

1 **Ageing leads to nonspecific antimicrobial peptide responses in *Drosophila melanogaster***

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30 Keywords:

31 Ageing, Antimicrobial peptides, Pathogen resistance, Sexual dimorphism, Immune specificity,

32 Immune senescence

33 **ABSTRACT**

34 Evolutionary theory predicts a late-life decline in the force of natural selection, possibly
35 leading to late-life deregulations of the immune system. A potential outcome of such
36 immune-deregulation is the inability to produce specific immunity against target pathogens.
37 We tested this possibility by infecting multiple *Drosophila melanogaster* lines (with bacterial
38 pathogens) across age-groups, where either individual or different combinations of Imd- and
39 Toll-inducible antimicrobial peptides (AMPs) were deleted using CRISPR gene editing. We
40 show a high degree of non-redundancy and pathogen-specificity of AMPs in young flies: in
41 some cases, even a single AMP could confer complete resistance. In contrast, ageing led to a
42 complete loss of such specificity, warranting the action of multiple AMPs across Imd- and Toll-
43 pathways during infections. Moreover, use of diverse AMPs either had no survival benefits,
44 or even accompanied survival costs post-infection. These features were also sexually
45 dimorphic: females expressed a larger repertoire of AMPs than males, but extracted
46 equivalent survival benefits. Finally, age-specific expansion of the AMP-pool was associated
47 with downregulation of negative-regulators of the Imd-pathway and a potential damage to
48 renal function, as features of poorly-regulated immunity. Overall, we could establish ageing
49 as an important driver of nonspecific AMP responses, across sexes and bacterial infections.

50

51 INTRODUCTION

52 Ageing often leads to physiological senescence, including immune senescence, characterised
53 by exaggerated and over-reactive pro-inflammatory responses (Stout-Delgado et al., 2009;
54 Khan et al., 2017). In several insects (e.g. fruit flies and flour beetles), older individuals show
55 increased expression of antimicrobial peptides (AMPs) (Zerofsky et al., 2005); higher
56 haemolymph antibacterial activity or phenoloxidase response (PO) after infection, without
57 any significant survival benefits (Khan et al., 2016). Instead, increased immunity often induces
58 lethal immunopathological damage in older individuals, increasing their mortality rate (Khan
59 et al., 2017; Badinloo et al., 2018). Similar effects are also reported in vertebrates, where an
60 increase in chronic inflammatory state with age leads to maladaptive impacts of the innate
61 immune system (Shaw et al., 2013). For example, older mice die faster owing to an elevated
62 level of interleukin-17 and neutrophil activation, causing hepatocyte necrosis (Stout-Delgado
63 et al., 2009). By and large, older individuals are thus more likely to experience the detrimental
64 effects of overactive immunity in both invertebrate and vertebrate species.

65 Age-specific hyper-activation of immunity is consistent with the evolutionary theory of ageing
66 which predicts a progressive decline in the force of natural selection with age (Williams, 1957;
67 Hamilton, 1966)— natural selection that optimizes organismal physiology for development
68 and reproduction early in life, can become too weak to effectively regulate the late-life
69 performance in older individuals (Maklakov and Chapman, 2019). For example, poor
70 regulatory mechanisms in several evolutionarily conserved signalling pathways such as
71 insulin/ insulin-like growth factor signalling can result in suboptimal levels of gene expression
72 in late life, with myriad negative health effects (Kenyon, 2010; Flatt and Partridge, 2018;
73 Carlsson et al., 2021). Such changes in conserved signalling pathways might also interfere with

74 the optimal induction and regulation of costly immune pathways in aged individuals. Several
75 experiments on age-specific changes in the expression of negative regulators of immunity
76 support this hypothesis (Neves and Sousa-Victor, 2020): e.g., reduced expression of anti-
77 inflammatory cytokine interleukin-10 not only causes over-activation of cytotoxic
78 inflammatory pathways in older mice, but also promotes their muscular, cardiovascular and
79 metabolic dysfunction (Mohanty et al., 2015). In older mice and humans, a rapid age-specific
80 decline of another immunomodulatory molecule, MANF, increases the levels of pro-
81 inflammatory cytokines and activated macrophages (Mohanty et al., 2015; Neves et al., 2016).
82 These changes in immunity and fitness effects are thus an outcome of age-related
83 malfunctioning of regulatory units of immune pathways.

84 A further potential manifestation of such a deregulated ageing immune system is the
85 progressive loss of specificity to pathogens. Younger individuals can optimise their immune
86 responses by acting selectively on pathogens with a limited set of immune effectors (Moret,
87 2003, Hanson et al. 2019). In contrast, older individuals, owing to their poorly regulated
88 immunity, might show nonspecific activation of higher number of immune effectors against
89 an equivalent dose of antigenic exposure. An extended immune repertoire can also
90 collectively increase the cytotoxicity of immune responses, elevating the risk of morbidity and
91 mortality with ageing (Khan et al., 2017; Badinloo et al., 2018). Indeed, prior experiments with
92 older mice showed that pathways leading to increased production of antigen non-specific
93 antibodies can enhance the risk of autoimmune responses with no improvement in pathogen
94 clearance ability or survival (Bruce et al., 2009). However, experiments measuring the
95 functional expansion of the available immune repertoire with ageing and their role in overall
96 infection outcome is currently missing.

97 In the present work, we tested the impact of ageing on specific interactions between immune
98 effectors and bacterial infections, using multiple *D. melanogaster* lines where different
99 combinations of AMPs from the Imd and Toll pathways were knocked out by CRISPR/Cas9
100 gene editing (Hanson et al., 2019). We targeted AMPs as they have been recently shown to
101 possess a high degree of non-redundancy, non-interchangeability and specificity against a
102 range of pathogens in young flies (Hanson et al., 2019). Only a small subset of the total AMP
103 repertoire provides the most effective protection against specific pathogens so that in some
104 cases, even a single AMP is sufficient to control the growth of specific pathogens: e.g., Imd-
105 pathway responsive AMP *Diptericins* (or *Drosocin*) against *Providencia rettgeri* (or
106 *Enterobacter cloacae*) infection. Such specificity of AMP responses might also indicate
107 potentially higher adaptive values associated with using fewer immune effectors in young
108 individuals, thereby, avoiding the net fitness costs of general immune activation (Moret,
109 2003). Indeed, earlier experiments suggest that toxic levels of AMP expression, due to
110 suppression of negative immune regulators of Imd-pathway (or increased transcriptional
111 activation of its positive regulators) in young flies, can lead to reduced lifespan or extensive
112 neurodegeneration causing faster ageing (Kounatidis et al., 2017). We speculate that the
113 general loss of regulation in an ageing immune system might also accompany loss of such
114 controlled specific AMP actions, deploying more AMPs to counter equivalent infection levels,
115 but without any added survival benefits.

116 In addition, the age-specific role of AMPs can be sex-specific with a strong sex-by-age
117 interactions (Belmonte et al., 2020). For instance, overexpression of relish or Toll-responsive
118 defensin can reduce male lifespan more than that of females in *Drosophila* (Badinloo et al.,
119 2018). Also, previous studies with flies infected with *P. rettgeri* indicated *Drosophila* males

120 had higher *Diptericin* expression (Duneau et al., 2017). A relatively higher expression of
121 *Diptericin* transcripts in males is perhaps needed to support its exclusive role against *P.*
122 *rettgeri* infection, whereas low expression in females opens up possibilities where *Diptericin*
123 is either dispensable or requires compensatory actions of other AMPs. However, there are no
124 direct experiments to test these possibilities of sex-specific expansion of AMP use.

125 **MATERIALS AND METHODS**

126 I. Fly strains and maintenance

127 To test the role of ageing on AMP-driven specific immunity, we used multiple *Drosophila*
128 *melanogaster* lines where different combinations of multiple and individual AMPs were
129 knocked out mostly using the CRISPR/Cas9 gene editing or homologous recombination
130 (details described in Hanson et al., 2019; also see Fig. S1). We used null mutants for 10 of the
131 14 known *Drosophila* AMPs that are expressed upon systemic infection. These include
132 mutations from six single gene families including *Defensin* (*Def*^{SK3}), *Attacin C* (*AttC*^{Mi}), *Attacin*
133 *D* (*AttD*^{SK3}), *Drosocin* (*Dro*^{SK4}) *Metchnikowin* (*Mtk*^{R1}) and *Drosomycin* (*Drs*^{R1}) loci and two small
134 deletion removing *Diptericins* *DptA* and *DptB* (*Dpt*^{SK1}), or the gene cluster containing *Drosocin*
135 and *Attacins* *AttA* and *AttB* (*Dro-AttAB*^{SK2}). The iso-w¹¹¹⁸ (DrosDel isogenic) wild-type was used
136 as the genetic background for mutant isogenization (see Ferreira et al., 2014; Hanson et al.,
137 2019). We also used 'ΔAMPs' flies where independent mutations were recombined into a
138 background lacking 10 inducible AMPs. However, we note that the impact of ΔAMPs could be
139 due to AMPs having specific effects or combinatorial action of multiple co-expressed AMPs.
140 To tease apart these effects, we also included various combined mutants where different
141 groups of AMPs were deleted based on the pathways that they are controlled by: (1) Group
142 B - flies lacking AMPs such as *Drosocin*, *Diptericins* and *Attacins* (*AttC*^{Mi}; *AttD*^{SK1}; *Dro*^{SK4}; *Dro-*
143 *AttAB*^{SK2}) (exclusively regulated by Imd-pathway) (2) Group C - flies lacking the two Toll-
144 regulated antifungal peptide genes *Metchnikowin* and *Drosomycin* (*Mtk*^{R1}; *Drs*^{R1}) (mostly
145 regulated by Toll-pathway). We also referred to flies with single mutations lacking *Defensin*
146 (*Def*^{SK3}) (co-regulated by Imd- and Toll-pathway) as group A. Finally, we also included fly line

147 where group-A, B and C mutants were combined to generate flies lacking AMPs either from
148 groups A and B (AB), or A and C (AC), or B and C (BC).

149 We maintained all fly stocks and experimental individuals on a standard cornmeal diet also
150 known as Lewis medium (Siva-Jothy et al., 2018) at a constant temperature of 25°C on a 12 :
151 12 hour light: dark cycle at 60% humidity. To generate the experimental flies, we reared flies
152 at a larval density of ~70 eggs/ 6ml food. We collected adult males and females as virgins and
153 held at a density of 25 flies/sex/food vial for the experiment described below. Female iso-
154 w^{118} flies undergo reproductive senescence within 25 days post-eclosion (Reproductive
155 output measured for 18-hours; Mean \pm SE: 3-day-old= 6.75 ± 0.77 vs 24-day-old= 3.17 ± 0.53 ,
156 $P<0.001$). Hence, in our experiments, we used 3 and 25-day-old individuals (post-eclosion) as
157 'young' and 'old' adults, respectively. We transferred the adults to fresh food vials every 3
158 days, during the entire experimental window. By screening the single mutants, along with
159 combined genotypes, we were able to compare the changes in specific immunity as a function
160 of possible interactions between AMPs of different groups' vs function of individual AMPs
161 with ageing.

162 II. Infection protocol and the assay for post-infection survival

163 For all the infection, we either used Gram-negative bacteria *Providencia rettgeri* or
164 *Pseudomonas entomophila*. Both are natural pathogens of *Drosophila* that activate the IMD
165 pathway (Myllymäki et al., 2014) and could impose significant mortality (Galac and Lazzaro,
166 2011; Dieppois et al., 2015). To quantify post-infection survivorship, we infected flies (septic
167 injury method) in the thorax region with a 0.1 mm minutien pin (Fine Science Tools) dipped
168 into a bacterial suspension made from 5 mL overnight culture (optical density of 0.95,
169 measured at 600 nm) of either *Providencia rettgeri* or *Pseudomonas entomophila* adjusted to

170 OD of 0.1 and 0.05 respectively (See SI methods for details). In total, we infected 160-280
171 flies/sex/infection treatment/bacterial pathogen/age-group/fly genotypes and then held
172 them in food vials in a group 20 individuals (For each treatment, sex, age-group, pathogen
173 type, we thus had 8-14 replicate food vials). We carried out sham infection with a pin dipped
174 in sterile phosphate buffer solution (1X PBS).

175 We then recorded their survival every 4-hours (± 2) for 5 days. Due to logistical challenges of
176 handling a large number of flies, we infected each sex and age-groups with *P. rettgeri* (or *P.*
177 *entomophila*) separately in multiple batches, where they were handled as — (i) Groups AB,
178 BC, AC; (ii) Group-A, B & C; (iii) Imd-responsive and (iv) Toll-responsive single mutants for *P.*
179 *rettgeri*; or (i) Groups AB, BC, AC, A, B & C; (ii) Imd-responsive and (iii) Toll-responsive single
180 mutants for *P. entomophila*. Every time, we also assayed iso-*w¹¹¹⁸* flies as a control to facilitate
181 a meaningful comparison across different batches. Therefore, although sexes and age-groups
182 for each mutant were not directly comparable, their relative effects with respect to control
183 iso-*w¹¹¹⁸* were estimated across sexes, age-groups and pathogen types. Note that we
184 compared each mutant separately with iso-*w¹¹¹⁸* flies, since we only wanted to capture their
185 changes in infection susceptibility relative to control flies. For each batch of flies, across
186 pathogen types, sexes and age-groups, we analysed the survival data with a mixed effects Cox
187 model, using the R package 'coxme' (Therneau, 2015). We specified the model as: survival ~
188 fly lines (individual AMP mutant lines vs iso-*w¹¹¹⁸*) + (1|food vials), with fly lines as a fixed
189 effect and replicate food vials as a random effect. Since none of the fly lines had any mortality
190 after sham-infection, we were able to quantify the susceptibility of each infected mutant lines
191 (AMP knockouts) with respect to control flies (iso-*w¹¹¹⁸* group) as the estimated hazard ratio
192 of infected AMP mutants versus control flies (hazard ratio = rate of deaths occurring in

193 infected AMP mutants /rate of deaths occurring in iso- w^{1118} group). A hazard ratio
194 significantly greater than one indicated a higher risk of mortality in the AMP mutant
195 individuals.

196 Note that the above experimental design allowed us to repeat the assay for post-infection
197 survival of young and old iso- w^{1118} flies infected with *P. rettgeri* (or *P. entomophila*) in 4 (or 3)
198 independently replicated experiments. We thus estimated the effects of ageing on their post-
199 infection survival, using a mixed effects Cox model specified as: survival ~ age + (1|food vials),
200 with age as a fixed effect, and food vials as random effects.

201 III. Assay for bacterial clearance

202 Mortality of control flies (iso- w^{1118}) injected with the experimental infection dose began
203 around 24-hours and 20-hours after infection with *P. rettgeri* and *P. entomophila* respectively
204 (Fig. S2A). We therefore used these time-points to estimate the bacterial load across the age-
205 groups as a measure of the pathogen clearance ability across AMPs (see SI methods for
206 detailed protocol). We homogenized flies in a group of 6 in the sterile PBS (n= 8-15 replicate
207 groups/sex/treatment/age-group/fly lines), followed by plating them on Luria agar. Due to
208 logistical challenges with large number of experimental flies, we handled each sex, age-group
209 and pathogen type separately and in multiple batches as described above.

210 Also, similar to post-infection survival data, we were only interested in comparing the changes
211 in bacterial load for each mutant line relative to control iso- w^{1118} flies across experimental
212 groups. We thus analysed the bacterial load data of each mutant genotype with iso- w^{1118} flies
213 separately across age-groups, sexes and pathogen types. Since residuals of bacterial load data
214 were non-normally distributed (confirmed using Shapiro-Wilks's test), we log-transformed

215 the data, but residuals were still non-normally distributed. Subsequently, we analysed the
216 log-transformed data, using a generalised linear model best fitted to gamma distribution, with
217 fly lines (i.e., control iso- w^{118} line vs individual AMP knockout line) as a fixed effect.

218 IV. Assay for the Malpighian tubule activity, as a proxy for immunopathological
219 damage

220 Malpighian tubules (MTs), the fluid-transporting excretory epithelium in all insects, are prone
221 to increased immunopathology following an immune activation due to their position in the
222 body and the fact that they cannot be protected with an impermeable membrane due to their
223 functional requirement (Dow et al., 1994; Khan et al., 2017). Previous experiments have
224 shown that risk of such immunopathological damage can increase further with ageing in
225 mealworm beetle *Tenebrio molitor* (Khan et al., 2017). It is possible that nonspecific AMP
226 responses with ageing in *Drosophila* was also associated with increased immunopathological
227 damage to MTs. We thus estimated the fluid transporting capacity of functional MTs
228 dissected from experimental females at 4-hours after immune challenge with 0.1 OD *P.*
229 *rettgeri* (n=12-20 females/ infection treatment/age-group), using a modified 'oil drop'
230 technique as outlined in previous studies (Dow et al. 1994; Li et al., 2020) (also see SI
231 methods).

232 This method provides a functional estimate of their physiological capacity by assaying the
233 ability to transport saline across the active cell wall into the tubule lumen. The volume of the
234 secreted saline droplet is negatively correlated with the level of immunopathological damage
235 to MTs. Since we collected the flies across the age-groups on different days, we analysed the

236 MT activity data as a function of infection status for each age-group separately, using a
237 generalized linear mixed model best fitted to a quasibinomial distribution.

238 V. Gene expression assay

239 Finally, we note that transcription of negative regulators of Imd-pathway such as *pirk* and
240 *caudal* are important to ensure an appropriate level of immune response following infection
241 with gram-negative bacterial pathogens, thereby avoiding the immunopathological effects
242 (Lee and Ferrandon, 2011; Kleino and Silverman, 2014). While *pirk* interferes with the
243 interaction of *PGRP-LC* and -*LE* with the molecule Imd to limit the activation of the Imd
244 pathway, *caudal* downregulates the expression of AMPs (Lee and Ferrandon, 2011). To
245 examine whether non-specific expansion of AMP repertoire was associated with the lower
246 expression of these negative regulators, we estimated their relative expression level in both
247 young and old iso-*w*¹¹¹⁸ individuals infected with *P. rettgeri* at 24 hours post-infection, by
248 using qPCR (as outlined in Prakash et al., 2021) (n= Total 15-21 flies in a group of 3
249 homogenized in Trizol reagent/ Infection treatment/ age-group and sex-combination).

250 In addition, we also estimated the expression of the Imd-pathway NF- κ B transcription factor
251 *Relish* and peptidoglycan recognition protein - *PGRP-LC*, both act as positive regulators of Imd-
252 pathway (Lemaitre and Hoffmann, 2007; Myllymäki et al., 2014) and hence, can serve as a
253 proxy for overactivated Imd pathway and higher AMP expression in older flies (Badinloo et
254 al., 2018) (also see SI methods). We analysed the gene expression data using ANOVA (see SI
255 methods section-iv for details).

256 **RESULTS**

257 We began our observation with high mortality and increased bacterial load in AMP-deficient
258 flies (Δ AMPs) infected with *P. rettgeri*, regardless of their sex and age (Fig. 1A, 1B, 1E, 1F;
259 Table S2, S3), suggesting that AMPs are critically important to prevent pathogen growth,
260 thereby increasing the post-infection survival costs (Fig. 1A, 1B, 1E, 1F; Fig. S3, S4; Table S2,
261 S3). Older iso-*w¹¹¹⁸* control females infected with *P. rettgeri* also showed higher mortality (Fig.
262 S2A, S2B; Table S4A) and increased bacterial load (Fig. S2C, Table S4B) than their younger
263 counterparts, suggesting negative effects of ageing on fitness and pathogen clearance ability.
264 By contrast, both young and old males had similar post-infection mortality rate (Fig. S2A, S2B;
265 Table S4A) with comparable bacterial load (Fig. S2C; Table S4B), indicating that ageing did not
266 impact male's ability to survive post-infection and clear pathogens, at least at the infection
267 dose used in our experiments. Nevertheless, in the subsequent assays, these results from
268 male flies infected with *P. rettgeri* enabled us to directly compare the relative effects of
269 deleting different Imd- vs Toll-responsive AMPs across age-groups (against a common
270 baseline).

271 I. **Ageing leads to an expansion of the required AMP repertoire against *P. rettgeri***
272 **infection**

273 To gain a broad understanding of how AMP specificity changes with age, we first tested
274 mutants lacking different groups of AMPs either from Imd- (e.g., group B) or Toll-pathways
275 (e.g., group C) (pathway-specific), or combined mutants lacking pathway-specific mutants in
276 different combinations (e.g., group AB, BC or AC) (See Fig. S1 for description of mutants). As
277 reported in a previous study by Hanson and co-workers (Hanson et al., 2019), young males
278 lacking group-AB and -BC AMPs were highly susceptible to *P. rettgeri* infection (Fig. 1A; Table

279 S2), and this was generally associated with 10-100-fold increased bacterial loads in these
280 mutants relative to the iso-*w¹¹¹⁸* control (Fig. 1B; Table S3). Subsequent assays with pathway-
281 specific (i.e., Imd- or Toll-pathway) AMP combinations (group A, B or C) confirmed that such
282 effects were primarily driven by Imd-regulated group-B AMPs that were shared between both
283 AB and BC combinations (Fig. 1C; Fig. S3; Table S2), and equally driven by increased bacterial
284 load (Fig. 1D; Table S3). We found a comparable pattern in young females as well, except that
285 flies lacking group BC combinations of AMPs were not negatively affected by infection (Fig.
286 1E, 1F, 1G, 1H; Fig. S4; Table S2, S3).

287 In contrast to young flies, most of the pathway-specific or combined mutants became highly
288 susceptible to *P. rettgeri* infection with age, except females of group-A mutants flies lacking
289 *Def*. This would suggest a possible sexually dimorphic effect of *Defensin* in *P. rettgeri* infection,
290 which appear to be important for males, but not females (Fig. 1C, 1G; Fig. S3, S4; Table S2).
291 Regardless of this slight variation across sexes, our results clearly demonstrated that only
292 having functional Imd-regulated group-B AMPs was not sufficient to protect older flies against
293 *P. rettgeri* infection. Also, these results indicated that a single AMP *Dpt*-driven protection
294 against *P. rettgeri* infection, as suggested by Hanson et al. (2019), may not be applicable to
295 older flies (also see Unckless et al., 2016). High susceptibility of older mutants lacking group
296 A or C AMPs (Fig. 1C, 1G; Table S2) and increased bacterial growth (Fig. 1D, 1H; Table S3)
297 therein clearly indicated that other AMPs responsive to Gram-positive bacteria (e.g., *Def*) or
298 fungal pathogens (e.g., *Mtk*, *Drs*) might be needed as well.

299 II. ***Dpt*-specificity against *P. rettgeri* infection is sex-specific and disappears with age**
300 Next, we decided to test the role of individual AMPs deleted in the pathway-specific or
301 compound mutants across age-groups and sexes. Interestingly, *Dpt* provided complete

302 protection against *P. rettgeri* only in young males, but not in females or older males (Fig. 2A,
303 2E; Table S5). This was verified by using fly lines where *DptA* and *DptB* are introduced on an
304 AMP-deficient background (ΔAMPs^{+Dpt}). *Dpt* reintroduction could fully restore survival as that
305 of wild-type flies only in young males, and this was associated with a decrease in CFUs
306 compared to the *Dpt* deletion mutant (Fig. 2B, Table S6). However, reintroduction of
307 functional *DptA* and *DptB* (ΔAMPs^{+Dpt}) in young or old females flies did not result in lower
308 CFUs (Fig. 2F; Table S6) and these flies remained highly susceptible to *P. rettgeri* (Fig. 2E; Table
309 S2). Young (or old) females also showed increased bacterial loads and associated higher
310 susceptibility when other Imd-regulated group-B AMPs such as *AttC* and *Dro-Att* (or *Dro* and
311 *Dro-Att* in old females) were deleted (Fig. 2E, 2F; Table S5, S6). Older females lacking *AttC*
312 showed increased infection susceptibility as well, but did not have increased bacterial load
313 (Fig. 2E, 2F; Table S5, S6). Older ΔAMPs^{+Dpt} males, on the other hand, could limit the bacterial
314 burden as low as that of the control iso-*w¹¹¹⁸* flies (Fig. 2B; Table S6), but still showed very
315 high post-infection mortality (Fig. 2A; Table S5). These results from older males thus
316 suggested that the ability to clear pathogens might not always translate into an improved
317 ability to survive after infection (Fig. 2A, 2B; Table S5, S6).

318 Why did females always require AMPs other than *Dpt* after *P. rettgeri* infection? Although the
319 mechanisms behind sex-specific expansion of AMP repertoire are unknown, a possible
320 explanation is that females show inherently lower expression level of *Dpt* relative to males .
321 Consequently, they may require the joint expression of other AMPs to complement the lower
322 *Dpt* expression, thereby enhancing the protection against *P. rettgeri* infection. Indeed, a
323 previous study has already demonstrated lower *Dpt* expression in iso-*w¹¹¹⁸* females than
324 males after *P. rettgeri* infection (see Duneau et al., 2017), although the causal link between

325 reduced *Dpt* expression and proportional increase in the compensatory action of other AMPs
326 is not yet experimentally validated.

327 Also, both males and females showed further extension to a Toll-responsive AMP repertoire
328 with ageing. In addition to the role of *Def* (included in group-A) as described above in older
329 males (but not in older females; compare Fig. 1D, 1H; Table S3), older flies of both sexes also
330 showed increased microbe loads and increased mortality when Toll-regulated AMPs from
331 such as *Drs* and *Mtk* were deleted (Fig. 2D, 2H; Table S6), raising a possibility of crosstalk
332 between Toll and Imd immune-signalling pathways (Duneau et al., 2017; Nishide et al., 2019).
333 Taken together, these results describe ageing as a major driver behind the loss of specificity
334 of AMP responses.

335 Additionally, we also note that a few other mutations such as deletion of *Dro* and *Dro-Att*,
336 which otherwise had no effects on the survival of *P. rettgeri*-infected young males, caused
337 significant increase in the bacterial load (Fig. 2A, 2B; Table S5, S6). Together, these results not
338 only underscored the multifaceted role of AMPs, but also provided functional resolution at
339 the level of single AMPs such as *Dpt* which in addition to playing the canonical role in resisting
340 the infection, also aided in withstanding the effects of increased pathogen growth, caused by
341 the dysfunction of other AMPs (Fig. 2A, 2B; Table S5, S6).

342 **III. Expansion of the required AMP repertoire does not improve, and even reduces,
343 survival in both older males and females infected with *P. entomophila***

344 To test if age-related loss of AMP specificity was specific to *P. rettgeri*, or also occurred with
345 other infections, we investigated the AMP repertoire in young and old flies infected with the
346 Gram-negative bacteria *P. entomophila*. Similarly, older flies required a larger repertoire of

347 AMPs (Fig. 3A, 3C; Table S7) and yet, died faster than young flies (old vs young: 4-fold vs 2-
348 fold; Fig. 3A, 3C). In contrast to younger flies, where only group-B, -AB and -BC mutants were
349 susceptible to *P. entomophila* infection, all the other pathway-specific or combined mutants
350 of older males and females were also highly sensitive to infection (Fig. 3A, 3C; Table S7).
351 However, further experiments with single AMP mutants revealed that the antibacterial
352 protection in both young males and females was still limited only to the exclusively Imd-
353 regulated group-B AMPs, where several of them individually caused significant increase in
354 microbe loads and reduction in post-infection survival (Fig. 4A, 4B, 4E, 4F; Table S9, S10). In
355 contrast, older flies also needed additional action of Toll-regulated AMP *Drs* (Fig. 4C, 4G; Table
356 S9), though it is striking that that increased mortality was not associated with increased
357 microbe loads relative to iso-*w¹¹¹⁸* in this case (Fig. 4D, 4H; Table S10). Overall, this is
358 comparable to *P. rettgeri* infection where potential crosstalk between Toll & Imd immune-
359 signalling pathways has already been implicated with ageing (Fig. 2; Table S5, S6). Also, the
360 broad similarity between age-specific expansion and cross-reactivity of AMP repertoire
361 against two different pathogens indicated the possibility where non-specificity can indeed be
362 a generalised feature of an ageing immunity. Moreover, the increased mortality in older flies
363 infected with *P. entomophila*, despite involving a higher number of AMPs, was perhaps an
364 indication of their exacerbated cytotoxic effects with age (Badinloo et al., 2018).

365 **IV. Ageing-induced expansion of the required AMP repertoire was associated with**
366 **downregulation of negative immune regulators and a trend of reduced renal**
367 **purging post-infection**

368 The expansion of the AMP repertoire in older flies could reflect a compensatory action to
369 balance the lower per capita efficiency of their individual AMPs. This would enable flies to

370 maintain an equivalent post-infection survival as that of younger flies against similar infection
371 dose (e.g., old vs young males infected with *P. rettgeri*; Fig. 1A; Table S2). However, any
372 benefits of recruiting multiple AMPs, may have been outweighed by the costs of expressing
373 them (suggested in Badinloo et al., 2018) as higher immune activity, in general, accelerates
374 the ageing process by imposing immunopathological damage to vital organs such as
375 Malpighian tubules (MTs) (Khan et al., 2017). We expected that expansion of the AMP
376 repertoire might have similar consequences in our experimental older flies as well. This is
377 closely reflected by our results where *P. rettgeri* infection significantly reduced renal function
378 in older females, measured as MT secretion (Fig. 5A; Table S11). Since functional MTs are
379 needed to purge excessive ROS produced during immune responses as a physiological
380 adaptation to prevent tissue damage in *Drosophila* (Li et al., 2020), reduced MT activity might
381 not only exacerbates the effects of pathogenic infection, but can also causes late-life costs
382 (Khan et al., 2017).

383 Finally, we also found ageing-associated downregulation of the major negative regulators of
384 Imd-signalling such as *Caudal* & *Pirk* in older flies (Fig. 5B; Table S12), which has been
385 previously linked to the production of toxic levels of AMP production, causing reduced
386 lifespan, locomotor defects and extensive neurodegeneration (Kounatidis et al., 2017;
387 Prakash et al., 2021). Based on these results, we speculate that the observed expansion of
388 AMP repertoire with age is therefore most likely to represent suboptimal body condition,
389 characterized by poorly regulated immune system and increased physiological costs.

390 **Discussion**

391 Recent studies performed functional validation of *Drosophila* AMPs, revealing remarkable
392 specificity and non-redundant interactions with subsets of pathogens that they target
393 (Hanson et al., 2019). In the present work, we analysed these specific AMPs responses
394 primarily as a function of ageing that alters the regulation and relative investment in immune
395 responses (Khan et al., 2016, 2017). We used two bacterial entomopathogen *P. rettgeri* and
396 *P. entomophila* to induce various level of AMP responses inside a fly host, ranging from a
397 single AMP to pathway-specific expression [e.g., Imd vs Toll; (Hanson et al., 2019)]. Further,
398 although sex profoundly impacts the relative use of AMPs (Duneau et al., 2017), previous
399 studies addressing AMP specificity have almost entirely focussed on males (Unckless et al.,
400 2016; Hanson et al., 2019). We thus also included both males and females in our experiments
401 to test the sex-specific effect of ageing on AMP functions. In fact, we showed that the
402 efficiency of these AMP responses is strictly age-driven with high degree of sexual
403 dimorphism. For example, the classic *Dpt*-driven protection against *P. rettgeri*, as shown by
404 previous studies (Hanson et al., 2019), is only limited to young males, whereas females also
405 needed other Imd-regulated AMPs. Although the reason is unclear, we speculate that
406 multiple AMPs were needed possibly to compensate the inherently lower expression level of
407 *Dpt* transcript in females than males (Duneau et al., 2017; also shown by Prakash, 2022).
408 However, regardless of sex and pathogen, ageing led to a more drastic expansion of AMP
409 repertoire— instead of deploying only canonical expression of Imd-responsive AMPs to
410 counter Gram-negative bacterial infections, older males and females also used AMPs from
411 Toll pathways.

412 Surprisingly, despite using more diverse AMPs, late-life expansion either did not confer any
413 survival benefits (during *P. rettgeri* infection in older males) or was associated with survival
414 costs (after *P. entomophila* infection). We thus speculate that the nonspecific use of AMPs
415 with ageing was unnecessary, perhaps indicating an immune system failing to control over-
416 reactive immune responses with potentially immunopathological effects (Stout-Delgado et
417 al., 2009; Goldstein, 2010; Khan et al., 2017; Badinloo et al., 2018). This notion was further
418 supported by reduced expression levels of negative regulators of immune responses such as
419 *caudal* and *pirk* in older flies, which have been previously implicated in over-activating Imd-
420 signalling and AMP expression. In addition, reduced renal function or Malpighian tubule
421 activity in infected older flies suggested that expanded AMP repertoire might not be able to
422 prevent the plausible physiological costs of bacterial infection. Although not verified
423 experimentally, we suspect a causal role of overactivated AMPs here. This is because (a)
424 overactive and simultaneously expressed multiple AMPs can impose cytotoxic effects
425 (Badinloo et al., 2018), and (b) reduced Malpighian tubule activity is already a known
426 manifestation of immunopathological costs caused by overactive insect immune components
427 (Sadd and Siva-Jothy, 2006; Khan et al., 2017), reducing fitness by accumulating toxic
428 metabolites (Li et al., 2020).

429 We also note an alternative possibility where age-specific increase in AMP expression could
430 have been beneficial. For example, since ageing can lead to accumulation of diverse microbes
431 in the body cavity (Ren et al., 2007; Arias-Rojas and Iatsenko, 2022), this might warrant the
432 overexpression of multiple AMPs to tackle the antigenic diversity of many microbial species
433 to maintain the health (Ren et al., 2007; Badinloo et al., 2018). Indeed, previous experiments
434 have found that highly expressed Imd-responsive AMPs such as *CecA1* and *Dro* were needed

435 to maintain health while extending the lifespan in *Drosophila* (Loch et al., 2017). However,
436 benefits of non-specific, highly expressed immune responses may still not be able to outweigh
437 the net costs of overreactive immune responses. In fact, detrimental effects of overreactive
438 immunity with ageing has been supported by recent analyses linking weaker strength of
439 purifying selection in older individuals and high frequency of non-synonymous and disease-
440 causing mutations (Cheng and Kirkpatrick, 2021). This in turn can lead to poorly-regulated
441 gene expression network in older animals with increased cancer risk in a range of species,
442 including humans. Taken together, non-specific AMP responses with ageing is thus a more
443 likely feature of a deregulated immune system of older individuals (Kounatidis et al., 2017).

444 Finally, the use of diverse array of AMP deletion mutants allowed us to capture enormous
445 functional diversity of AMPs, revealing dynamic age- and sex-specific changes in their
446 pathogen clearance ability. Older individuals showed increased divergence between
447 individual AMPs vs their combined action (e.g., *Dpt* vs group-B mutants in older females),
448 possibly indicating greater complexity associated with higher number of AMPs in use vs their
449 various interactions. Although we did not find much evidence of synergism or additive effects
450 between individual AMPs (but see the older males infected with *P. entomophila*),
451 indispensability of each AMPs to maintain the fitness post-infection in older flies suggested
452 the mutually non-exclusive and intertwined nature of their activity with ageing. We hope that
453 these results will motivate future studies to investigate the deeper mechanistic details of
454 nonspecific AMP function with ageing. Also, with growing importance of AMPs in developing
455 novel antibiotics and autoimmune disease research, identifying age or sex as major sources
456 of variability in AMP functions and fitness impacts might have significant importance for
457 therapeutic and gerontological research.

458 **Acknowledgements**

459 We thank Basabi Bagchi, Joy Bose, and Srijan Seal for feedback on the manuscript. We are
460 grateful to Bruno Lemaitre and Mark A. Hanson for generously providing us the fly lines. We
461 thank Srijan Seal, Katy M. Monteith, Raghav Pavan Thunga and Devshuvam Banerji for
462 laboratory assistance.

463 **Author contribution**

464 IK conceived the experiments; IK, AP, BS designed the experiments; AP, BS, and SS performed
465 the experiments; AP, BS and IK analysed the data; IK and PV acquired the funding and
466 provided resources and consumables. IK and AP drafted the manuscript with additional input
467 and comments from BS, SS and PV. All authors agreed on the final version of the manuscript.

468 **Funding**

469 We thank the grant supplement from SERB-DST India (No. ECR/2017/003370) to I. Khan and,
470 Ashoka University for supporting this research. The research was also funded by a Society in
471 Science Branco Weiss fellowship and a Chancellor's Fellowship (University of Edinburgh) both
472 awarded to P. Vale. A. Prakash was funded by a Darwin Trust PhD Studentship (U. Edinburgh).

473

474 **Competing interest**

475 None

476 **References**

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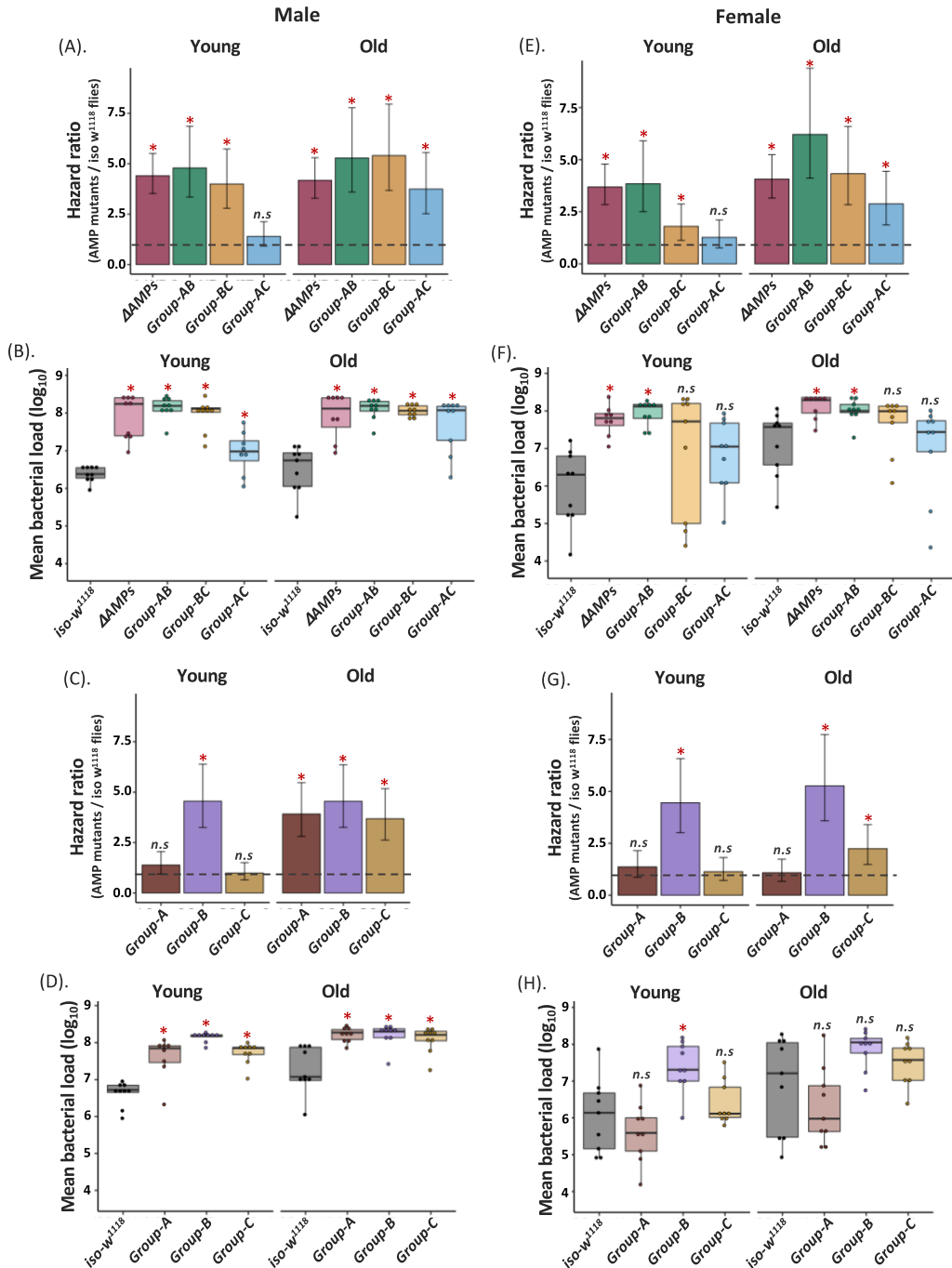
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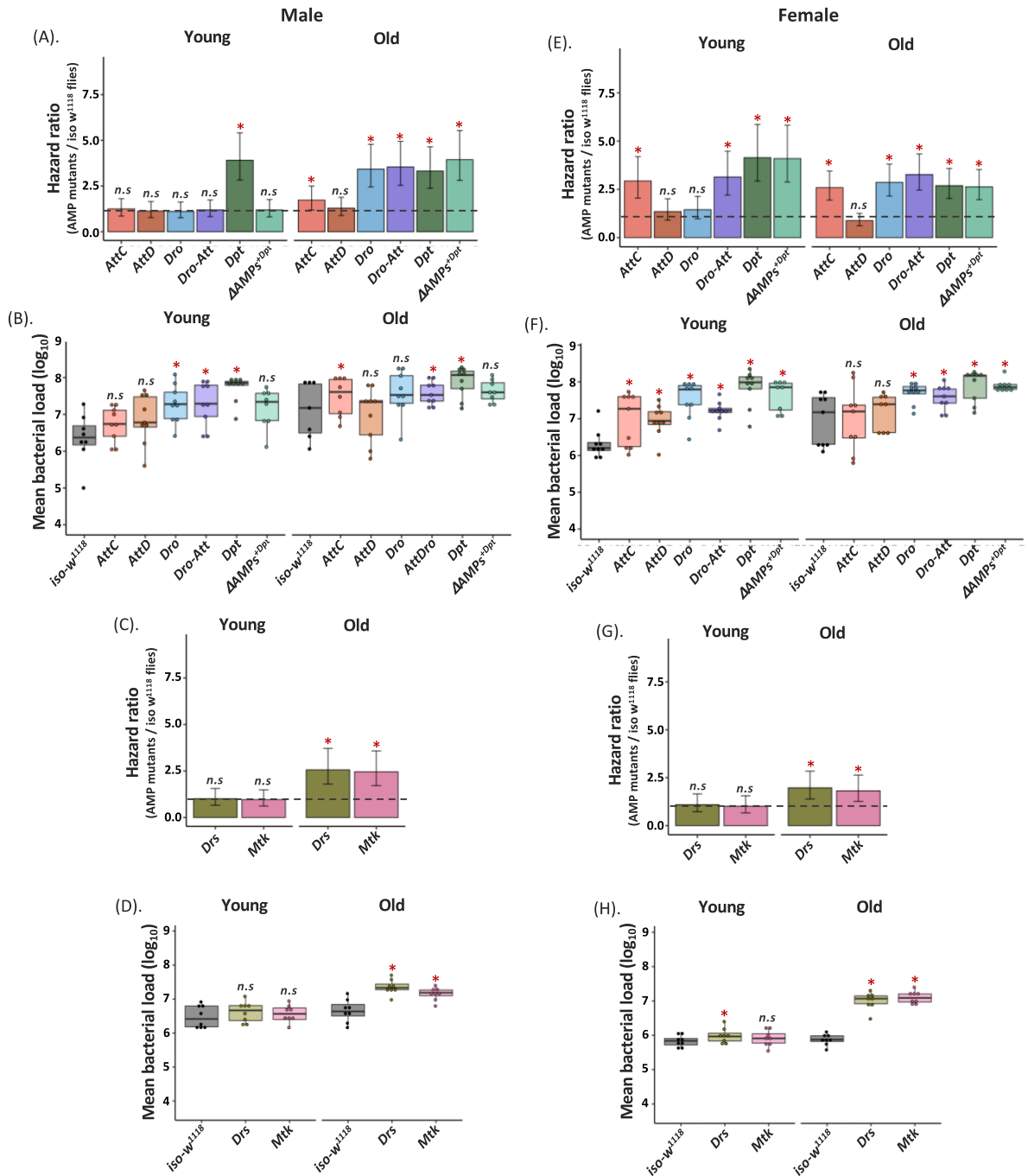
597 **Figures**

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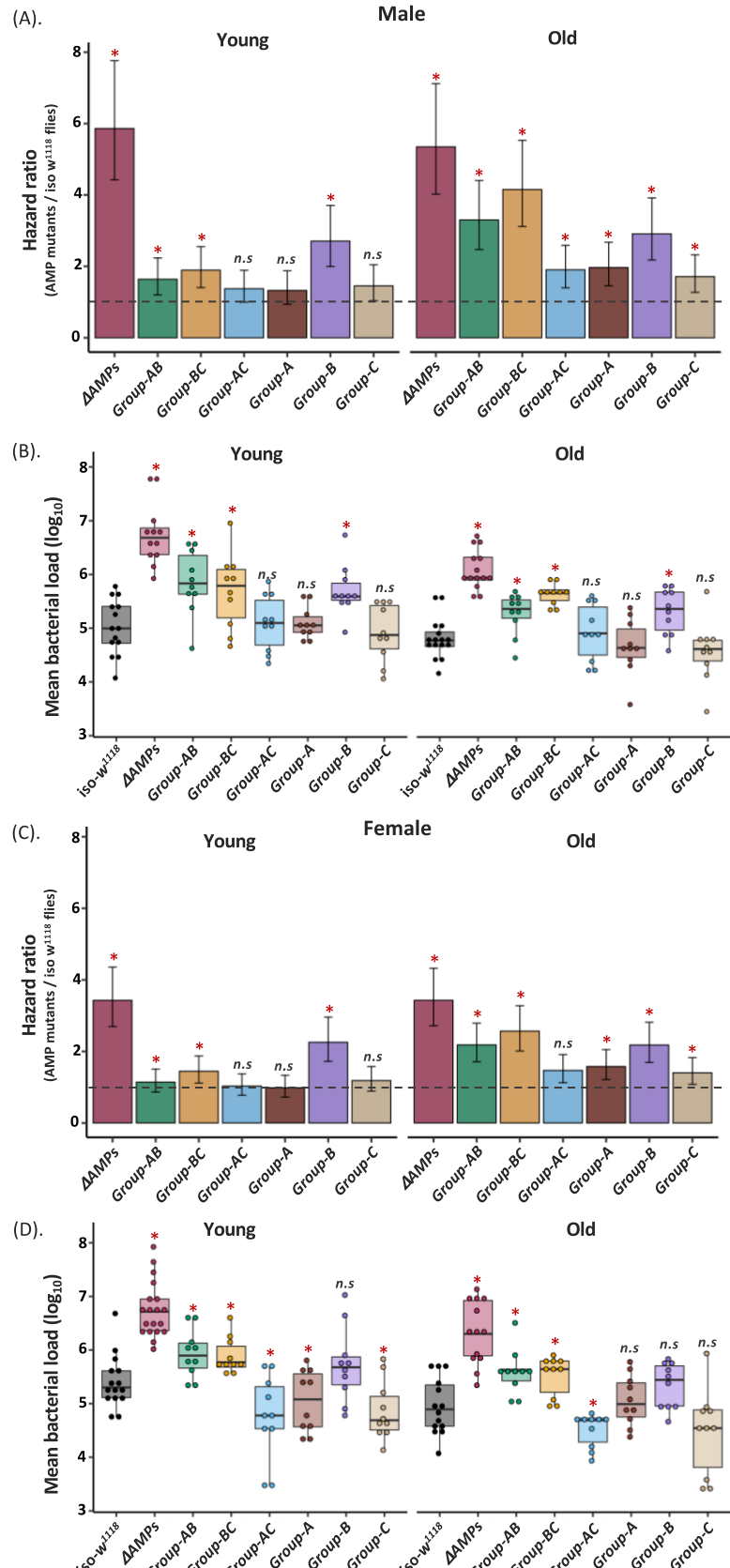
599 **Figure 1. Infection with *Providencia rettgeri* in multiple AMP-knockouts.** The estimated
600 hazard ratios calculated from survival curves (160-180 flies/sex/infection treatment/ age-
601 group/fly line; see Fig. S3, S4) and bacterial load (n= 8-9 replicate groups/sex/treatment/age-
602 group/fly line) measured at 24-hours after *P. rettgeri* infection across sexes and age-groups.
603 Hazard ratios for double combination of AMP-knockouts (i.e., group-AB, BC, & AC; see Fig. S1
604 for details about the fly lines) in males **(A)** and females **(E)**. Bacterial loads for double
605 combination of AMP-knockouts in males **(B)** and females **(F)**. Hazard ratios for single
606 combination of AMP Knockouts (e.g., group- A, B & C) in males **(C)** and females **(G)**. Bacterial
607 load for single combination of compound of AMP-knockouts in males **(D)** and females **(H)**. In
608 panels A, C, E, G, hazard ratios significantly greater than 1 (hazard ratio =1; shown as
609 horizontal dashed grey lines), indicated by asterisk (*), suggests higher infection susceptibility
610 of mutant flies than the iso-*w¹¹¹⁸* control flies. In panels B, D, F, H, each data point represents
611 the bacterial load of flies pooled in a group of 6. Mutant fly lines that had significantly
612 different bacterial load from wild-type iso-*w¹¹¹⁸* are indicated by asterisks. ns = not significant.
613 Group A- flies lacking *Defensin*; Group B - flies lacking AMPs such as *Drosocin*, *Diptericins* and
614 *Attacins*; Group C - flies lacking *Metchnikowin* and *Drosomycin*; Group-A, B and C mutants
615 were combined to generate flies lacking AMPs either from groups A and B (AB), or A and C
616 (AC), or B and C (BC).



618 **Figure. 2. Infection with *Providencia rettgeri* in individual AMP-knockouts.** The estimated
619 hazard ratios calculated from survival curves (160-180 flies/sex/infection treatment/ age-
620 group/fly line; see Fig. S3, S4) and bacterial load (n= 8-9 replicate groups/sex/treatment/age-
621 group/fly line) measured at 24 hours after *P. rettgeri* infection across sexes and age-groups.
622 Hazard ratios for Imd-responsive single AMP (e.g., *Dpt*, *AttC*, *AttD*, *Dro*; see Fig. S1 for details
623 about the fly lines) and Att-Dro knockouts in males (**A**) and females (**E**). Bacterial load of Imd-
624 responsive single AMP and Att-Dro knockouts in males (**B**) and females (**F**). Hazard ratios for
625 Toll-responsive single AMP knockouts (e.g., *Drs* & *Mtk*) in males (**C**) and females (**G**). Bacterial
626 loads of Toll-responsive single AMP knockouts in males (**D**) and females (**H**) respectively. In
627 panels A, C, E, G, hazard ratios significantly greater than 1 (hazard ratio =1; shown as
628 horizontal dashed grey lines), indicated by asterisk (*), suggests higher infection susceptibility
629 of mutant flies than the iso-*w¹¹¹⁸* control flies. In panels B, D, F, H, each data point represents
630 the bacterial load of flies pooled in a group of 6. Mutant fly lines that had significantly
631 different bacterial load from wild-type iso-*w¹¹¹⁸* are indicated by asterisks (*). ns = not
632 significant.

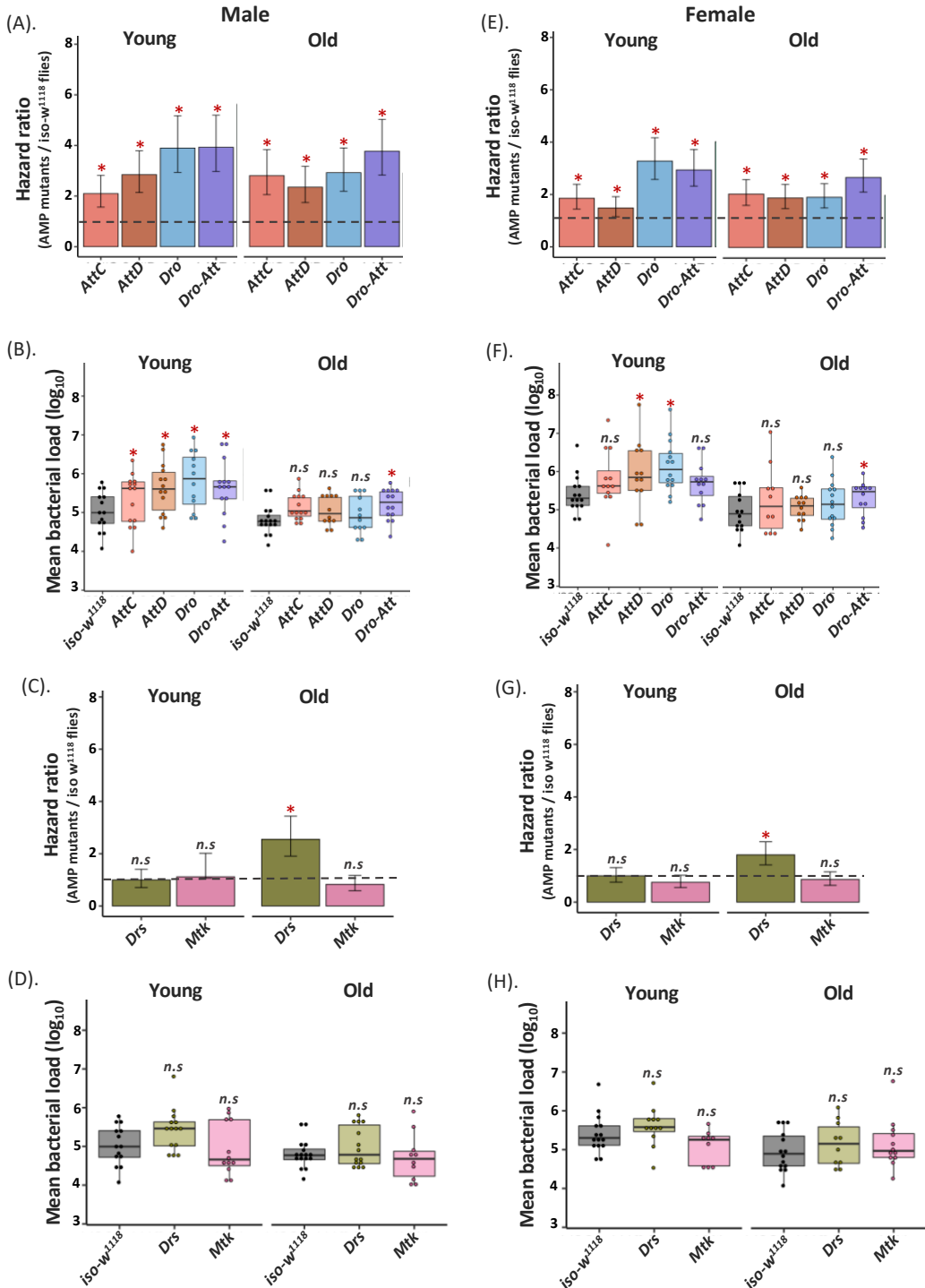


634 **Figure 3. Infection with *Pseudomonas entomophila* in multiple AMP-knockouts.** The
635 estimated hazard ratios calculated from survival curves (180-280 flies/treatment/age-
636 groups/sex/fly line; see SI Fig. S5, S6) and bacterial load (n= 9-15 replicate
637 groups/sex/treatment/age-group/fly line) measured at 20-hours after *P. entomophila*
638 infection across sexes and age-groups. Hazard ratios for double (i.e., group-AB, BC, & AC) and
639 single combination (i.e., group-A, B, C) of AMP-knockouts in males (**A**) and females (**C**).
640 Bacterial loads for double and single combination of AMP-knockouts in males (**B**) and females
641 (**D**). In panels A & C hazard ratios significantly greater than 1 (hazard ratio =1; shown as
642 horizontal dashed grey lines), indicated by asterisk (*), suggests higher infection susceptibility
643 of mutant flies than the iso-*w¹¹¹⁸* control flies. In panels B & D each data point represents the
644 bacterial load of flies pooled in a group of 6. Mutant fly lines that had significantly different
645 bacterial load from wild-type iso-*w¹¹¹⁸* are indicated by asterisks. ns = not significant. See Fig
646 1 or the main text for the description of different fly groups.

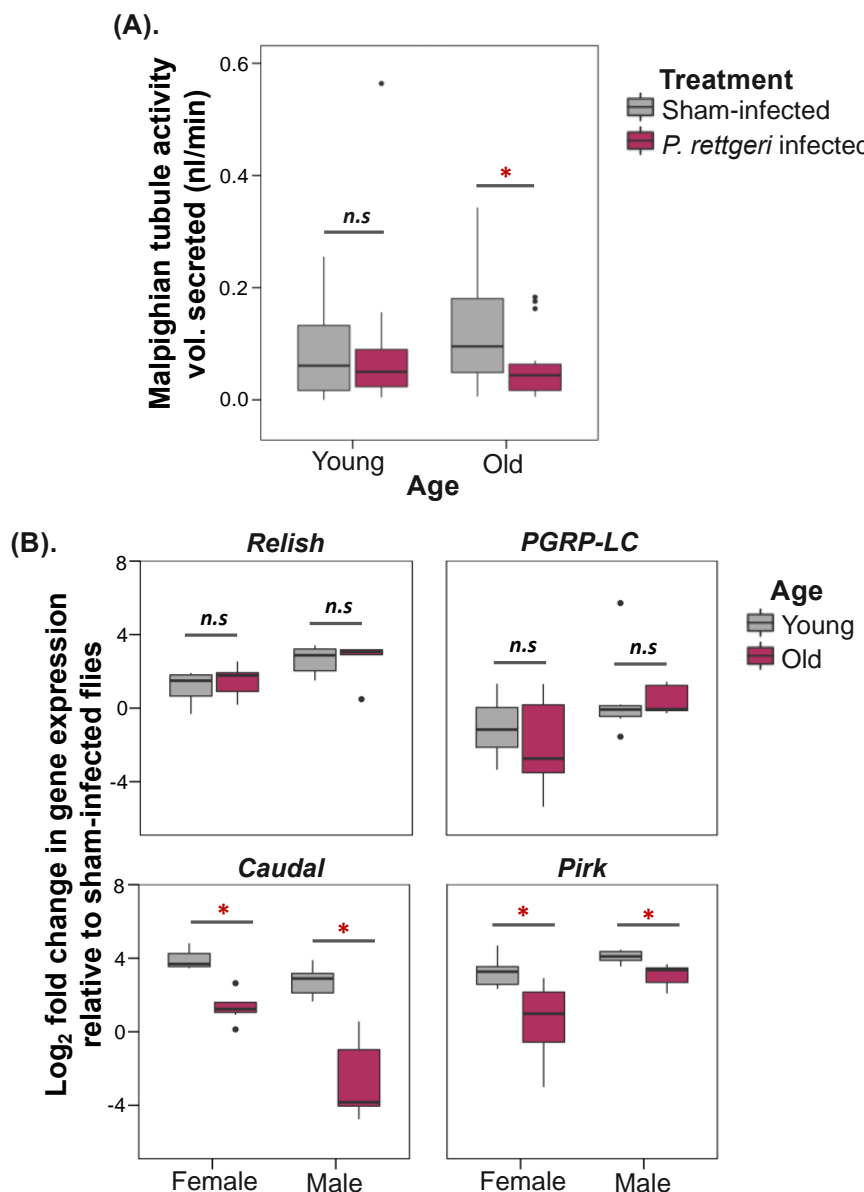


648 **Figure 4. Infection with *Pseudomonas entomophila* in individual AMP-knockouts.** The
649 estimated hazard ratios calculated from survival curves (180-280 flies/sex/infection
650 treatment/ age-group/fly line; see SI Fig. S5, S6) and bacterial load (n= 9-15 replicate
651 groups/sex/treatment/age-group/fly line) measured at 20 hours after *P. entomophila*
652 infection across sexes and age-groups. Hazard ratios for Imd-responsive single AMP (e.g.,
653 *AttC*, *AttD*, *Dro*; see Fig. S1 for details about the fly lines) and *Att-Dro* knockouts in males **(A)**
654 and females **(E)**. Bacterial load of Imd-responsive single AMP and *Att-Dro* knockouts in males
655 **(B)** and females **(F)**. Hazard ratios for Toll-responsive single AMP knockouts (e.g., *Drs* & *Mtk*)
656 in males **(C)** and females **(G)**. Bacterial loads of Toll-responsive single AMP knockouts in males
657 **(D)** and females **(H)** respectively. In panels A, C, E, G, hazard ratios significantly greater than
658 1 (hazard ratio =1; shown as horizontal dashed grey lines), indicated by asterisk (*), suggests
659 higher infection susceptibility of mutant flies than the iso-*w¹¹¹⁸* control flies. In panels B, D, F,
660 H, each data point represents the bacterial load of flies pooled in a group of 6. Mutant fly lines
661 that had significantly different bacterial load from wild-type iso-*w¹¹¹⁸* are indicated by
662 asterisks (*). ns = not significant.

663



665 **Figure 5. Ageing-associated immune dysregulation and immunopathology. (A)** Malpighian
666 tubule (MT) activity ($n = 12-20$ females/infection treatment/age-group), as a proxy for
667 immunopathological damage, measured at 4-hours after infection with 0.1 OD of *P. rettgeri*.
668 Statistically significant difference between groups are indicated by asterisk (*). **(B)** Expression
669 of positive (*Relish*, *PGRP-LC*) and negative (*Caudal*, *Pirk*) regulators of Imd-pathway across
670 sexes and age-groups after *P. rettgeri* infection, relative to an internal control *rp49* ($n =$ Total
671 15-21 flies homogenized in Trizol in a group of 3/Infection treatment/ age-group/ sex-
672 combination). ns = not significant



673

674

675

676 **SUPPLEMENTARY METHODS**

677 **i. Bacterial culture preparations & systemic infections**

678 We used two gram-negative bacterial pathogens *Providencia rettgeri* and *Pseudomonas entomophila*,
679 with a broad host range infecting insects, nematodes, plants and also higher vertebrates (Siva-Jothy
680 et al., 2018; Troha and Buchon, 2019). In fruit flies, *P. rettgeri* and *P. entomophila* infection can result
681 in severe pathology, eventually causing death (Vodovar et al., 2005; Dieppois et al., 2015; Galac and
682 Lazzaro, 2011). In brief, we used overnight grown cultures of *P. rettgeri* & *P. entomophila* at 37°C and
683 30°C respectively with shaking at 120 rpm obtained from pure bacterial isolates.

684 We used LB [Luria Bertani broth; see (Siva-Jothy et al., 2018) for recipe] as a growth media for the
685 bacterial cultures. At the mid-log phase ($OD_{600} = 0.95$), we harvested the bacterial cells by
686 centrifugation at 5000 rpm for 5 min at 4°C and re-suspended the bacterial pellet in 1X PBS (phosphate
687 buffer saline). We adjusted the final inoculum to $OD_{600} = 0.1$ for *P. rettgeri* & 0.05 OD for *P.*
688 *entomophila* for all systemic infections. Briefly, we pricked the adult flies (young 3-5 day old, old 25-
689 28 day old) in the thorax region (Khalil et al., 2015) using 0.1 mm minutein needle (Fine science tools
690 Ltd.). The OD 0.1 *P. rettgeri* and 0.05 OD of *P. entomophila*. For mock infections, we replaced the
691 bacterial cultures with sterile 1x PBS.

692 **ii. Bacterial load measurements**

693 To test whether ageing reduces host's ability to suppress the bacterial growth, we quantified bacterial
694 load in each of the fly lines at 24-hours (or 20-hours) after *Providencia rettgeri* (or *Pseudomonas*
695 *entomophila*) infection across different age groups (3- vs 25-day-old) and sexes (as described in
696 Prakash et al., 2021). First, we thoroughly washed individual flies (pooled in a group of 6 in 1.5ml
697 centrifuge tube) with 70% ethanol for 30-seconds to remove the surface bacteria, followed by rinsing
698 them twice with sterile distilled water. We then homogenized the surface sterilized individuals using
699 a clean motorized pestle for approximately a minute in 180 μ l 1X PBS broth. We performed serial
700 dilution of the fly homogenate up to 10⁻⁵-fold and added 4 μ L of the aliquot onto LB agar plate. We
701 incubated the plates overnight for 18-hours at 30°C and counted the resultant bacterial colonies. None
702 of sham-infected control fly homogenates produced any colonies on LB agar plates.

703 **iii. Assay for the Malpighian tubule activity**

704 Insect Malpighian tubules are functionally analogous to mammalian kidney, playing vital roles in
705 mediating osmoregulation, detoxification, and excretion (Chapman, 1998; Li et al., 2020). Their activity
706 is of particular importance during infection because of their increased vulnerability to damage during
707 immune activation against target pathogens (Sadd and Siva-Jothy, 2006). In fact, previous studies have
708 provided an *in vitro* functional estimate of the ability of isolated tubules to transport saline across the
709 active cell wall into the tubule lumen to demonstrate a large reduction in MT function due to
710 immunopathological damage associated with immune activation in mealworm beetles *Tenebrio*
711 *molitor* (Sadd and Siva-Jothy, 2006; Khan et al., 2017). Here, we used Ramsay assay (as described in
712 Dow et al., 1994; Li et al., 2020) to estimate the fluid transporting capacity of MTs harvested from
713 experimental flies after 4-hours of 0.1 OD *P. rettgeri*, as a proxy for immunopathology owing to
714 immune response. Briefly, the flies were dissected in Schneider's insect medium where intact MTs
715 were first detached from the gut. One end of tubules were immersed in a mixture of Schneider's
716 medium and insect saline (bathing buffer), whereas the other end was pulled out of the buffer and
717 wrapped around a minutein pin (0.1 mm). We allowed the fluid to secrete for 3-hours from the
718 common ureter, following which the volume of secreted fluid was quantified. We considered the
719 volume of the secreted droplet to be negatively correlated with the degree of immunopathological

720 harm to MTs during bacterial infection (Khan et al., 2017). We analysed the MT activity data using a
721 generalised linear model best fitted to quasi-binomial distribution.

722 **iv. Gene expression studies**

723 The expression of major Imd-pathway regulators was quantified by qRT-PCR for both young and older
724 individuals, as described previously in Prakash et al. (2021). We randomly selected a subset of young
725 and old iso-*w¹¹¹⁸* flies infected with 0.1 OD *P. rettgeri* for RNA extraction at 24h post-infection. We
726 randomly removed selected flies (3 flies per vial) at time point 24 hours post exposure to *P. rettgeri*.
727 We then homogenised the whole flies into 1.5 μ l microcentrifuge tubes containing 80 μ l of TRIzol
728 reagent (Ambion, Life Technologies) using sterile micro-pestles (15-21 replicate flies homogenized in
729 a group of 3 flies/ sex/ age-groups). The tubes containing the homogenate were kept frozen at -80°C
730 till we could extract the RNA following the manufacturer's protocol (Zymo research Ltd).

731 After confirming the purity of the eluted samples using a Nanodrop 2000 Spectrophotometer (version
732 3.8.1), we performed the reverse transcription (RT), where cDNA was synthesized from 2 μ l of the
733 eluted RNA using M-MLV reverse transcriptase (Promega) and random hexamer primers, followed by
734 1: 1 dilution in nuclease free water. We then performed quantitative qRT-PCR using an Applied
735 Biosystems machine with a Fast SYBR Green Master Mix (Invitrogen). For each replicate, we set up the
736 10 μ l reaction mix containing 1.5L of 1:1 diluted cDNA, 5 μ l of Fast SYBR Green Master Mix and a 3.5 μ l
737 of a primer stock containing both forward and reverse primer at 1 μ M suspended in nuclease free
738 water (final reaction concentration of each primer was 0.35 μ M; see Table S1 for qPCR primers used
739 in this study). For each cDNA sample across gene of interests, we had two technical replicates. We
740 calculated the mean threshold cycle (Ct) for the further analyses as described in Livak and Schmittgen
741 (2001).

742 We analysed the gene expression data by first calculating the Δ Ct value for the expression of gene of
743 interest vs the house-keeping gene *rp49*, across sexes after *P. rettgeri* (or sham) infection. We then
744 calculated the fold change in gene expression in infected flies relative to the sham-infected flies. We
745 found the data to be non-normally distributed (confirmed using Shapiro-Wilks's test). We thus log-
746 transformed data to fit into a normal distribution and analysed the data using an ANOVA. For each
747 gene of interest, we specified the model as: Fold-change~ Age, with age as fixed effects across genes
748 and sexes separately.

749 **SUPPLEMENTARY FIGURES**

750 **Figure S1.** List of fly lines used in the study including background or control (iso-*w¹¹¹⁸*), fly lines lacking
751 single (individual) or multiple (combinations) of pathway-specific AMP knockouts that are Imd, Toll or
752 co-regulated.

753

- DrosDel iso-*w¹¹¹⁸* control flies
- Δ AMPs = lacking 10/14 known fly AMPs from Defensin, Drosocin, Attacin, Diptericin, Metchnikowin and Drosomycin gene families.

■ Imd-pathway regulated AMPs
■ Toll-pathway regulated AMPs
■ Co-regulated by IMD- & Toll-pathway

Fly lines lacking individual AMPs:

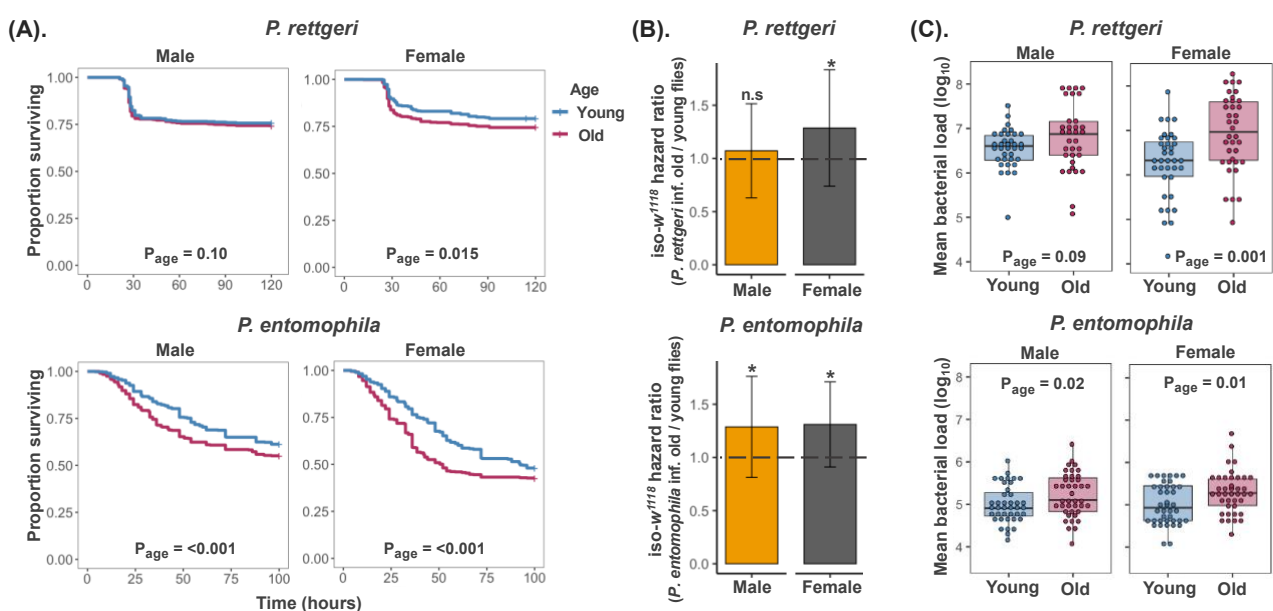
Attacin-C (*AttC^{Mi}*)
Attacin-D (*AttD^{SK3}*)
Diptericin (*Dpt^{SK1}* -*DptA* & *DptB*)
Drososin (*Dro^{SK4}*)
Drosomycin (*Drs^{R1}*)
Metchnikowin (*Mtk^{R1}*)

Fly lines lacking multiple combinations of AMPs:

Dro-Att^{SK2}= *Drosocin* and *AttacinA*, *AttacinB*
Group-A= *Defensin*
Group-B= *Drosocin*, *Diptericin* and *Attacin*
Group-C= *Metchnikowin* and *Drosomycin*
Group-AB=Group A and group B
Group-BC=Group B and group C
Group-AC=Group A and group C

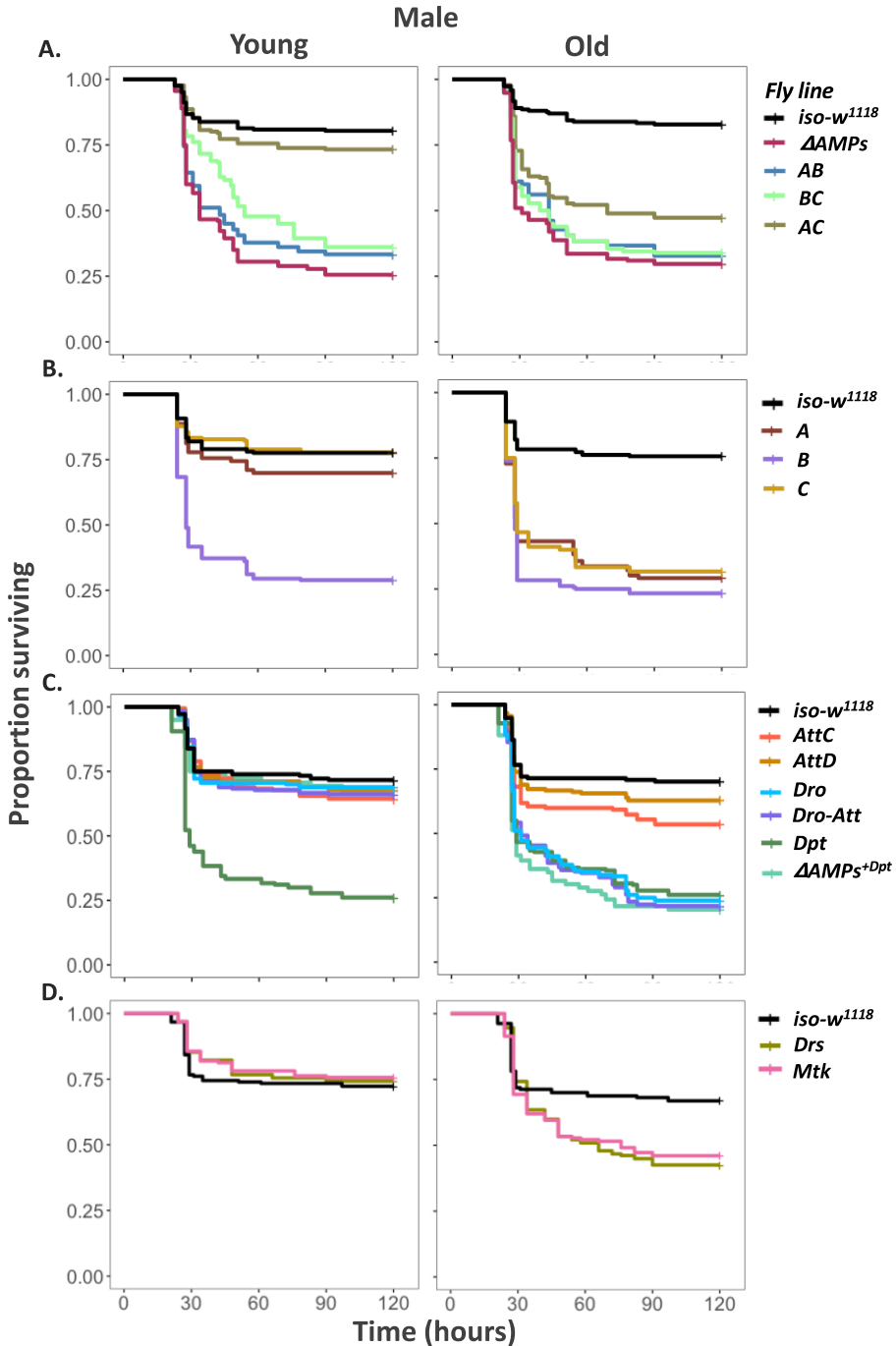
754

755 **Figure S2.** Post-infection survival and bacterial load data of iso- w^{1118} flies across age-groups **(A)**
756 Survival curves for iso- w^{1118} male and females infected with *P. rettgeri* or *P. entomophila* across age-
757 groups (n= 160-280 flies/infection treatment/ age-group/ sex/ replicate experiment). Survival data
758 were analysed with a mixed effects Cox model [model: survival ~ age + (1|food vials + 1| replicate
759 experiments), with 'age' as fixed effects and, 'food vials' and 'replicate experiments' as random
760 effects. **(B)**. Estimated hazard ratios for old vs young iso- w^{1118} males and females. A hazard ratio
761 significantly greater than 1 indicates higher susceptibility of older iso- w^{1118} flies to infection than their
762 younger counterparts. **(C)**. Bacterial load measured as colony forming units (CFUs) after 24- (or 20-)
763 hours of *P. rettgeri* (or *P. entomophila*) infection (n=8-15 replicate groups/treatment/age-groups/sex/
764 pathogen/ replicate experiments). Bacterial load data (log transformed) for each sexes were analysed
765 using a generalized linear mixed model (GLMM) with Gamma distribution, using 'age' and 'replicate
766 experiment' as a 'fixed' and 'random effect' respectively.
767



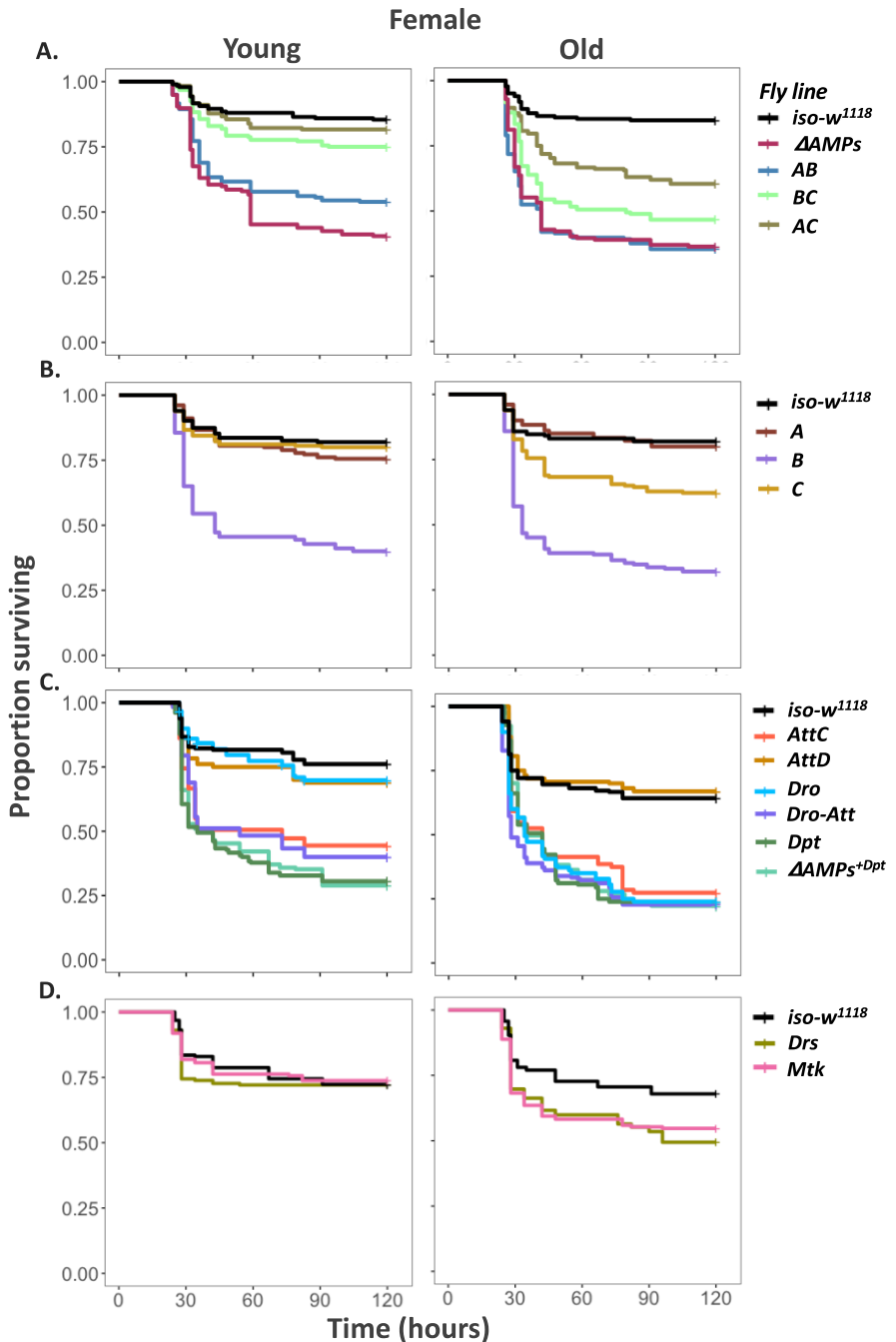
768

769 **Figure S3.** Survival curves for young and old males following *P. rettgeri* infection. Control iso-*w¹¹¹⁸* flies
770 vs (A) Combined mutants and Δ AMPs; (B) Pathway-specific mutants; (C) Single mutants lacking Imd-
771 responsive AMPs (i.e., *Dpt*, *AttC*, *AttD*, *Dro*), Δ AMPs^{+Dpt} and *Dro*-*Att* (D) Single mutants lacking Toll-
772 responsive AMPs (e.g., *Drs* & *Mtk*) (n= 160-180 flies/infection treatment/ fly lines/age combination).



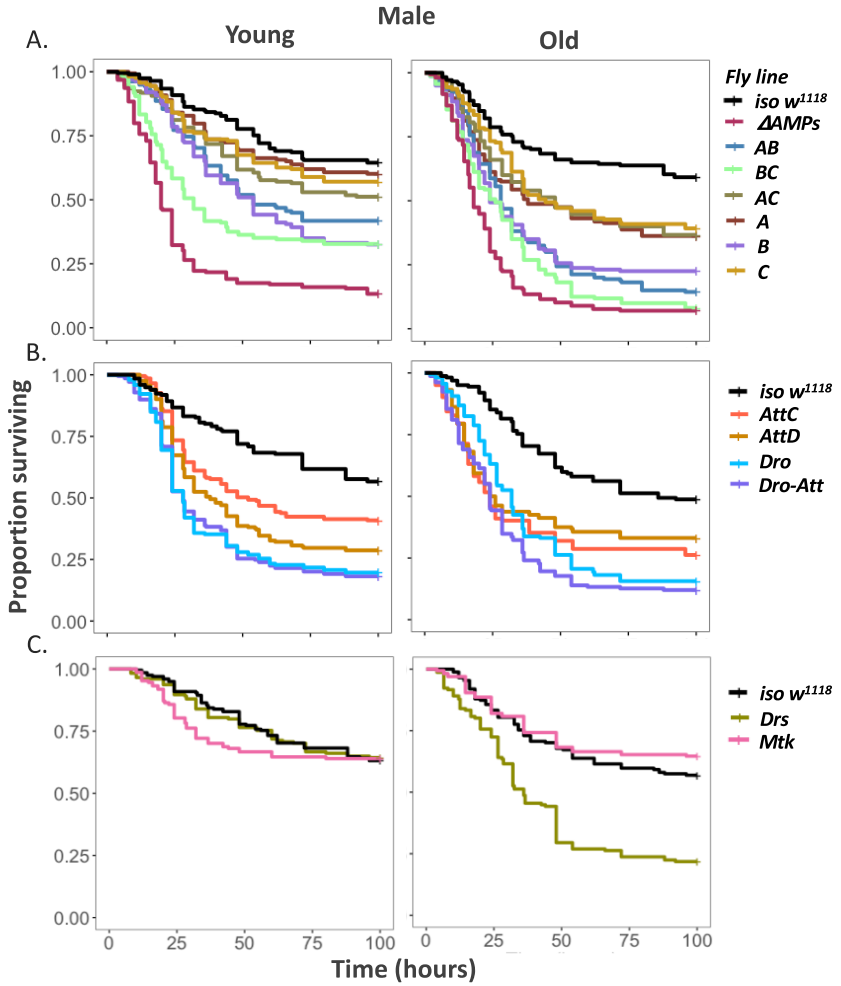
773

774 **Figure S4.** Survival curves for young and old females following *P. rettgeri* infection. Control iso-*w¹¹¹⁸*
775 flies vs **(A)** Combined mutants and Δ AMPs; **(B)** Pathway-specific mutants; **(C)** Single mutants lacking
776 Imd-responsive AMPs (i.e., *Dpt*, *AttC*, *AttD*, *Dro*), Δ AMPs^{+Dpt} and *Dro*-*Att* **(D)** Single mutants lacking
777 Toll-responsive AMPs (e.g., *Drs* & *Mtk*). (n= 160-180 flies/infection treatment/ fly lines/age
778 combination).



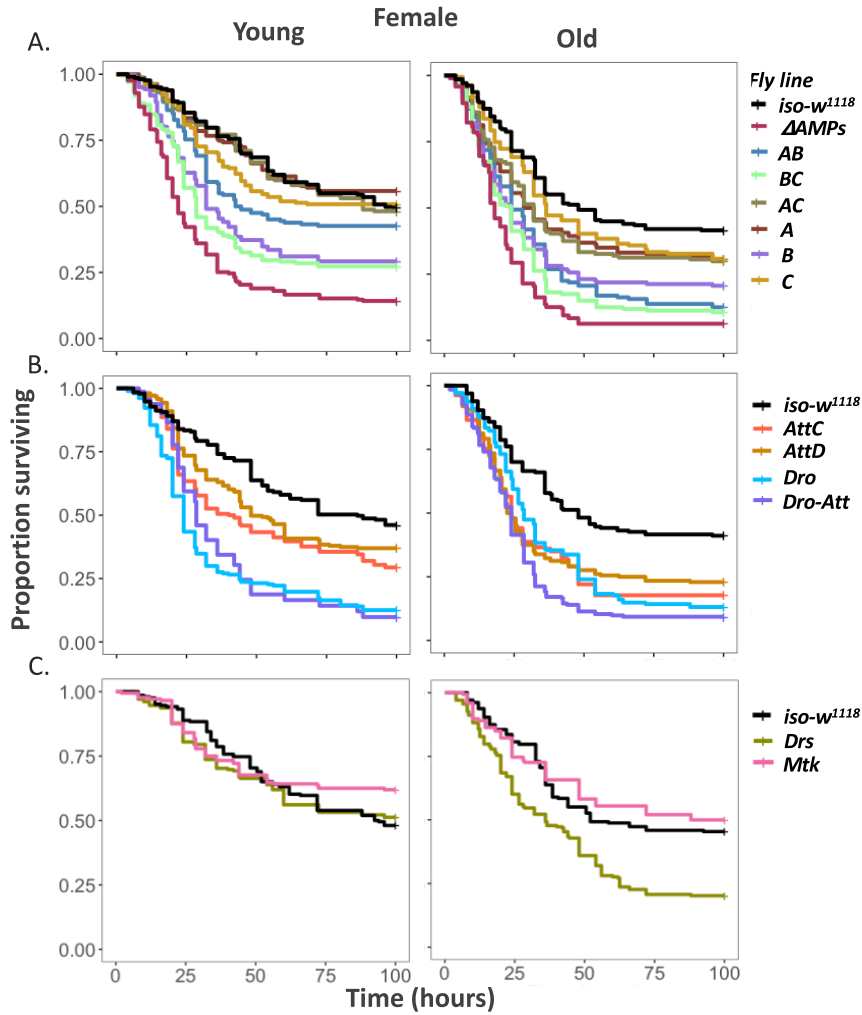
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780

781 **Figure S5.** Survival curves for young and old males following *P. entomophila* infection. Control iso-
782 *w¹¹¹⁸* flies vs (A) Combined and pathway-specific mutants, and Δ AMPs; (B) Single mutants lacking Imd-
783 responsive AMPs (i.e., *AttC*, *AttD*, *Dro*), and *Dro-Att*; (C) Single mutants lacking Toll-responsive AMPs
784 (e.g., *Drs* & *Mtk*) (n= 180-280 flies/infection treatment/ fly lines/age combination).



785

786 **SI Figure 6.** Survival curves for young and old females following *P. entomophila* infection. Control iso-
787 *w¹¹¹⁸* flies vs **(A)** Combined and pathway-specific mutants, and Δ AMPs; **(B)** Single mutants lacking Imd-
788 responsive AMPs (i.e., *AttC*, *AttD*, *Dro*), and *Dro-Att*; **(C)** Single mutants lacking Toll-responsive AMPs
789 (e.g., *Drs* & *Mtk*) (n= 180-280 flies/infection treatment/ fly lines/age combination).



790

791 **SUPPLEMENTARY TABLES**

792 **Table S1:** List of qPCR primers used in the study

	Genes	Sequence	Obtained from
1.	<i>Rp49_F</i>	5' ATGCTAAGCTGCGCACAAATG 3'	(Gupta and Vale, 2017)
	<i>Rp49_R</i>	5' GTTCGATCCGTAACCGATGT 3'	
2.	<i>relish_F</i>	5' CAAGAAGAACGAGGATGCC 3'	(Zaidman-Rémy et al., 2006)
	<i>relish_R</i>	5' GTATACAGGCGGTGCAAGTG 3'	
3.	<i>PGRP-LC_F</i>	5' TTGAACCAAAGTAAGATCAGAGAT 3'	(Zaidman-Rémy et al., 2006)
	<i>PGRP-LC_R</i>	5' GTCCAGATATATTGTTGAATT 3'	
4.	<i>caudal_F</i>	5' CGGCGGATAACTTCGTTCA 3'	(Ryu et al., 2008)
	<i>caudal_R</i>	5' CTGCTGTTGGTCTCGTTGAG 3'	
5.	<i>pirk_F</i>	5' CTCAAGGTGGTCAAGTCCCT 3'	(Kleino et al., 2008)
	<i>pirk_R</i>	5' CTCCGTGCCGTATCGTTAGA 3'	

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794

795 **Table S2.** Summary of mixed effects Cox model to estimate the changes in the survival of individual
796 fly lines with compound AMP mutations (i.e., A, B, C, AB, BC & AC), across sexes and age-groups,
797 relative to control iso-*w¹¹¹⁸* flies after *P. rettgeri* infection. We specified the model as: survival ~ fly
798 line (individual mutant lines vs iso-*w¹¹¹⁸*) + (1|food vials), with 'fly line' as a fixed effect and 'replicate
799 food vials' as a random effect.

800

Sex	Age	Fly lines	Loglik	χ^2	Df	P
Male	Young	<i>ΔAMPs</i>	-929.97	116.98	1	<0.001
			<i>Random effects</i>	<i>Std dev</i>		
			Food vials	0.007		
		AB	-867.68	90.847	1	<0.001
			<i>Random effects</i>	<i>Std dev</i>		
			Food vials	0.0.02		
	Old	BC	-846.22	79.525	1	<0.001
			<i>Random effects</i>	<i>Std dev</i>		
			Food vials	0.17		
		AC	-504.96	2.2842	1	0.13
			<i>Random effects</i>	<i>Std dev</i>		
			Food vials	0.14		
		A	-573.05	2.5879	1	0.10
			<i>Random effects</i>	<i>Std dev</i>		
			Food vials	0.004		
		B	-936.88	104.36	1	<0.001
			<i>Random effects</i>	<i>Std dev</i>		
			Food vials	0.01		
		<i>ΔAMPs</i>	-501.69	0.004	1	1
			<i>Random effects</i>	<i>Std dev</i>		
			Food vials	0.003		
		AB	-743.61	106.3	1	<0.001
			<i>Random effects</i>	<i>Std dev</i>		
			Food vials	0.17		

		Food vials	0.01		
	BC	-814.13	98.733	1	<0.001
		<i>Random effects</i>	<i>Std dev</i>		
		Food vials	0.01		
	AC	-719.61	51.638	1	<0.001
		<i>Random effects</i>	<i>Std dev</i>		
		Food vials	0.004		
	A	-1016.4	87.631	1	<0.001
		<i>Random effects</i>	<i>Std dev</i>		
		Food vials	0.01		
	B	-980.79	101.74	1	<0.001
		<i>Random effects</i>	<i>Std dev</i>		
		Food vials	0.005		
	C	-912.58	80.3	1	<0.001
		<i>Random effects</i>	<i>Std dev</i>		
		Food vials	0.01		
Female	Young	ΔAMPS	-644.46	80.563	1 <0.001
			<i>Random effects</i>	<i>Std dev</i>	
			Food vials	0.02	
		AB	-616.08	45.162	1 <0.001
			<i>Random effects</i>	<i>Std dev</i>	
			Food vials	0.01	
		BC	-434.53	6.3086	1 0.012
			<i>Random effects</i>	<i>Std dev</i>	
			Food vials	0.003	
		AC	-355.40	0.8767	1 0.34
			<i>Random effects</i>	<i>Std dev</i>	
			Food vials	0.01	
		A	-444.25	1.8543	1 0.17
			<i>Random effects</i>	<i>Std dev</i>	
			Food vials	0.007	
		B	-758.83	81.386	1 <0.001
			<i>Random effects</i>	<i>Std dev</i>	
			Food vials	0.01	
		C	-392.30	3.636	1 0.56
			<i>Random effects</i>	<i>Std dev</i>	
			Food vials	0.01	
Old	Old	ΔAMPS	-661.41	90.838	1 <0.001
			<i>Random effects</i>	<i>Std dev</i>	
			Food vials	0.02	
		AB	-764.86	114.51	1 <0.001
			<i>Random effects</i>	<i>Std dev</i>	
			Food vials	0.46	
		BC	-678.05	58.733	1 <0.001
			<i>Random effects</i>	<i>Std dev</i>	
			Food vials	0.11	
		AC	-586.80	28.226	1 <0.001
			<i>Random effects</i>	<i>Std dev</i>	
			Food vials	0.23	
		A	-399.61	0.3776	1 0.53

		<i>Random effects</i>	<i>Std dev</i>		
		Food vials	0.007		
<i>B</i>		-842.26	101.48	1	<0.001
		<i>Random effects</i>	<i>Std dev</i>		
<i>C</i>		Food vials	0.01		
		-571.10	17.644	1	<0.001
		<i>Random effects</i>	<i>Std dev</i>		
		Food vials	0.01		

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802

803 **Table S3.** Summary of a generalized linear mixed model on log transformed bacterial load data
 804 (confirmed and fitted with a gamma distribution) for individual fly lines with compound mutations
 805 (i.e., A, B, C, AB, BC & AC) across sexes and age-groups, relative to control iso-*w¹¹¹⁸* flies after *P. rettgeri*
 806 infection. We specified the model as: log transformed bacterial load ~ fly line (individual mutant lines
 807 vs iso-*w¹¹¹⁸*), with 'fly line' as fixed and 'replicate food vials' as random effects across sexes and age-
 808 groups separately.
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Sex	Age	Fly line	F ratio	Df	P
Male	Young	<i>ΔAMPs</i>	62.2077	1	<0.001
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		<i>AB</i>	221.2613	1	<0.001
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		<i>BC</i>	112.2566	1	<0.001
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		<i>AC</i>	11.9602	1	0.003
	Old	<i>Random effect</i>	<i>std error</i>		
		Food vials	0.0002		
		<i>A</i>	26.7677	1	0.0001
		<i>Random effect</i>	<i>std error</i>		
		Food vials	0.0002		
		<i>B</i>	150.7016	1	<0.001
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		<i>C</i>	65.4964	1	<0.001
		<i>Random effect</i>	<i>std error</i>		
		Food vials	0.0001		
Male	Old	<i>ΔAMPs</i>	21.4858	1	0.0003
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		<i>AB</i>	41.3092	1	<0.001
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		<i>BC</i>	42.3453	1	<0.001
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		<i>AC</i>	14.2853	1	0.001
		<i>Random effect</i>	<i>std error</i>		

		Food vials	<0.001		
		A	15.5048	1	0.001
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		B	13.9127	1	0.001
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		C	9.8094	1	0.006
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
Female	Young	Δ AMPs	19.1118	1	0.0006
		<i>Random effect</i>	<i>std error</i>		
		Food vials	0.0005		
		AB	26.475	1	0.0001
		<i>Random effect</i>	<i>std error</i>		
		Food vials	0.0002		
		BC	2.0876	1	0.16
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		AC	3.4533	1	0.08
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
Female	Old	A	1.3195	1	0.26
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		B	9.8945	1	0.006
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		C	0.7393	1	0.4
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		Δ AMPs	8.1367	1	0.01
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		

	<i>Random effect</i>	<i>std error</i>
Food vials	<0.001	

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Table S4. (A) Summary of the mixed effects Cox model analyses on the age-specific survival data of *iso-w¹¹¹⁸* males and females, infected with *P. rettgeri* and *P. entomophila*. For each pathogen species and sex, we specified the model as: Post-infection survival ~ Age + (1|Replicate experiment), with 'age' as a fixed effect and 'replicate experiment' as a random effect. **(B)** Summary of a generalized linear mixed-effects model, fitted to Gamma distribution, for age-specific bacterial load in *iso-w¹¹¹⁸* males and females, after exposure to *P. rettgeri* and *P. entomophila* infection. For each pathogen species and sex, we specified the model as: Bacterial load ~ Age + (1|Food vial) + (1|Replicate experiment), with 'age' as a fixed effect, 'food vials' and 'replicate experiment' as random effects.

A. Pathogen	Sex	Source	Loglik	Chi sq.	Df	P
<i>P. rettgeri</i>	Male	Age	-2617.6	2.6135	1	0.10
		<i>Random effect</i>	<i>Std dev</i>			
		Food vials	0.14			
		Replicate experiment	0.16			
	Female	Age	-2505.6	5.8313	1	0.015
		<i>Random effect</i>	<i>Std dev</i>			
		Food vials	0.07			
		Replicate experiment	0.17			
<i>P. entomophila</i>	Male	Age	-3083.8	24.269	1	<0.001
		<i>Random effect</i>	<i>Std dev</i>			
		Food vials	0.27			
		Replicate experiment	0.20			
	Female	Age	-4508.0	23.584	1	<0.001
		<i>Random effect</i>	<i>Std dev</i>			
		Food vials	0.19			
		Replicate experiment	0.20			

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B. Pathogen	Sex	Source	F ratio	Df	p
<i>P. rettgeri</i>	Male	Age	2.956	1	0.09
		<i>Random effect</i>	<i>Std dev.</i>		
		Replicate experiment	0.0009		
		Age	12.56		0.0007
	Female	<i>Random effect</i>	<i>Std dev.</i>	1	0.0007
		Replicate experiment	0.0002		
		Age	5.147		0.026
		<i>Random effect</i>	<i>Std dev.</i>		
<i>P. entomophila</i>	Male	Replicate experiment	0.001	1	0.026
		Age	6.628		
		<i>Random effect</i>	<i>Std dev.</i>		
	Female	Replicate experiment	0.0003	1	0.012
		Age	5.147		
		<i>Random effect</i>	<i>Std dev.</i>		

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Table S5. Summary of the mixed effects Cox model analyses to estimate the changes in the survival of individual Imd- and Toll-pathway specific mutants, *ΔAMPs^{ΔDpt}* and *Dro-Att* across sexes and age-groups, relative to control *iso-w¹¹¹⁸* flies after *P. rettgeri* infection. For each fly line across sexes and

829 age-groups, we specified the model as: survival ~ fly line (individual fly lines vs iso-w¹¹¹⁸) + (1|food
 830 vials), with 'fly line' as a fixed effect and 'replicate food vials' as a random effect.
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Sex	Age	Mutants	Loglik	χ^2	Df	P
Male	Young	AttC	-655.69	1.377	1	0.24
			<i>Random effects</i>	<i>Std dev</i>		
		AttD	Food vials	0.003		
			-623.08	0.4223	1	0.51
		Dro	<i>Random effects</i>	<i>Std dev</i>		
			Food vials	0.011		
		Dpt	-612.05	0.3547	1	0.55
			<i>Random effects</i>	<i>Std dev</i>		
		ΔAMPs^{+Dpt}	Food vials	0.01		
			-582.66	0.8597	1	0.35
		Dro-Att	<i>Random effects</i>	<i>Std dev</i>		
			Food vials	0.003		
		Drs	-639.45	0.8147	1	0.36
			<i>Random effects</i>	<i>Std dev</i>		
		Mtk	Food vials	0.003		
			-530.18	0.4169	1	0.51
		Mtk	<i>Random effects</i>	<i>Std dev</i>		
			Food vials	0.003		
Male	Old	AttC	-652.03	9.6893	1	0.001
			<i>Random effects</i>	<i>Std dev</i>		
		AttD	Food vials	0.03		
			-623.07	2.5085	1	0.11
		Dro	<i>Random effects</i>	<i>Std dev</i>		
			Food vials	0.005		
		Dpt	-967.92	65.577	1	<0.001
			<i>Random effects</i>	<i>Std dev</i>		
		ΔAMPs^{+Dpt}	Food vials	0.004		
			-949.55	62.044	1	<0.001
		Dro-Att	<i>Random effects</i>	<i>Std dev</i>		
			Food vials	0.004		
		Drs	-861.39	75.056	1	<0.001
			<i>Random effects</i>	<i>Std dev</i>		
		Mtk	Food vials	0.006		
			-797.95	69.954	1	<0.001
		Drs	<i>Random effects</i>	<i>Std dev</i>		
			Food vials	0.004		
		Mtk	-815.79	12.942	1	0.0003
			<i>Random effects</i>	<i>Std dev</i>		
			Food vials	0.003		
		Mtk	-773.24	10.206	1	0.001

Female	Young	AttC	Random effects	Std dev	
			Food vials	0.004	
		AttD	-789.20	39.231	1 <0.001
			Random effects	Std dev	
		Dro	Food vials	0.006	
			-566.75	2.133	1 0.14
		Dpt	Random effects	Std dev	
			Food vials	0.003	
		ΔAMPs ^{+Dpt}	-548.26	3.3923	1 0.065
			Random effects	Std dev	
		Dro-Att	Food vials	0.01	
			-903.67	76.398	1 <0.001
		Drs	Random effects	Std dev	
			Food vials	0.003	
		Mtk	-829.03	73.907	1 <0.001
			Random effects	Std dev	
		Dro	Food vials	0.004	
			-827.71	48.025	1 <0.001
		Dpt	Random effects	Std dev	
			Food vials	0.01	
		Drs	-573.33	0.0986	1 0.75
			Random effects	Std dev	
		Mtk	Food vials	0.004	
			-536.23	0.0158	1 0.89
		Dro-Att	Random effects	Std dev	
			Food vials	0.004	
Female	Old	AttC	-1117.7	46.083	1 <0.001
			Random effects	Std dev	
		AttD	Food vials	0.005	
			-717.91	0.9219	1 0.33
		Dro	Random effects	Std dev	
			Food vials	0.007	
		Dpt	-1150.0	55.462	1 <0.001
			Random effects	Std dev	
		Drs	Food vials	0.003	
			-1152.8	49.958	1 <0.001
		ΔAMPs ^{+Dpt}	Random effects	Std dev	
			Food vials	0.003	
		Dro-Att	-1059.5	46.363	1 <0.001
			Random effects	Std dev	
		Drs	Food vials	0.003	
			-1155.2	65.681	1 <0.001
		Mtk	Random effects	Std dev	
			Food vials	0.003	
		Dro-Att	-824.05	11.458	1 0.0007
			Random effects	Std dev	
		Drs	Food vials	0.004	
			-771.29	7.4092	1 0.006
		Mtk	Random effects	Std dev	
			Food vials	0.004	

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Table S6. Summary of a generalized linear mixed model on log transformed bacterial load data (confirmed and fitted with a gamma distribution) for individual fly lines with Imd- & Toll-pathway specific, and *Dro-Att* mutations across sexes and age-groups, relative to control iso-*w¹¹¹⁸* flies after *P. rettgeri* infection. We specified the model as: log transformed bacterial load ~ fly line (individual mutant lines vs iso-*w¹¹¹⁸*) with 'fly line' and 'replicate food vials' as fixed and random effects across sexes and age-groups separately.

Sex	Age	Fly line	f ratio	df	p
Male	Young	AttC	1.7874	1	0.2
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		AttD	2.4788	1	0.13
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		Dro	11.0361	1	0.005
		<i>Random effect</i>	<i>std error</i>		
		Food vials	0.0002		
		Dro-Att	10.259	1	0.006
		<i>Random effect</i>	<i>std error</i>		
		Food vials	0.0002		
		Dpt	24.6683	1	0.0002
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		AMPS ^{+Dpt}	0.2392	1	0.63
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		Drs	0.7916	1	0.38
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		Mtk	0.3658	1	0.55
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		AttC	12.4439	1	0.004
Male	Old	<i>Random effect</i>	<i>std error</i>		
		Food vials	0.001		
		AttD	0.144	1	0.71
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		Dro	3.2405	1	0.09
		<i>Random effect</i>	<i>std error</i>		
		Food vials	0.0004		
		Dro-Att	8.1086	1	0.01
		<i>Random effect</i>	<i>std error</i>		
		Food vials	0.0006		
		Dpt	6.4361	1	0.02
		<i>Random effect</i>	<i>std error</i>		
		Food vials	0.0001		
		AMPS ^{+Dpt}	4.3503	1	0.058
		<i>Random effect</i>	<i>std error</i>		

		Food vials	0.0003		
		<i>Drs</i>	23.4568	1	0.0003
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		<i>Mtk</i>	13.1573	1	0.002
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
Female	Young	<i>AttC</i>	6.0366	1	0.02
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		<i>AttD</i>	11.6271	1	0.003
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		<i>Dro</i>	46.0708	1	<0.001
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		<i>Dro-Att</i>	31.1687	1	<0.001
Female	Old	<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		<i>Dpt</i>	53.239	1	<0.001
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		<i>AMPs^{+Dpt}</i>	57.8499	1	<0.001
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		<i>Drs</i>	5.3592	1	0.03
		<i>Random effect</i>	<i>std error</i>		
Female	Old	Food vials	<0.001		
		<i>AttC</i>	0.0098	1	0.92
		<i>Random effect</i>	<i>std error</i>		
		Food vials	0.0005		
		<i>AttD</i>	0.5483	1	0.47
		<i>Random effect</i>	<i>std error</i>		
		Food vials	0.0002		
		<i>Dro</i>	15.2034	1	0.001
		<i>Random effect</i>	<i>std error</i>		
		Food vials	0.0005		
Female	Old	<i>Dro-Att</i>	7.3335	1	0.01
		<i>Random effect</i>	<i>std error</i>		
		Food vials	0.0003		
		<i>Dpt</i>	11.9248	1	0.003
		<i>Random effect</i>	<i>std error</i>		
		Food vials	0.0001		
		<i>AMPs^{+Dpt}</i>	20.1858	1	0.0005
		<i>Random effect</i>	<i>std error</i>		
		Food vials	0.0005		
		<i>Drs</i>	117.8731	1	<0.001
Female	Old	<i>Random effect</i>	<i>std error</i>		

		Food vials	<0.001			
		<i>Mtk</i>	307.8285	1	<0.001	
		<i>Random effect</i>	<i>std error</i>			
		Food vials	<0.001			

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Table S7. Summary of mixed effects Cox model to estimate the changes in the survival of individual fly lines with compound AMP mutations (i.e., A, B, C, AB, BC & AC), across sexes and age-groups, relative to control iso-*w¹¹¹⁸* flies after *P. entomophila* infection. We specified the model as: survival ~ fly line (individual mutant lines vs iso-*w¹¹¹⁸*) + (1|Food vials), with 'fly line' as a fixed effect and 'replicate food vials' as a random effect.

Sex	age	genotype	loglik	χ^2	Df	P
Male	Young	ΔAMPs	-1225.2	153.08	1	0.002
			<i>Random effects</i>	<i>Std dev</i>		
			Food vials	0.0004		
		AB	-895.93	22.54	1	<0.001
			<i>Random effects</i>	<i>Std dev</i>		
			Food vials	0.004		
		BC	-996.87	53.62	1	0.002
			<i>Random effects</i>	<i>Std dev</i>		
			Food vials	0.02		
		AC	-861.94	8.553	1	0.003
			<i>Random effects</i>	<i>Std dev</i>		
			Food vials	0.03		
		A	-732.93	0.1783	1	0.67
			<i>Random effects</i>	<i>Std dev</i>		
			Food vials	0.01		
		B	-981.89	45.449	1	<0.001
			<i>Random effects</i>	<i>Std dev</i>		
			Food vials	0.19		
		C	-791.30	2.6932	1	0.10
			<i>Random effects</i>	<i>Std dev</i>		
			Food vials	0.004		
Male	Old	ΔAMPs	-1090.3	136.08	1	0.00021
			<i>Random effects</i>	<i>Std dev</i>		
			Food vials	0.30		
		AB	-1087	74.174	1	0.0002
			<i>Random effects</i>	<i>Std dev</i>		
			Food vials	0.004		
		BC	-1117.1	109.89	1	0.0002
			<i>Random effects</i>	<i>Std dev</i>		
			Food vials	0.02		
		AC	-898.99	16.873	1	0.003
			<i>Random effects</i>	<i>Std dev</i>		
			Food vials	0.01		
		A	-940.02	20.315	1	<0.001
			<i>Random effects</i>	<i>Std dev</i>		
			Food vials	0.006		
		B	-1033.5	54.008	1	<0.001

				<i>Random effects</i>	<i>Std dev</i>		
				Food vials	0.01		
				<i>C</i>	-980.23	14.057	1 0.0001
				<i>Random effects</i>	<i>Std dev</i>		
				Food vials	0.008		
				-1557.4	104.31	1	0.002
				<i>Random effects</i>	<i>Std dev</i>		
				Food vials	0.003		
				<i>AB</i>	-1116.4	5.432	1 0.019
				<i>Random effects</i>	<i>Std dev</i>		
				Food vials	0.003		
				-1272.0	37.70	1	0.0082
				<i>Random effects</i>	<i>Std dev</i>		
				Food vials	0.005		
				<i>AC</i>	-1072	0.148	1 0.70
				<i>Random effects</i>	<i>Std dev</i>		
				Food vials	0.003		
				-1009.6	1.0044	1	0.31
				<i>Random effects</i>	<i>Std dev</i>		
				Food vials	0.16		
				<i>B</i>	-1210.6	29.765	1 <0.001
				<i>Random effects</i>	<i>Std dev</i>		
				Food vials	0.018		
				-1090.7	2.6955	1	0.10
				<i>Random effects</i>	<i>Std dev</i>		
				Food vials	0.005		
				<i>ΔAMPs</i>	-1580.4	107.24	1 0.0021
				<i>Random effects</i>	<i>Std dev</i>		
				Food vials	0.007		
				<i>AB</i>	-1388.1	48.83	1 0.0002
				<i>Random effects</i>	<i>Std dev</i>		
				Food vials	0.006		
				<i>BC</i>	-1392.6	67.93	1 0.002
				<i>Random effects</i>	<i>Std dev</i>		
				Food vials	0.015		
				<i>AC</i>	-1204.6	8.252	1 0.004
				<i>Random effects</i>	<i>Std dev</i>		
				Food vials	0.003		
				<i>A</i>	-1269.6	7.9978	1 0.004
				<i>Random effects</i>	<i>Std dev</i>		
				Food vials	0.004		
				<i>B</i>	-1336.2	29.789	1 <0.001
				<i>Random effects</i>	<i>Std dev</i>		
				Food vials	0.003		
				<i>C</i>	-1265.7	3.9814	1 0.046
				<i>Random effects</i>	<i>Std dev</i>		
				Food vials	0.004		

851 **Table S8.** Summary of a generalized linear mixed model on log transformed bacterial load data
 852 (confirmed and fitted with a gamma distribution) for individual fly lines with compound mutations
 853 (i.e., A, B, C, AB, BC & AC) across sexes and age-groups, relative to control iso- w^{1118} flies after *P.*
 854 *entomophila* infection. We specified the model as: log transformed bacterial load ~ fly line (individual
 855 mutant lines vs iso- w^{1118}), with 'fly line' as fixed and 'replicate food vials' as random effects across
 856 sexes and age-groups separately.
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Sex	Age	Fly line	f ratio	df	p
Male	Young	AMPs	62.851	1	<0.001
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		AB	13.7047	1	0.001
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		BC	5.4655	1	0.02
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		AC	0.0129	1	0.91
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		A	0.1218	1	0.73
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		B	10.6354	1	0.003
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		C	0.6261	1	0.43
		<i>Random effect</i>	<i>std error</i>		
		Food vials	0.0001		
Male	Old	AMPs	80.1819	1	<0.001
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		AB	7.5861	1	0.01
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		BC	33.3257	1	<0.001
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		AC	0.2875	1	0.59
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		A	0.6311	1	0.43
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		B	8.2924	1	0.008
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		C	1.6474	1	0.21
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
Female	Young	AMPs	62.969	1	<0.001

		<i>Random effect</i>	<i>std error</i>			
		Food vials	<0.001			
		<i>AB</i>	6.3347	1	0.01	
		<i>Random effect</i>	<i>std error</i>			
		Food vials	0.00013			
		<i>BC</i>	6.1418	1	0.02	
		<i>Random effect</i>	<i>std error</i>			
		Food vials	0.0001			
		<i>AC</i>	9.2819	1	0.005	
		<i>Random effect</i>	<i>std error</i>			
		Food vials	0.0003			
		<i>A</i>	7.6082	1	0.01	
		<i>Random effect</i>	<i>std error</i>			
		Food vials	<0.001			
		<i>B</i>	1.1206	1	0.3	
		<i>Random effect</i>	<i>std error</i>			
		Food vials	<0.001			
		<i>C</i>	7.7927	1	0.01	
		<i>Random effect</i>	<i>std error</i>			
		Food vials	<0.001			
Female	Old	<i>AMPs</i>	3.5078	1	0.0002	
		<i>Random effect</i>	<i>std error</i>			
		Food vials	0.0001			
		<i>AB</i>	10.2129	1	0.004	
		<i>Random effect</i>	<i>std error</i>			
		Food vials	<0.001			
		<i>BC</i>	8.5458	1	0.007	
		<i>Random effect</i>	<i>std error</i>			
		Food vials	<0.001			
		<i>AC</i>	5.7803	1	0.02	
		<i>Random effect</i>	<i>std error</i>			
		Food vials	<0.001			
		<i>A</i>	0.0586	1	0.81	
		<i>Random effect</i>	<i>std error</i>			
		Food vials	<0.001			
		<i>B</i>	3.6222	1	0.07	
		<i>Random effect</i>	<i>std error</i>			
		Food vials	<0.001			
		<i>C</i>	1.1598	1	0.3	
		<i>Random effect</i>	<i>std error</i>			
		Food vials	0.0001			

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Table S9. Summary of the mixed effects Cox model analyses to estimate the changes in the survival of individual Imd- and Toll-pathway specific and *Dro-Att* mutations across sexes and age-groups, relative to control iso-*w¹¹¹⁸* flies after *P. entomophila* infection. For each fly line across sexes and age-groups, we specified the model as: survival ~ fly line (individual fly lines vs iso-*w¹¹¹⁸*) + (1|food vials), with 'fly line' as a fixed effect and 'replicate food vials' as a random effect.

Sex	age	genotype	loglik	<i>x²</i>	Df	P
Male	Young	AttC	-1156.1	13.81	1	0.0002

			<i>Random effects</i>	<i>Std dev</i>	
			Food vials	0.004	
<i>AttD</i>			-1266.4	41.284	1 <0.001
			<i>Random effects</i>	<i>Std dev</i>	
<i>Dro</i>			Food vials	0.004	
			-1293.2	82.966	1 <0.001
<i>Dro-Att</i>			<i>Random effects</i>	<i>Std dev</i>	
			Food vials	0.31	
<i>Drs</i>			-1388.2	84.631	1 <0.001
			<i>Random effects</i>	<i>Std dev</i>	
<i>Mtk</i>			Food vials	0.003	
			-709.92	0.2541	1 0.61
<i>Old</i>	<i>AttC</i>		<i>Random effects</i>	<i>Std dev</i>	
			Food vials	0.0	
<i>AttD</i>			-1014.8	21.112	1 <0.001
			<i>Random effects</i>	<i>Std dev</i>	
<i>Dro</i>			Food vials	0.002	
			-1132.5	52.959	1 <0.001
<i>Dro-Att</i>			<i>Random effects</i>	<i>Std dev</i>	
			Food vials	0.005	
<i>Drs</i>			-1135.7	78.785	1 <0.001
			<i>Random effects</i>	<i>Std dev</i>	
<i>Mtk</i>			Food vials	0.008	
			-1044.2	45.125	1 <0.001
<i>Female</i>	<i>Young</i>	<i>AttC</i>	<i>Random effects</i>	<i>Std dev</i>	
			Food vials	0.001	
<i>AttD</i>			-750.24	1.612	1 0.20
			<i>Random effects</i>	<i>Std dev</i>	
<i>Dro</i>			Food vials	0.004	
			-1329.8	15.971	1 <0.001
<i>Dro-Att</i>			<i>Random effects</i>	<i>Std dev</i>	
			Food vials	0.003	
<i>Drs</i>			-1330.3	5.9269	1 0.014
			<i>Random effects</i>	<i>Std dev</i>	
<i>Female</i>	<i>Young</i>	<i>AttD</i>	Food vials	0.01	
			-1524.5	84.181	1 <0.001
<i>Dro</i>			<i>Random effects</i>	<i>Std dev</i>	
			Food vials	0.01	
<i>Dro-Att</i>			-1643.9	87.659	1 <0.001
			<i>Random effects</i>	<i>Std dev</i>	
<i>Drs</i>			Food vials	0.004	
			-1182.1	0.0018	1 1
<i>Female</i>	<i>Young</i>		<i>Random effects</i>	<i>Std dev</i>	
			Food vials	0.004	

			<i>Mtk</i>	-985.38	3.7318	1	0.054
			<i>Random effects</i>	<i>Std dev</i>			
			Food vials	0.004			
Female	Old	<i>AttC</i>	-1453.1	33.386	1	<0.001	
			<i>Random effects</i>	<i>Std dev</i>			
		<i>AttD</i>	Food vials	0.004			
			-1441.2	23.373	1	<0.001	
		<i>Dro</i>	<i>Random effects</i>	<i>Std dev</i>			
			Food vials	0.003			
		<i>Dro-Att</i>	-1435.7	31.6	1	<0.001	
		<i>Random effects</i>	<i>Std dev</i>				
		<i>Food vials</i>	0.004				
		<i>Drs</i>	-1546.7	69.786	1	<0.001	
			<i>Random effects</i>	<i>Std dev</i>			
			Food vials	0.008			
		<i>Mtk</i>	-1515.5	28.931	1	<0.001	
			<i>Random effects</i>	<i>Std dev</i>			
			Food vials	0.004			
		<i>Random effects</i>	<i>Std dev</i>				
		Food vials	0.003				

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868 **Table S10.** Summary of a generalized linear mixed model on log transformed bacterial load data
869 (confirmed and fitted with a gamma distribution) for individual fly lines with Imd- & Toll-pathway
870 specific, and *Dro-Att* mutations across sexes and age-groups, relative to control iso-*w¹¹¹⁸* flies after *P.*
871 *entomophila* infection. We specified the model as: log transformed bacterial load ~ fly line (individual
872 mutant lines vs iso-*w¹¹¹⁸*), with 'fly line' as fixed and 'replicate food vials' as random effects across
873 sexes and age-groups separately.

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Sex	Age	Fly line	f ratio	df	p
Male	Young	<i>AttC</i>	5.4807	1	0.02
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		<i>AttD</i>	7.3457	1	0.01
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		<i>Dro</i>	10.9639	1	0.003
		<i>Random effect</i>	<i>std error</i>		
		Food vials	0.0001		
		<i>Dro-Att</i>	7.4034	1	0.01
		<i>Random effect</i>	<i>std error</i>		
		Food vials	0.0001		
		<i>Drs</i>	4.0698	1	0.054
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		<i>Mtk</i>	0.1824	1	0.67
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
Male	Old	<i>AttC</i>	0.8264	1	0.37

		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		AttD	2.4385	1	0.13
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		Dro	0.704	1	0.4
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		Dro-Att	7.8316	1	0.009
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		Drs	1.2319	1	0.27
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		Mtk	0.2088	1	0.65
		<i>Random effect</i>	<i>std error</i>		
		Food vials	0.0001		
Female	Young	AttC	3.5928	1	0.07
		<i>Random effect</i>	<i>std error</i>		
		Food vials	0.0004		
		AttD	8.6402	1	0.007
		<i>Random effect</i>	<i>std error</i>		
		Food vials	0.0002		
		Dro	19.452	1	0.0002
		<i>Random effect</i>	<i>std error</i>		
		Food vials	0.0002		
		Dro-Att	2.711	1	0.11
		<i>Random effect</i>	<i>std error</i>		
		Food vials	0.0001		
		Drs	1.4662	1	0.23
		<i>Random effect</i>	<i>std error</i>		
		Food vials	0.0001		
		Mtk	4.0912	1	0.055
		<i>Random effect</i>	<i>std error</i>		
		Food vials	0.0001		
Female	Old	AttC	1.1095	1	0.3
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		AttD	0.3591	1	0.55
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		Dro	1.0402	1	0.31
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		Dro-Att	7.0194	1	0.01
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		Drs	0.7705	1	0.38
		<i>Random effect</i>	<i>std error</i>		
		Food vials	<0.001		
		Mtk	0.7199	1	0.4

	<i>Random effect</i>	<i>std error</i>
Food vials		<0.001

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Table S11. Summary of a generalized linear model (best fitted to a quasi-binomial distribution) for Malpighian tubule activity as a function of infection status across age-groups. For each age group, we specified the model as Malpighian tubule activity~ Infection status, with infection status as a fixed effect across age-groups.

Age	Df	χ^2	P
Young	1	0.0814	0.77
Old	1	6.179	0.012

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Table S12. Summary of ANOVA on the gene expression data in males and females. For each sex and genes, we specified the model as: Fold-change in gene expression ~ age, with 'age' as a fixed effect.

Gene	Sex	Df	SS	F-ratio	P
<i>Relish</i>	Male	1	0.004	0.003	0.95
	Female	1	2.64	0.594	0.45
<i>PGRP-LB</i>	Male	1	3.65	0.347	0.56
	Female	1	2.64	0.594	0.45
<i>Caudal</i>	Male	1	119.01	60.12	<0.001
	Female	1	23.75	53.97	<0.001
<i>Pirk</i>	Male	1	3.506	15.04	0.002
	Female	1	24.36	8.708	0.01

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