentration on mosquito performance. Ae. albopictus larvae were 114 successfully deprived from environmental microorganisms, with the exception of 115 intracellular ones such as Wolbachia, while allowing larval development up to the adult 116 stage. In order to disentangle the relative and combined importance of both microbiota and 117 diet on mosquito traits, the performance of gnotobiotic Ae. albopictus was investigated 118 along a gradient of nutrient concentrations. Water microbial community composition and 119 relative abundance were also determined across diet and inoculum conditions.

#### 120 **RESULTS**

121

122 Altered microbiota larvae exhibit a diet-dependent juvenile developmental 123 pattern compared to conventional siblings upon constant bacterial inoculum 124 concentration. Ae. albopictus larvae naturally hosting the intracellular bacterium 125 Wolbachia but deprived from environmental microorganisms were generated by egg 126 surface sterilization (FIG S1), hereafter referred as altered microbiota (AM) larvae. A very 127 low (about 8,600 fold less as estimated by 16S qPCR) residual amount of bacterial DNA in 128 AM rearing water was still detected compare to conventional (CONV) water after egg 129 sterilization (FIG S3). It was composed of bacterial OTUs either poorly abundant 130 (Aeromonas, Dysgonomonas, Pseudomonas) and/or only found in highly diluted inoculum 131 condition (Bacteroides, Stenotrophomonas) of CONV water samples (FIG S3 and FIG 4). 132 Of note, AM larvae were stalled at larval stage only reached adult stage when incubated in 133 darkness (data not shown) recapitulating axenic mosquito larvae development pattern 134 from previous studies (36, 41).

135 The impact of diet concentration in rearing water on larval development was 136 estimated using AM larvae or siblings re-associated upon hatching with a conventional 137 microbial inoculum (CONV) composed of 5 to 6 major bacterial species (FIG S2). AM and 138 CONV larvae were exposed to a gradient of diet from 0.1% to 20% concentration (FIG S4). 139 Diet concentration modulates larvae development depending on the microbial status (Wald  $\chi^2$ ,  $P_{\text{diet x microbial status}} = 3.97e^{-05}$ ). Overall, CONV larvae developed from 12% to 0.5% diet 140 141 concentration whereas AM larvae developed from 20% to 5% although larval viability 142 reached higher values in AM compare to CONV (FIG S4). Indeed, when exposed to a single, low dilution (10<sup>-3</sup>) of inoculum, the proportion of CONV larvae that reached pupal 143 144 stage remained below 25% regardless of diet concentration. In addition, an increase in water turbidity was noticed the day after inoculation with larval death occurring prior fourthinstar.

147 To disentangle the impact of microbiota and diet concentration on mosquito larvae 148 development, a set of three experiments was conducted in which larvae were exposed to a single dose of bacterial inoculum, but more diluted (10<sup>-6</sup>) to limit larval mortality while 149 randomizing microbiota variation by loading a different bacterial inoculum in each 150 151 experiment. Four diet concentrations spanning the viable range for AM and CONV larvae 152 (2, 5, 10 and 12%) were tested. Results showed that larval development differs according 153 to experiments (FIG 1), maybe due to variations in egg batch or composition of microbial 154 inoculum (FIG S2). When controlling for random experiment effect, larvae-to-pupae 155 viability depended on the interaction between microbial status and diet concentration (Wald  $\chi^2$ ,  $P_{\text{diet x microbial status}} < 2.2e^{-16}$ ) (FIG 1A). AM larvae survival from 5 to 12% diet 156 157 concentration was similar but lower than at 2% whereas CONV larvae viability at 2 and 5% 158 diet concentration was similar but higher than at 10 and 12% (FIG 1A). Unlike larvae, pupae-to-adult viability was not impacted by these two variables (Wald  $\chi^2$ ,  $P_{diet} = 0.16$ , 159 160 P<sub>microbial status</sub> = 0.31, P<sub>diet x microbial status</sub> = 0.51) (FIG 1B). Larval viability was higher in AM 161 compared to CONV except at the lowest diet concentration of 2% (FIG 1A). Among the 162 larvae that reached pupal stage, we measured the time (in days) needed to reach 50% of 163 the final number of pupae (Day<sub>50</sub>) as a proxy of development time. As observed for larvae-164 to-pupae viability, mosquito juvenile development time depended on the interaction between microbial status and diet concentration (Anova,  $P_{\text{diet x microbial status}} < 2.2e^{-16}$ ) (FIG 165 166 1C). Overall, CONV larvae always developed faster (about 7 days) than AM counterparts. 167 The Day<sub>50</sub> of CONV larvae at 2, 5 and 10% diet concentrations were similar but smaller 168 than at 12%, whereas the Day<sub>50</sub> of AM larvae at 2% diet concentration was higher than at 169 5%, which in turn was higher than at 10 and 12% (FIG 1C).

170 No significant difference was observed in the proportion of adults from each sex according to microbial status, diet concentration or their interaction (Wald  $\chi^2$ ,  $P_{diet} = 0.72$ , 171  $P_{\text{microbial status}} = 0.50, P_{\text{diet x microbial status}} = 0.067)$  (FIG 2A). Adult wing length was significantly 172 173 impacted by the interaction between microbial status and diet concentration (Anova, P<sub>diet x</sub> microbial status = 0.009) (FIG 2B). CONV females presented larger wings compare to males, 174 175 and within each sex, wing length was not different according to diet concentration. In AM 176 conditions, females displayed higher wing length compared to males except at 2% diet concentration, with all males presenting similar wing length regardless of diet 177 178 concentration. Within AM females, a similar wing length was measured at 10 and 12%. 179 However, AM females were larger at 12%, compared to 5 and 2% diet concentrations.

180

Bacterial load and diet concentration in larval rearing water are strong 181 182 predictors of Ae. albopictus juvenile performance. We showed that at constant initial 183 load, bacteria promote larval performance depending on diet concentration. In nature, 184 microbial load and diet concentration are likely to vary in larval breeding sites. Therefore, 185 we asked if and how concurrent variation in bacterial inoculum and diet concentration could shape mosquito juvenile performance. Within a single experiment, CONV larvae 186 were exposed to three different dilutions of inoculum  $(10^{-4}, 10^{-6} \text{ and } 10^{-8})$  and three diet 187 188 concentrations (1, 5 and 12%) prior to measure larval viability, development time and 189 microbiota composition. Concentrations of inoculum and diet were chosen according to 190 previous conditions showing the most contrasted effects on larval performance (FIG 1). 191 Three independent batches of inoculum (B1, B2 and B3) were used. Analysis of the 192 bacterial community variance of the water of each batch showed differences in bacterial 193 composition (FIG S2). When controlling for inoculum batch variation, results showed that 194 larval viability is impacted by the interaction between inoculum and diet concentration (Wald  $\chi^2$ ,  $P_{\text{diet x inoculum}} = 6.86e^{-7}$ ) (FIG 3A). At 10<sup>-4</sup> inoculum dilution, larval viability is the 195

196 lowest overall, with 1 and 5% diet concentrations presenting similar but higher viability than 12%. Larval viability is higher at  $10^{-6}$  compared to  $10^{-4}$  inoculum dilution, except for 197 the 12% diet concentration. At 10<sup>-6</sup> inoculum dilution, no difference in larval viability was 198 199 found according to diet concentration although the pattern shows a decrease in larval viability as diet concentration increased, as previously observed (FIG 3A and 1A). At 10<sup>-8</sup> 200 inoculum dilution, larval viability pattern shows an opposite trend compared to 10<sup>-6</sup>, with 201 202 larval viability increasing with diet concentration. Viability at 1% and 5% diet concentration 203 is similar but lower compare to 12% that displays a significantly higher viability compare to 204 all the conditions tested (FIG 3A). The median larval development time depends on both 205 microbial inoculum and diet concentration although the interaction was not significant (Wald  $\chi^2$  on data transformed by 1/x<sup>2</sup> function,  $P_{diet} = 0.018$ ,  $P_{inoculum} = 1.10e^{-14}$ ,  $P_{diet x}$ 206  $_{inoculum}$  = 0.11). Day<sub>50</sub> increased upon 10<sup>-4</sup> and 10<sup>-8</sup> inoculum dilution compared to 10<sup>-6</sup>, 207 although it varied with the diet concentration. The shortest Day<sub>50</sub> was measured at 10<sup>-6</sup> 208 inoculum and 5% diet concentration, being significantly lower compared to 10<sup>-4</sup> / 12% and 209  $10^{-4}$  / 1% conditions as well as all  $10^{-8}$  diet concentrations (FIG 3B). Taken together, our 210 211 results indicate that Ae. albopictus larval performance cannot be fully understood without 212 considering the combined impact of diet and bacterial inoculum, that have carry-over 213 effects on adult wing length. While specific larvae survival patterns were observed upon 214 diet x microbiota interaction, the development time seems less constraint by this 215 interaction with a trend for bacteria-associated to develop faster overall especially upon 216 higher microbial inoculum although some diet concentration specific variations still 217 remained.

218

Initial bacterial inoculum and diet concentration shape aquatic habitat
 microbiota composition with bacterial taxa significantly correlated with Ae.
 *albopictus* performance. Eleven bacterial OTUs were identified at the genus level (with

222 >5% relative abundance) within the rearing water that served as microbial inoculum for 223 CONV larvae (FIG S2). Six out of these 11 OTUs (Brevundimonas, Delftia, 224 Flavobacteriaceae, Pseudomonas, Sphingobacteriaceae and Sphingobacterium) were 225 found five days later in the water of CONV larvae (FIG 4A). Visualisations of the relative 226 abundances of the 26 most predominant genera suggested a strong structuration of 227 bacterial microbiota according to initial inoculum concentration, and to a lesser extent, diet 228 concentration (FIG 4A). Beta diversity (microbial diversity between samples) of water 229 samples was represented using non-metric multidimensional scaling (NMDS) with Bray 230 Curtis dissimilarity as the distance metric (FIG 4B-C). When assessing the effect of diet concentration (1, 5 or 12%), inoculum dilution ( $10^{-4}$ ,  $10^{-6}$  or  $10^{-8}$ ) and batch of inoculum 231 (B1, B2, B3), the triple interaction was significant (PERMANOVA, Pinoculum x diet x batch = 232 0.005,  $R^2 = 0.0047$ ) (FIG 4B). When trying to disentangle the effect of individual factors on 233 234 bacterial structure, inoculum dilution had a strong impact when plotted against diet 235 concentration (FIG 4C). These data indicate that complex interactions occur between diet 236 concentration and that the initial composition and abundance of bacterial inoculum shape 237 community structure in larval rearing water. Overall, the bacterial microbiota is 238 homogenous upon when inoculating a high bacterial load whereas a lower amount of 239 inoculum triggers a shift toward a more variable bacterial community in rearing water. 240 Correlation matrix between OTUs abundance (in sequencing reads) and larval traits, including development time (median  $Day_{50}$ ), inoculum dilution ( $10^{-4}$ ,  $10^{-6}$  and  $10^{-8}$ ), diet 241 242 concentration (1, 5 and 12%) or larval viability (in %) was performed (FIG 4D). Twenty-two 243 OTUs displayed a significant correlation (Spearman, P < 0.05) with a correlation coefficient 244 below -0.4 or above 0.4 that we used as threshold. Only one OTU, assigned to 245 *Bacteroides* genera, correlated with  $Day_{50}$  (FIG 4D). Most of the OTUs correlating in 246 relative abundance with one or more larval traits had a low relative abundance with the 247 exception of Sphingobacterium (relative abundance correlated with larval viability),

*Chryseobacterium* (abundance correlated with both larval viability and inoculum dilution)
and *Bosea* (abundance correlated with larval viability and diet concentration) (FIG 4A&D).
Details of significant correlation patterns between OTUs relative abundance and larval
traits, diet and inoculum concentration for OTUs with the highest relative abundance are
shown in FIG S5.

253

## 254 **DISCUSSION**

255

256 Our study demonstrated that microbiota and diet in aquatic larval habitat drive 257 ecological performance of Ae. albopictus larvae with carry-over effects on adult 258 mosquitoes. This study is the first, to the best of our knowledge, to address Ae. albopictus 259 development using larvae deprived from external microorganisms (AM) and siblings re-260 associated with conventional bacteria (CONV). AM larvae of Ae. albopictus developed up 261 to the adult stage, as observed in axenic from Aedes aegypti (36) or Drosophila 262 melanogaster (11). If the presence of living but non-cultivable bacteria cannot be ruled out, 263 AM larvae delayed development time and shorter wing length recapitulated developmental 264 phenotypes of axenic Ae. aegypti (36). More broadly, these results resonate with studies in 265 axenic flies and mice reinforcing the role of microbiota as a promoter of juvenile 266 development in metazoan (57, 58).

Our results showed that diet and bacteria mostly impact larvae-to-pupae transition. Bacteria from rearing water promoted larval development upon nutrient scarcity but, above a given nutrient concentration, impaired larval viability while the opposite trend was shown in AM larvae. Upon high nutrient concentration, bacterial presence seemed toxic for mosquito larvae although at low diet concentration, it promoted larval development through an increased viability and a shorter development time. Together, it underlies the importance of bacteria and diet interaction on mosquito fitness in *Ae. albopictus*. Bacterial 274 growth in rearing water depends on the strain and diet considered although total bacterial 275 load increases with diet concentration (43, 46). Maintenance of a bacterial homeostasis in 276 mosquitoes is under tight immunological control (65). Therefore, we can hypothesize that 277 diet concentration mediates an increase in water bacterial load that disrupt bacterial 278 homeostasis, leading to a fitness decrease in mosquito.

279 Despite larval development time depended on diet x microbiota interaction, CONV 280 development time remained around 6 to 7 days with only a small increase at the highest 281 diet concentration tested. This suggests that within a range of diet concentration, bacteria 282 from rearing water can buffer nutrient scarcity that maintains a short larval development 283 time as in Drosophila (37). However, development time of CONV mosquito larvae was 284 steadily ~5 days shorter than AM siblings even at the highest diet concentration. 285 conversely to *Drosophila* for which this developmental delay is observed in nutrient-poor 286 conditions but abolished on rich diet (11). Although the use of microorganisms as nutrients 287 by mosquitoes is generally pointed out (66), our results suggest that, beyond a food 288 source, bacteria are actively providing Ae. albopictus with compounds as supported by the 289 identification of folate and riboflavin provisioning by bacteria as an essential mechanism 290 for mosquito development in Ae. aegypti (41, 42). Further experiments comparing 291 metabolically inactive (e.g. heat-killed) and active bacterial exposure in axenic larvae will 292 provide additional evidences to understand host-microbe interactions in mosquito.

A previous study showed that diet concentration impacts microbiota composition in *Ae. aegypti* larvae and adults (46). Our results unravelled a more complex interaction involving both the diet concentration and the bacterial load inoculum in shaping bacterial communities in larval water. Results showed that nutrient-rich water samples colonized with a highly concentrated inoculum have a strongly similar bacterial community compared to nutrient poor samples exposed to a diluted bacterial inoculum. Several OTUs were preferentially retrieved in water following exposure to a highly diluted (10<sup>-8</sup>) inoculum 300 supporting the idea of a more random bacterial community construction upon low initial 301 bacterial dose. Interestingly, the four OTUs that correlated to diet concentration were more 302 abundant at the lowest diet concentration suggesting that nutrient density and/or 303 competition could impact their load upon higher diet concentration. Notably, one OTU from 304 the Comamonadaceae family with a higher abundance in rearing water upon low (1%) diet 305 concentration displayed the same abundance/diet concentration pattern in Ae. aegypti 306 (46). OTU from genus Bosea was more abundant when diet concentration was low (1%), 307 and its abundance correlated with larvae viability although not in a linear way. Present in 308 sylvatic Ae. aegypti larval habitats, Bosea can trigger a decrease in adult lifespan upon 309 larval exposure (45). Chryseobacterium and Sphingobacterium abundance negatively 310 correlated with larvae viability. If Chryseobacterium was often encountered in mosquito 311 microbiota and restored larval development in mono-association (67), our results suggest 312 that its abundance is key for Ae. albopictus performance. Sphingobacterium was 313 associated with a developmental delay and shorter wing length in Ae. aegypti (43). Our 314 data suggest that it can also impair larval viability in a density-dependent manner. 315 Altogether, our data indicate a complex interplay between bacterial founders and nutrients 316 load that shape bacterial community structure and impact mosquito performance. While it 317 is known that given bacterial strains can compensate for mosquito larvae auxotrophies 318 notably for vitamins (41, 42), data are still lacking to fully understand complex multipartite 319 host-microbe interactions in mosquitoes. For instance, Lactoplantibacillus plantarum and 320 Acetobacter pomorum triggered a metabolic cooperation that complemented Drosophila 321 larval auxotrophies through metabolites that are not produced by bacteria in a mono-322 association context (68). This underlines that diet-microbiota interaction is key as diet 323 selects for diet-adapted strains, influences the growth of microorganisms and cooperation 324 in the niche which in turn impacts host performance as shown in fly and mammals (4, 5, 325 69). The situation can be further complexified if we consider other microorganisms from

*Ae. albopictus* microbiota that can be directly involved in mosquito metabolism as recently uncovered for fungi (70) or indirect effects such as arbovirus-mediated modulation of bacterial diversity in adult mosquitoes (71).

329 The development of gnotobiology (76) and holidic diets coupled to in-depth 330 transcriptomic and metabolic analysis will help toward the understanding of mosquito 331 interaction with their biotic and abiotic environment (41). Interestingly, recent work showed 332 opposite conclusions on the impact of microbiota on larval transcriptome, advocating for 333 additional studies that compare transcriptomes of axenic and mono-associated larvae with 334 field-derived bacterial isolates, upon diet and bacterial inoculum variations (39, 40). More 335 broadly, it has potential major applications for alternative vector control strategies for 336 instance by manipulating mosquito oviposition behaviour through bacteria and diet in the 337 rearing water in order to limit mosquito density (75). Altogether, this work provides a better 338 understanding of the ecological determinants of symbiosis in a medically relevant 339 organism while opening interesting avenues for alternative vector control strategies. It also 340 uncovers a fragile equilibrium between diet, bacteria and mosquito fitness, opening 341 questions about the genetic and environmental basis of mosquito traits notably involved in 342 vectorial capacity.

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#### 344 MATERIAL AND METHODS

345

Mosquito colony maintenance.  $F_9$  and  $F_{10}$  from an *Ae. albopictus* colony (referred as VB) established from field mosquitoes collected in 2018 in Villeurbanne and Pierre-Bénite (France) were used. Larvae were maintained at 26°C with dechlorinated water and Tetramin fish food. Adults were raised at 28°C, 80% relative humidity, 16:8 light:dark photoperiod in mass rearing. Females were blood fed on mice in accordance with the Institutional Animal Care and Use Committee from Lyon1 University (Apafis #31807-2021052715018315). Egg papers were stored at 28°C for up to two months.

353

**Diet plugs preparation.** Sterile agar diet plugs of various concentrations were prepared. Finely grinded tropical fish flakes (TetraMin) were mixed with sterile water at final concentrations of 20%, 12%, 10%, 8%, 5%, 2%, 1%, 0.5% and 0.1% (w:v). Bacteriological agar (Conda) was added at a final concentration of 1.6% (w:v). Diet suspensions were autoclaved (120°C, 20 min), poured into 90 mm petri dishes (20 mL per dish) and stored at 4°C for up to 3 days. Die-cut of 0.6 g agar food plugs using a 15-mL sterile tube (Falcon) allowed a precise control of food quantity.

361

362 Preparation of conventional microbial inoculum. Larvae hatched from surface-363 sterilized eggs were re-associated with a microbiota representative of the insectary-reared 364 siblings to obtain conventional (CONV) larvae. Briefly, VB eggs hatched were in hatched 365 for 2h at -20 ATM in a vacuum chamber then 200 first instar larvae were transferred in 24 366 x 32 x 9 cm plastic trays with 1.5 L of dechlorinated water. After 7 days at 26°C with 0.1 g / 367 tray every two days of grinded fish flakes (Tetramin) supplemented with yeast extract 368 (Biover) (3:1 w:w), 50 mL of water (without larvae) from 3 independent trays were pooled 369 to constitute the CONV inoculum. When testing for inoculum batch effect, 3 batches were prepared from independent egg papers. To prepare CONV larvae first instars from surface-sterilized eggs were exposed immediately upon hatching to CONV inoculum at selected dilution (from 10<sup>-3</sup> to 10<sup>-8</sup>) in sterile water (Gibco). Bacterial composition of CONV inoculum is provided (FIG S2) while no eukaryotic microorganisms were detected by 18S PCR.

375

376 Conventional and altered microbiota mosquito production. After hatching 377 larvae acquire their microbiota through the ingestion of microorganisms present on the egg 378 surface or in the rearing water. Ae. albopictus larvae with altered microbiota (AM) were 379 obtained by surface sterilization of mosquito eggs (FIG S1). In 6-well plates, a 0.6 g agar 380 food plug at the selected concentration was added per well with 5 mL of sterile water (for 381 AM condition) or 5 ml of CONV inoculum at selected dilutions (for CONV condition). A total 382 of 3 larvae / well was added and plates were incubated at 28°C in complete darkness for 383 up to 22 days. The well is the biological replicate unit and 6 to 18 wells were tested for 384 each condition. For each experiment, agar plugs and PBS (in which surface-sterilized 385 larvae hatched) were incubated on LBm agar for 7 days at 30°C to control for the absence 386 of cultivable contaminants. Wells were observed daily and discarded in presence of turbid 387 water. As a majority of microorganisms are not cultivable, random wells were observed for 388 each plate under the microscope to assess the absence of microbial contaminants.

389

Larvae viability and development time. Presence of pupae was recorded daily at fixed hours. Larvae-to-pupae viability represents the percentage of larvae that reached pupal stage. Development time represents the time (in days) needed to reach 50% of the total number of pupae (Day<sub>50</sub>). On the day of emergence, pupae were individually transferred with ~300  $\mu$ L of rearing water in a sterile tube for emergence. Adults were sexed before storage together with water of emergence at -80°C. bioRxiv preprint doi: https://doi.org/10.1101/2022.11.16.516852; this version posted November 17, 2022. The copyright holder for this preprint (which was not certified by peer review) is the author/funder. All rights reserved. No reuse allowed without permission.

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Wing length measurement. Wing length is a proxy for *Ae. albopictus* adult performance (79). Adults stored at -80°C were thawed and both wings were dissected under a Leica M80 stereomicroscope. Wings were included Eurapal (Roth) on a 10-wells, epoxy-coated glass slide (Labelians). Slides were photographed at 20x magnification with Leica MC170 HD camera. Images were analysed with ImageJ (version 2.1.0/1.53c). Wing length was measured between the intersection of the second and third vein and the intersection of the seventh vein with the wing border using ImageJ (80).

404

DNA isolation from larvae rearing water. Five days post exposure with CONV 405 406 inoculum, 500 µL of rearing water were stored at -20°C prior to total DNA isolation. 407 Samples were centrifuged for 20 min, 4°C, at 17,000 G and the pellet containing the 408 microorganisms was used to extract DNA with the DNeasy Blood & Tissue kit as 409 recommended (Qiagen). DNA concentration was estimated by Qubit dsDNA HS kit 410 (Thermo Fisher scientific) and samples were stored at -20°C. A blank control was 411 performed by using only DNA lysis buffer to control DNA microbial contaminations arising 412 from kit reagents or the environment during DNA isolation.

413

414 **Quantitative PCR of 16S DNA.** The 16S DNA load was measured by qPCR using 415 the Itaq SYBR green supermix kit (Bio-Rad), 784F (5'-AGGATTAGATACCCTGGTA-3') 416 and 1061R (5'-CRRCACGAGCTGAC') primers and 5 ng of template DNA isolated from 417 water samples. The 16  $\mu$ L reaction comprised 0.48  $\mu$ L of each primer at 10  $\mu$ M, 8  $\mu$ L of 418 Master mix and 5.04  $\mu$ L of PCR grade water. After a single denaturation step at 95°C for 3 419 min, a two-step amplification was performed including 10 s at 95°C followed by 30 s at 420 60°C, for 40 cycles on a Bio-Rad CFX96 machine. The number of 16S copies per  $\mu$ L was 421 calculated using serial dilutions of an *Acinetobacter* PCR amplicon from 10<sup>8</sup> to 10 422 copies/μL.

423

424 Microbial amplicon sequencing. Identification of bacteria and eukaryotes was 425 based on PCR amplification of a ~280 bp fragment of the V5-V6 variable region from the 426 16S rRNA gene (52) and a ~430 bp fragment of the 18S gene (81). Duplicate PCR for 427 each sample were done using the 5X Hot BIOAmp (Biofidal, Vaulx-en-Velin, France – 428 http://www.biofidal.com). Duplicates PCR products were pooled and 5 µL were separated 429 by electrophoresis on 1.5% agarose gel supplemented with 2.5 µL of clear sight DNA stain for 17 min at 100V. All the 18S PCR were negative. For the 16S, all PCR were positive at 430 431 expected size. A total of 188 libraries from 16S amplicons were constructed, including 432 controls. Sequencing was done on Illumina MiSeq (2x300 bp, paired-end) at Biofidal. In 433 total, 20,764,855 reads were obtained and demultiplexed. Sequence quality control and 434 analysis were carried out using the FROGS pipeline (82) as described (83). Taxonomic 435 affiliation was performed with SILVA database 138.1 for bacteria (84) with Mothur pipeline 436 (85) at a 80% minimum bootstrap using a naïve Bayesian classifier (86). Sequences were 437 grouped into operational taxonomic units (OTUs) by clustering at 97% similarity. No OTUs 438 were detected in the blank extraction or PCR controls. To compare samples, normalization 439 was performed at 11,780 sequences. A total of 184 operational taxonomic units (OTUs) 440 was obtained. OTUs with a relative abundance less than 10 times greater than that 441 observed in the negative control were removed (52). All FastQ files were deposited in the 442 EMBL European Nucleotide Archive (https://www.ebi.ac.uk/ena) under the project 443 accession number PRJEB57586. The small amount of residual bacterial DNA detected in 444 water from altered microbiota larvae was guantified by gPCR, and taxonomic identity and 445 relative abundance of the 5 OTUs detected shown (FIG S3).

447 Statistical analysis. Analyses and graphical representations were done on R (http://www.r-project.org/). Larvae- and pupae-to-adult viability (binary response variable) 448 449 were analysed by generalized linear mixed-effects (GLMM) (87). GLMM with a binomial 450 distribution and a probit link function were fitted by maximum likelihood (Laplace 451 approximation). The development time (Day<sub>50</sub>) and wing length (in mm) were analysed 452 using linear mixed-effects (LMM) models fit by restricted maximum likelihood while 453 controlling for normal distribution of the residuals. Microbial status and diet concentration 454 in interaction represented fixed effects while experiment or batch of inoculum were 455 included as random effect. The GLMM and LMM models were conducted under Ime4 456 package version 1.1-25 (88). The inference of explanatory variable on variations of the response variables was tested with a Wald  $\chi^2$  test or an ANOVA for binomial and 457 458 continuous data respectively. Post-hoc comparisons after GLMM/LMM were performed 459 with *emmeans* package version 1.5.2-1 (89) to estimate pairwise differences between diet 460 concentrations and microbial status using Tukey-HSD test with p-value correction for 461 multiple comparisons. Adonis-ANOVA and Non-Metric Multidimensional Scaling ordination 462 were performed with the ade4 and vegan packages (90, 91). Non-linear Spearman 463 correlations were performed with the Hmisc package (92). Other R packages were used 464 for data organization and representation such as *plyr* version 1.8.6 (93) and *gaplot2* 465 version 3.3.2 (94).

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# 707 Legend of figures

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710 FIG 1 Juvenile development pattern of Ae. albopictus larvae upon a diet gradient in 711 presence or absence of bacteria. Development pattern of CONV (black) and AM (blue) 712 larvae at four diet concentrations (2, 5, 10 and 12%). Three independent experiments (#1 713 to #3 as indicated by color code) were performed, with one to three 6-well plates per 714 experiment (3 larvae per well) for each microbial status and diet concentration 715 combination. (A) Larval viability expressed as the proportion (in %) of mosquito larvae that 716 reached pupal stage. Each dot represents the mean viability per 6-well plate. (B) Pupal 717 viability expressed as the proportion (in %) of mosquito pupae that reached adult stage. 718 Each dot represents the mean viability per 6-well plate. (C) Larval development time into 719 pupae is represented as the time (in days) needed to reach 50% of the final number of 720 pupae ( $Day_{50}$ ). Different letters indicate statistically significant viability following Tukey-721 HSD *post-hoc* pairwise comparison.

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723 FIG 2 Adult sex ratio and wing size upon a diet gradient in presence or absence of 724 bacteria. Adults derived from CONV or AM larvae reared at different diet concentrations (2, 725 5, 10 and 12%) were sorted by sex and individual wings length was measured. Three 726 independent experiments (#1 to #3) were performed, with one to three 6-well plates per 727 experiment (3 larvae per well) for each microbial status and diet concentration 728 combination. The number of adults available varied according to the juvenile mortality rate 729 of each combination. (A) Proportion of female and male adult mosquitoes. The number of 730 individuals (n) for each condition is indicated. (B) Adult wings length (in mm). Each dot 731 represents the mean length of both wings for an individual. From 5 to 17 individuals were 732 analysed depending on the sex, microbial status and diet concentration. Different letters 733 indicate statistically significant viability following Tukey-HSD post-hoc pairwise 734 comparison.

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**FIG 3** Juvenile performance of *Ae. albopictus* upon concomitant variation of bacterial inoculum and diet concentration. CONV larvae exposed to three microbial inoculum concentrations (10<sup>-4</sup>, 10<sup>-6</sup> and 10<sup>-8</sup>) issued from three independent batches (B1, B2 and B3, color coded) and three inocula and diet (1, 5, and 12%) concentrations were monitored up to pupal stage. (A) Larval viability expressed as the proportion (in %) of larvae that reached pupal stage at different inoculum and diet concentrations. Each dot represents the mean viability per 6-well plate. (B) Larval development time expressed as the time (in days) needed to reach 50% of the final number of pupae. Each dot represents
the mean Day<sub>50</sub> per well. Different letters indicate statistically significant viability following
Tukey-HSD *post-hoc* pairwise comparison. For each batch of inoculum, one 6-well plate
was prepared (3 larvae per well) for each inoculum and diet concentration combination.

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748 **FIG 4** Bacterial community structuration in larval rearing water upon bacterial inoculum 749 and diet concentration gradients. (A) Relative abundances (in %) of bacterial operational 750 taxonomic units (OTUs) at the genus level for each combination of inoculum and diet 751 concentration, for the three batches of microbial inoculum (B1, B2, B3) used. OTUs 752 representing less than 5% in relative abundance were grouped (red, <5% Rel. ab.). (B,C) 753 Non-metric multidimensional scaling (NMDS) analysis of the Bray-Curtis dissimilarities as 754 a function of diet concentration and inoculum dilution (B) or the interaction between batch 755 of inoculum and inoculum dilution (C). (D) Heatmap representing Spearman correlation 756 coefficient between phenotype of interest (development time (Day<sub>50</sub>), inoculum dilution, 757 diet concentration or Larvae-to-pupae viability (L to P)) as a function of OUT abundance. Only OTUs with significative p-value ( $<1e^{-6}$ ) for correlation analysis and a correlation 758 759 coefficient < -0.4 or > 0.4 are shown. When genus level information was not available, the 760 family (F) or order (O) level is indicated.

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FIG S1 Protocol for surface sterilization of Ae. albopictus eggs. Protocol was adapted from 762 763 previous work (36) Ae. albopictus eggs (<2 months old) were used. Careful visual 764 inspection of egg papers under binocular was conducted prior to each experiment to avoid 765 concave eggs (as a proxy for unfertilized eggs) or egg papers with mold or mites. Within 766 each experiment, at least three different batches of embryos from the same generation but 767 laid at different days were used. Under safety cabinet, eggs were rinsed twice by dipping 768 the paper in sterile water to discard debris. Egg papers were then soaked in a petri dish 769 containing 70% ethanol solution (in sterile water) for 5 min then transferred in a 50-mL 770 falcon tube containing 30 mL of sodium hypochlorite (3% active chlorite) supplemented 771 with 4 mg/mL ampicillin for 5 min. Sodium hypochlorite immediately detached the eggs 772 from the paper, that was removed using clean forceps. Within the 5 min, eggs quickly 773 settled at the bottom of the tube allowing the complete removal of sodium hypochlorite 774 without the need of centrifugation. Eggs were resuspended in 25 ml of 70% ethanol and 775 incubated for 5 min. We ensured that all eggs remained immersed in ethanol and gently 776 agitated the tube to allow a complete contact of the eggs with ethanol solution. Eggs were 777 rinsed three times in sterile water (Gibco) for 5 min then 30 mL of sterile 1X PBS was added. The tube was closed with a 0.2 µm filtered cap and transferred in a vacuum
chamber outside the cabinet for 40 min at -20 ATM to allow hatching. The sterile first instar
larvae were transferred under the safety cabinet in a sterile petri dish and immediately
transferred in 6-well plates using a P1000 pipette according to the experimental design.

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**FIG S2 Taxonomic identity and** relative abundance of bacterial OTUs in water inoculum for CONV larvae. Relative abundance (in %) of operational taxonomic units (OUT) of bacteria at genus level (or family (F) level) in water of conventionally reared larvae that served as inoculum for CONV condition. Two DNA samples (S1, S2) per batch of inoculum (Batch 1, 2 and 3) were sequenced except for Batch 3 (only one sample). The two OTUs that were isolated by culture-dependent approach on LBm media are shown (\*). OTUs representing less than 5% in relative abundance were grouped (red, <5% Rel. ab.).

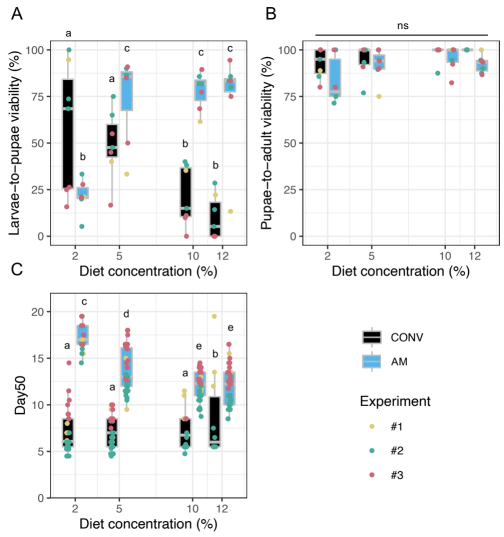
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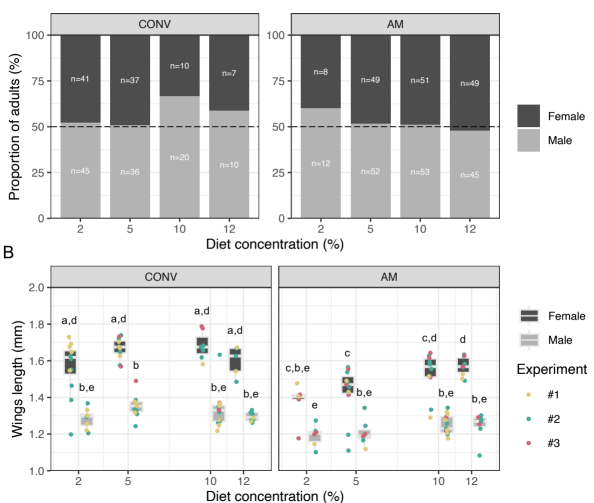
FIG S3 Bacteria composition and relative abundance of altered microbiota larvae rearing 791 792 water DNA samples. (A) Quantification of 16S copies number (log<sub>10</sub>) in DNA isolated from 793 water of conventional (CONV) and altered microbiota water samples after five days at 28°C. CONV samples (Cq values from 14 to 16, mean 5.5x10<sup>6</sup> copies) originated from 794 water containing larvae exposed to 10<sup>-8</sup> dilution of inoculum and incubated with 5% diet 795 concentration. The altered microbiota DNA samples (Cq values from 28 to 29.4, mean 796 6.39x10<sup>2</sup> copies) originated from water with diet but no larvae (S1, S2), water with larvae 797 798 and diet (S4, S5) or water without larvae nor diet (S3). A blank DNA isolation control (Cq 799 34.9, 10.5 copies) was done by replacing water with DNA template by lysis buffer (S6). (B) 800 Relative abundance of bacterial operational taxonomic units (OUT) at genus level in AM DNA water samples (S1 to S5). OUT representing less than 5% in relative abundance 801 802 were grouped (red, <1% Rel. ab.). No reads were obtained from the blank DNA control. 803 Presence/absence of larvae and diet is indicated for the five samples.

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**FIG S4** *Ae. albopictus* larval viability depends on microbial exposure and diet concentration. Larval viability expressed as the proportion (in %) of conventional (CONV) exposed to a low dilution (10<sup>-3</sup>) of microbial inoculum (A) or altered microbiota (AM) (B) larvae that reached pupal stage upon different diet concentrations (in %). Each dot represents the mean viability of 3 wells from two independent 6-well plates (3 larvae/well).

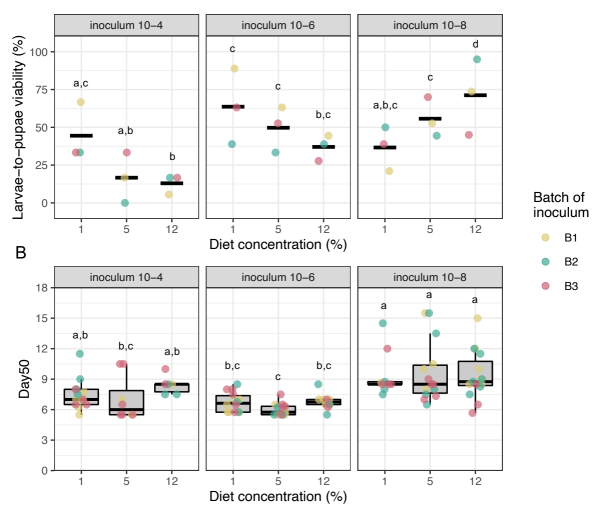
- 811 **FIG S5** Correlation patterns between relative abundance of selected OTUs and larval traits
- or experimental factors. Only the most abundant OTUs for which a significant correlation
- (Spearman, P < 0.05) with a correlation coefficient below -0.4 or above 0.4 was found are
- 814 displayed.





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