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Instructions for use

1	No evidence for contribution of sexually monomorphic wing pigmentation pattern
2	to mate choice in Drosophila guttifera
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14 Abstract

In many animal groups, sexually dimorphic ornaments are thought to evolve by 15 intraspecific competition or mate choice. Some researchers pointed out that sexually 16 monomorphic ornaments could also evolve by mate choice by both sexes or either sex. 17 18 Many species of fruit fly have sexually monomorphic wing pigmentation. However, involvement of their sexually monomorphic ornaments in mate choice has not been tested. 19 We aimed to examine whether the sexually monomorphic polka-dotted pattern on wings 20 21 of Drosophila guttifera contributes to mate choice. Because D. guttifera does not mate in the dark condition at all and no courtship sound has been observed, some visual 22 23 information is likely to be used in mating behaviour. We compared the number of matings 24 between individuals with and without wings, and found that presence of wings influenced mating behaviour in both sexes. We then compared the number of matings between 25 individuals bearing replaced wings, one group bearing conspecific *D. guttifera* wings and 26 another group bearing heterospecific D. melanogaster wings with no pigmentation 27 28 pattern. An effect of conspecific/heterospecific wings was only detected in mate choice by females. Comparison between wild-type and black-painted wings revealed no 29 evidence of a contribution of wing pigmentation pattern to mate choice in either sex. 30

31 Introduction

32 Evolution of sexually dimorphic ornaments has been explained in the theoretical 33 framework of sexual selection, such as Fisher's runaway process and the handicap principle (Fisher, 1915; Zahavi, 1975). In many animal groups, sexually dimorphic 34 35 ornaments were suggested to evolve by intrasexual competition and mate choice (Petrie, 36 Halliday, & Sanders, 1991; Petrie & Halliday, 1994; Andersson, 1982; Ryan, 1985; Theis, Salzburger, & Egger, 2012). Some researchers argued that sexually monomorphic 37 38 ornaments could evolve before sexually dimorphic ornaments under the evolutionary restriction of sexual dimorphism by genetic correlation or constraint between the two 39 40 sexes, and that dimorphism evolves by mutations enabling circumvention of the genetic 41 constraint on a long timescale (Lande, 1980). The details of genetic constraints are now 42 understood according to a concept of intralocus sexual conflict (Bonduriansky & Chenoweth, 2009). For example, king penguins, Aptenodytes patagonicus, have sexually 43 monomorphic ornaments, but female mate choice was observed to be stronger than male 44 45 mate choice (Pincemy, Dobson, & Jouventin, 2009). This suggests that sexually 46 monomorphic ornaments could evolve by a mate choice by either sex. As a counter explanation, the same traits could be involved in mate choices of both sexes that result in 47 evolution of a sexually monomorphic ornament (Kraaijeveld, Kraaijeveld-Smit, & 48 Komdeur, 2007). 49

50 In Drosophila (fruit flies), there are species with various pigmentation patterns 51 on wings (Koshikawa, 2020). Many species of *Drosophila* are known to have malespecific black pigmentation on the anterior-distal part of wings (Kopp & True, 2002). 52 Males of these species appear to display their wings in front of females (Prud'homme et 53 al., 2006). However, an effect of wing pigmentation on mate choice was not always 54 55 observed. Using three species with sexually dimorphic black spots on male wings 56 (Drosophila suzukii, D. biarmipes and D. subpulchrella), Roy and Gleason (2019) 57 examined whether females prefer males with spots or males without spots (that were made by CO₂ anaesthesia after eclosion). An effect of spots on mate preference was not 58 detected in their study. Fuyama (1979) revealed that males without spots as a result of 59 amputation showed lower mating frequency than intact males in D. suzukii when females 60 were kept in constant light to make them less accepting of mating. In D. biarmipes, which 61 has natural polymorphism of wing pigmentation, males with pigmentation on their wings 62 showed greater mating success than males without pigmentation (Hegde, Chethan, & 63 64 Krishna, 2005; Singh & Chatterjee, 1987). The effect of pigmentation was dependent on environmental conditions (Parkash, Lambhod, & Singh, 2013). 65 66

Despite having sexually monomorphic pigmentation on their wings, some fruit

fly males display their wings in front of females during their courtship. For example, in 67 Idiomyia grimshawi (synonym of Drosophila grimshawi), adults aggregate in leks and 68 males display with their elaborately pigmented wings (Spieth, 1966; Edwards, Doescher, 69 Kaneshiro, & Yamamoto, 2007). However, the function of this sexually monomorphic 70 71 pigmentation during courtship has not been studied experimentally. Many species of fruit 72 flies are known to have sexually monomorphic wing pigmentations (Patterson, 1943; Miller, Marshall, & Grimaldi, 2017; Werner, Steenwinkel, & Jaenike, 2020; Dufour, 73 74 Koshikawa, & Finet, 2020), but no functional testing of these monomorphic traits has 75 been reported.

76 Drosophila guttifera has been used as a research model for elucidating 77 pigmentation pattern formation (Werner, Koshikawa, Williams, & Carroll, 2010; 78 Koshikawa et al., 2015; Koshikawa, Fukutomi, & Matsumoto, 2017; Fukutomi, 79 Matsumoto, Agata, Funayama, & Koshikawa, 2017; Dion et al. 2020; Shittu, Steenwinkel, Koshikawa, & Werner 2020). This species has a sexually monomorphic polka-dotted 80 81 pattern on its wings, but the function of the pattern has not been explored. Because D. guttifera does not mate at all in a dark condition, visual information is likely to be used 82 83 in mating (Grossfield, 1966). In addition, courtship sound, such as wing vibration, was 84 not observed in this species (Spieth, 1952; Grossfield, 1977; Wen & Li, 2011).

The purpose of this study was to clarify whether the sexually monomorphic pigmentation pattern in *D. guttifera* contributes to mate choice. We examined the effect of the presence of wings and the polka-dotted pattern on wings on mate choice of both sexes by conducting mating experiments with cutting and replacing of wings.

89

90 Materials and methods

91 Fly

92 Drosophila guttifera inhabits North America and is related to the quinaria group 93 (Chialvo, White, Reed, & Dyer, 2019; Izumitani, Kusaka, Koshikawa, Toda, & Katoh, 2016). In this study, adults of the wild type D. guttifera (stock number 15130-1971.10, 94 provided by the Drosophila Species Stock Center at the University of California, San 95 Diego) kept in our laboratory were used. In addition to intact adults ("wild type"), adults 96 97 whose wings were cut within 24 hours after eclosion ("no wing") and adults whose wings were painted black 9-12 days after eclosion ("black wing") were prepared. Also, adults 98 99 whose wings were replaced by D. guttifera wings ("guttifera wing"), wings with an incomplete pigmentation pattern ("incomplete pattern"), wings of Drosophila 100 melanogaster Oregon-R ("melanogaster wing"), and black-painted wings of D. 101 102 melanogaster ("melanogaster black wing") were prepared and then maintained for 14-87

hours before experiments (the recipients were wild type D. guttifera 5-11 days after 103 104 eclosion). These flies were used in mate choice experiments (Figure 1). The strain with 105 an incomplete pigmentation pattern has a recessive mutation, but the genetic lesion of this mutation is unknown (Wataru Yamamoto, personal communication). Wings were 106 107 replaced using a cyanoacrylate adhesive (Aron Alpha, Konishi Co., Ltd., Japan). To 108 produce "black wing" flies, flies' entire wings were painted with a black ink marker (Mckie, Zebra Co., Ltd., Japan). All individuals were reared with standard 109 110 cornmeal/sugar/yeast/agar food (Fukutomi, Matsumoto, Funayama, & Koshikawa, 2018) under a photoperiod of 12:12 (light:dark) at 25°C. 111

- 112
- 113 Sexual maturity after eclosion

114 To prepare virgin individuals for mating experiments, we had to confirm that adults were not sexually mature within 24 hours after eclosion. Firstly, males and females 115 were separated into different vials within 4 hours after eclosion. Some vials were kept 7-116 117 10 days after eclosion. Four females 7–10 days after eclosion and four males 4 hours after eclosion were introduced per one vial (n=5). Likewise, four males 7-10 days after 118 119 eclosion and four females 4 hours after eclosion were introduced per one vial (n=6). In a control group, four females and four males 7–10 days after eclosion were introduced per 120 121 one vial (n=4). These three groups of vials were kept for 24 hours, and then adults in the 122 vials were removed. The number of larvae in the vials was counted 5-7 days after adults 123 had been removed.

For mate choice experiments, adult males and females within 24 hours after eclosion were collected and kept in groups of 1–20 adults of the same sex. Mate choice experiments were conducted 5–13 days after eclosion.

127

128 Recording of mating behaviour

All mate choice experiments were conducted in chambers made of acrylic resin 129 130 (10 mm in diameter and 5 mm in height). A small piece of fly food was placed in each chamber to prevent drying of flies during experiments. The chambers were put on a piece 131 132 of white paper, and then adults used in experiments were introduced into the chambers. 133 The chambers were then illuminated with a halogen lamp (PICL-NEX, Nippon P·I, Co., Ltd., Japan) and recorded with a web camera (C615n, Logicool Co., Ltd., Japan) or a 134 digital camera (HDR-PJ590V, Sony, Japan) continuously for 3 hours. The start time of 135 the recording was between 90 and 150 minutes after the light phase of the original rearing 136 condition. 137

138

139 Preference for wings

In a male mate choice experiment (Table 1; competition experiment with one 140 141 male and two females per one chamber) to examine the preference for wings, we observed whether males mated with "wild type" females or "no wing" females. The kinds of mating 142 143 were classified into four types—the first mating between one male and one "wild type" 144 female, the second mating between them, the first mating between one male and one "no wing" female, and the second mating between them. As another index of the preference 145 146 for wings, we observed whether males courted females and which female was courted 147 first. Also, we measured courtship latency and courtship duration and compared them. 148 We defined two types of courtship latencies and two types of courtship durations. Latency 149 1: the length of time (min) from the start of recording to the first courtship. Latency 2: the length of time from the start of recording to the first successful courtship followed by 150 mating. Duration 1: the length of the first courtship. Duration 2: the length of the first 151 152 successful courtship followed by mating. Latency 1 and Duration 1 mostly reflect only 153 male's preference. Latency 2 and Duration 2 reflect females' eagerness to mate as well as 154 male's preference, since they depend on the length of time until females accept to be 155 courted and mate. Latency 1 could be the same as Latency 2, and Duration 1 could be the same as Duration 2, when successful courtship followed by mating was immediately 156 observed after the start of recording. 157

158 In a female choice experiment (Table 1; competition experiment with one female 159 and two males per one chamber) to examine the preference for wings, we observed whether females mated with "wild type" males or "no wing" males. In this experiment, 160 161 the difference between the number of matings achieved by two males may be influenced 162 by female preference and also males' eagerness to mate. To estimate males' eagerness to 163 mate, we observed whether males courted females and which male courted first. Also, we 164 measured the courtship latency (Latency 1: the length of time from the start of recording 165 to the first courtship).

166

167 Preference for conspecific wings

In a male choice experiment (Table 1; competition experiment) to examine the preference for the polka-dotted pattern, we observed whether males mated with "*guttifera* wing" females or "*melanogaster* wing" females. The kinds of mating were classified into four types—the first mating between one male and one "*guttifera* wing" female, the second mating between them, the first mating between one male and the one "*melanogaster* wing" female, and the second mating between them. As another index of the preference for the polka-dotted pattern, we observed whether males courted females and which female was courted first. Also, courtship latencies (Latency 1 and 2) and
courtship durations (Duration 1 and 2) were measured.

In a female choice experiment (Table 1; competition experiment) to examine the preference for the polka-dotted pattern, we observed whether females mated with "*guttifera* wing" males or "*melanogaster* wing" males. To estimate males' eagerness to mate, we observed whether males courted females and which male courted first. Also, we measured the courtship latency (Latency 1).

182

183 Preference for the polka-dotted pattern

184 In a female choice experiment (Table 1; competition experiment) to examine the 185 preference for the precise number of polka dots, we observed whether females mated with 186 "guttifera wing" males or "incomplete pattern" males. Also, to examine the preference for dark colour of the entire wings, we observed whether females mated with "wild type" 187 males or "black wing" males. Finally, we observed whether females mated with 188 189 "melanogaster wing" males or "melanogaster black wing" males. In these three experiments, to estimate males' eagerness to mate, we observed whether males courted 190 191 females and which male courted first. Also, we measured the courtship latency (Latency 192 1).

193

194 Statistical analysis

195 All statistical analyses were performed using R ver. 3.5.2 (R Core Team, 2018). 196 In both male and female mate choice experiments, GLMs were used to investigate the 197 influence of the mating combination between the sexes, male age, female age and hours 198 after females had received replaced wings (if the experiment included wing replacement) 199 on the occurrence of mating. We included ages and hours after wing replacements in this 200 analysis, because ages and CO₂ anaesthesia are known to affect the behavior of some 201 Drosophila species (Avent, Price & Wedell, 2008; Colinet & Renault, 2012; Verspoor, 202 Cuss & Price, 2015). The occurrence of mating was treated as a response variable 203 assuming a binomial distribution. Mating combination, male age, female age and hours 204 after females had received replaced wings were treated as explanatory variables. As a link 205 function, logit function was used. In addition to GLMs, chi-squared tests were performed on the mating combination between the sexes without variables of ages. For the three 206 experiments to test the preference for the polka-dotted pattern, Bonferroni correction with 207 chi-squared test was performed to counteract the problem of multiple similar experiments 208 (Wright, 1992). When sample size was less than 40, Yates correction with the chi-squared 209 210 test was performed (Campbell, 2007). For comparison of courtship latencies and 211 courtship durations, we used the Student's *t*-test. To determine which type of female was

212 courted first and which type of male courted first, we used the exact binomial test.

213

214 **Results**

215 The courtship behaviour of *D. guttifera*

216 The courtship behaviour of both males and females was observed. Firstly, a male approached a female from behind, the side or the front. Next, the male chased the female 217 218 if the female ran away from the male. Then, the male tapped female body parts (mostly 219 top of the wing or head) with its foreleg. Next, the male extended its proboscis and licked 220 the female's ovipositor. Following this behaviour, the male intensely vibrated its 221 proboscis toward the female. No wing motion (extension, vibration, scissoring, flicking 222 or rotation) of the male was observed. A receptive female spread its wings widely and 223 then the male mounted the female. This courtship behaviour seems to be identical to those 224 observed in previous studies (Spieth, 1952; Grossfield, 1966; Grossfield, 1977).

225

226 Sexual immaturity at 24 hours after eclosion

Four males 7–10 days after eclosion and four females 4 hours after eclosion were introduced per vial and were removed from the vial after 24 hours. After 5–7 days, no larvae were observed in the vials (n=5). Also, no larvae were observed in vials into which four females 7–10 days after eclosion and four males 4 hours after eclosion were introduced and then removed (n=6). In contrast, the number of larvae was 15.8 ± 11.0 (mean \pm SD [standard deviation], n=4) in vials into which four males and four females (all 7–10 days after eclosion) were introduced and then removed.

We concluded that neither males nor females become sexually mature within 24 hours after eclosion. Therefore, in the following experiments, the adults within 24 hours after eclosion were collected as virgins. We kept virgins for more than 5 days in groups and used them in mate choice experiments.

238

239 Preference for wings

For male choice experiments, we conducted 51 experiments to examine preference for wings (Figure 2a). Some males mated with both of the two females. Such matings were recorded separately as the first mating and the second mating. Males never mated twice with the same female in our observations. In the first matings, males mated with "wild type" females 25 times and with "no wing" females 10 times. In the second matings, males mated with "wild type" females four times and with "no wing" females 11 times. Thus, at the first mating, males mated with "wild type" at higher rate than with ²⁴⁷ "no wing", and at the second mating males mated with "no wing" females at higher rate ²⁴⁸ than with "wild type" females (GLM, estimate=-2.7865, p=0.000682; chi-squared test, ²⁴⁹ p=0.0006539). Other explanatory variables were not statistically significant (GLM, ²⁵⁰ p>0.05).

251 We observed courtships as another index of preference for wings in these 51 252 experiments. Males courted females in 47 experiments in total. Among 15 experiments in which the male mated with both females, we observed courting both females in all 15 253 254 experiments. Among 20 experiments in which the male mated with one female, we 255 observed courting both females in 16 experiments and courting one female in four 256 experiments. Among 16 experiments in which the male did not mate, we observed 257 courting both females in eight experiments, courting one female in four experiments, and no courting in four experiments. Males courted first "wild type" females 29 times and 258 "no wing" females 18 times, which were not significantly different (exact binomial test, 259 p=0.1439). Courtship latencies were not significantly different between courtships with 260 261 "wild type" females (Latency 1, 29.5 ± 36.0 min [mean \pm SD]; Latency 2, 43.3 ± 35.5 min) and courtships with "no wing" females (Latency 1, 25.9 ± 25.8 min; Latency 2, 62.9262 263 \pm 47.4 min) (*t*-test, *p*=0.6054 and *p*=0.1067). Courtship durations were significantly different between courtships with "wild type" females (Duration 1, 0.49 ± 0.69 min; 264 Duration 2, 0.69 \pm 0.89 min) and courtships with "no wing" females (Duration 1, 1.19 \pm 265 266 2.18 min; Duration 2, 2.32 \pm 2.41 min) (*t*-test, *p*=0.04862 and *p*=0.001912). The significant difference of these courtship durations may indicate the difference of female's 267 268 receptive behaviour, and it could be a part of the reason why "no wing" females mated 269 less than "wild type" females (see Discussion).

For female choice experiments, we conducted 51 experiments to examine preference for wings (Figure 2b). Females mated with "wild type" males 27 times and with "no wing" males 15 times. Females never mated twice with the same male in our observations. Females mated with "wild type" males more frequently than with "no wing" males (GLM, estimate=1.0313, p=0.0158; chi-squared test, p=0.01577). Other explanatory variables were not statistically significant (GLM, p>0.05).

We observed courtships to estimate males' eagerness to mate in these 51 experiments. Males courted females in all 51 experiments. Among 42 experiments with successful matings, both males courted in 38 experiments and one male courted in four experiments. Among nine experiments without matings, both males courted in eight experiments and one male courted in one experiment. Females were courted first by "wild type" males in 22 experiments and by "no wing" males in 29 experiments, which were not significantly different (exact binomial test, p=0.4011). The courtship latency (Latency 1) of "wild type" males $(15.8 \pm 21.6 \text{ min})$ and that of "no wing" males $(18.0 \pm 22.0 \text{ min})$ were not significantly different (*t*-test, *p*=0.6192). Therefore, there was no evidence of a difference in males' eagerness to mate. In summary, females were found to prefer males with wings to males without wings.

- 287
- 288 Preference for conspecific wings

For male choice experiments, we conducted 50 experiments to examine the 289 290 preference for the polka-dotted pattern (Figure 3a). In the first matings, males mated with "guttifera wing" females 19 times and "melanogaster wing" females 14 times. In the 291 second matings, males mated with "guttifera wing" females eight times and D. 292 293 melanogaster wing females nine times. No significant difference was seen between the rate of mating of males with "guttifera wing" wing females, and that of males with 294 "melanogaster wing" females (GLM, estimate=0.24864, p=0.544; chi-squared test, 295 p=0.4237). Other explanatory variables were also not statistically significant (GLM, 296 297 *p*>0.05).

We observed courtships as another index of preference for the polka-dotted 298 299 pattern in these 50 experiments. Males courted females in 48 experiments in total. Among 300 17 experiments in which the male mated with both females, we observed courting both 301 females in all 17 experiments. Among 16 experiments in which the male mated with one 302 female, we observed courting both females in 13 experiments and courting one female in 303 three experiments. Among 17 experiments in which the male did not mate, we observed 304 courting both females in 10 experiments, courting one female in five experiments, and no courting in two experiments. Males courted first "guttifera wing" females in 28 305 experiments and "melanogaster wing" females in 20 experiments, which were not 306 significantly different (exact binomial test, p=0.3123). Courtship latencies were not 307 308 significantly different between courtships with "guttifera wing" females (Latency 1, 16.1 \pm 23.9 min; Latency 2, 41.7 \pm 41.6 min) and courtships with "melanogaster wing" 309 females (Latency 1, 14.7 \pm 17.6 min; Latency 2, 35.3 \pm 27.6 min) (*t*-test, *p*=0.7698 and 310 p=0.5498). Courtship durations were not significantly different between courtships with 311 312 "guttifera wing" females (Duration 1, 1.27 ± 3.51 min; Duration 2, 1.39 ± 1.86 min) and 313 courtships with "melanogaster wings" females (Duration 1, 0.61 ± 0.76 min; Duration 2, 314 1.30 ± 0.87 min) (*t*-test, *p*=0.2351 and *p*=0.8428).

For female choice experiments, we conducted 49 experiments to examine the preference for the polka-dotted *D. guttifera* wings (Figure 3b). Females mated with *"guttifera* wing" males 22 times and *"melanogaster* wing" males 12 times. Females mated with *"guttifera* wing" males significantly more frequently than with *"melanogaster* wing" males (GLM, estimate=0.94335, p=0.0361; chi-squared test, p=0.03382). Other explanatory variables were not statistically significant (GLM, p>0.05).

We observed courtships to estimate males' eagerness to mate in these 49 321 experiments. Males courted females in 48 experiments. Among 34 experiments with 322 323 successful matings, both males courted in 30 experiments and one male courted in four 324 experiments. Among 15 experiments without matings, both males courted in 12 experiments, one male courted in two experiments, and no courting was observed in one 325 326 experiment. Females were courted first by "guttifera wing" males in 23 experiments and "melanogaster wing" males in 25 experiments, which were not significantly different 327 328 (exact binomial test, p=0.8854). The courtship latency (Latency 1) of "guttifera wing" 329 males (12.8 \pm 24.0 min) and that of "melanogaster wing" males (11.7 \pm 25.9 min) were not significantly different (t-test, p=0.8351). Therefore, there was no evidence of the 330 difference of males' eagerness to mate. In summary, females were found to prefer 331 "guttifera wing" males to "melanogaster wing" males. In these experiments, however, 332 333 we could not distinguish whether females preferred conspecific wings or the polka-dotted pattern. For this reason, we performed additional experiments using conspecific but 334 335 differently patterned wings.

336

337 Preference for polka-dotted pattern

338 In these additional experiments, we conducted 30 experiments to examine females' preference for different pigmentation patterns (Figure 4a). Females mated with 339 340 "guttifera wing" males eight times and "incomplete pattern" males 10 times, and these rates were not significantly different (GLM, estimate=-0.328292, p=0.572; chi-squared 341 342 test with Yates correction p=0.7782; with Bonferroni correction, p=1). Other explanatory 343 variables were also not statistically significant (GLM, p>0.05). Also, we observed 344 courtships to estimate males' eagerness to mate in these 30 experiments. Males courted females in 29 experiments. Among 18 experiments with successful matings, both males 345 346 courted in 13 experiments and one male courted in five experiments. Among 12 347 experiments without matings, both males courted in eight experiments, one male courted in three experiments, and no courting was observed in one experiment. Females were 348 349 courted first by "guttifera wing" males in 15 experiments and "incomplete pattern" males in 15 experiments (exact binomial test, p=1). The courtship latency (Latency 1) of 350 351 "guttifera wing" males (9.54 \pm 11.0 min) and that of "incomplete pattern" males (8.49 \pm 12.5 min) were not significantly different (t-test, p=0.7525). In these results, we found no 352 evidence of a difference in males' eagerness to mate. 353

354

We then considered the possibility that females might prefer darkness of male

wings, not a particular pattern. We conducted 38 experiments to examine preference for 355 darkness of entire wings (Figure 4b). Females mated with "wild type" males 11 times and 356 "black wing" males 13 times, rates that were not significantly different (GLM, estimate=-357 0.2440, p=0.6219; chi-squared test with Yates correction, p=0.8051; with Bonferroni 358 359 correction, p=1). Other explanatory variables were also not statistically significant (GLM, p>0.05). Males courted females in all 38 experiments. Among 24 experiments with 360 successful matings, both males courted in all 24 experiments. Among 14 experiments 361 362 without matings, both males courted in 11 experiments and one male courted in three experiments. Females were courted first by "wild type" males in 17 experiments and 363 364 "black wing" males in 21 experiments, which were not significantly different (exact binomial test, p=0.6271). The courtship latency (Latency 1) of "wild type" males (19.7 ± 365 29.9 min) and that of "black wing" males $(9.33 \pm 13.9 \text{ min})$ were not significantly 366 different (*t*-test, p=0.06082). In these results, we found no evidence of a difference in 367 males' eagerness to mate. 368

369 Finally, we conducted 41 experiments to examine whether females preferentially mated with "melanogaster wing" males or "melanogaster black wing" males (Figure 4c). 370 Females mated with "melanogaster wing" males 12 times and "melanogaster black wing" 371 males 12 times, rates that were not different (GLM, estimate=0.008613, p=0.9861; chi-372 373 squared test, p=1; Bonferroni correction, p=1). Other explanatory variables were also not 374 statistically significant (GLM, p>0.05). Males courted females in all 41 experiments. Among 24 experiments with successful matings, both males courted in 22 experiments 375 376 and one male courted in two experiments. Among 17 experiments without matings, both 377 males courted in 13 experiments and one male courted in four experiments. Females were courted first by "melanogaster wing" males in 18 experiments and "melanogaster black 378 379 wing" males in 23 experiments, which were not significantly different (exact binomial 380 test, p=0.5327). The courtship latency (Latency 1) of "melanogaster wing" males (15.5 \pm 27.8 min) and that of "melanogaster black wing" males (16.1 \pm 29.3 min) were not 381 significantly different (*t*-test, p=0.9283). In these results, we found no evidence of the 382 difference in males' eagerness to mate. 383

384

Taking these results altogether, we found that females prefer *D. guttifera* wings 385 to D. melanogaster wings, but have no preference for pigmentation pattern or wing colouration. The difference between the preference for D. guttifera wings and D. 386 melanogaster wings seems to be caused by some other wing characteristics, such as wing 387 shape or smell (See Discussion). 388

389

In two sets of male mate choice experiments, the rates of mating were 0.686 391 (35/51 experiments in Figure 2a) and 0.660 (33/50 experiments in Figure 3a). The rates 392 of courtship were 0.922 (47/51 experiments in Figure 2a) and 0.960 (48/50 experiments 393 394 in Figure 3a). These results show that most males courted females, suggesting that 395 matings could be largely affected by eagerness of females, rather than that of males. As 396 most previous studies have indicated individual differences of mate preference (Bell, Hankison, & Laskowshi, 2009), there could be individual differences of eagerness in 397 398 females of *D. guttifera*.

399 In five sets of female choice experiments, the rates of mating were 0.824 (42/51 400 experiments in Figure 2b), 0.694 (34/49 experiments in Figure 3b), 0.600 (18/30 401 experiments in Figure 4a), 0.632 (24/38 experiments in Figure 4b) and 0.585 (24/41 experiments in Figure 4c). The rates of courtship were 1.000 (51/51 experiments in Figure 402 403 2b), 0.980 (48/49 experiments in Figure 3b), 0.967 (29/30 experiments in Figure 4a), 404 1.000 (38/38 experiments in Figure 4b) and 1.000 (41/41 experiments in Figure 4c). 405 Again, these results show that most males courted females, suggesting that the mating could be affected by eagerness in females, rather than that in males. 406

407 Except in the experiments to examine preference for wings (Figure 2b), the rates
408 of mating were low. This could be explained by effect(s) of artificial treatments – painting
409 wings with a black ink marker and replacing wings using a cyanoacrylate adhesive.

410 Lastly, we examined whether females mated with males that courted first, by 411 integrating five female choice experiments. Females mated with males that courted first 412 in 99 experiments. Females mated with males that did not court first in 44 experiments. 413 In summary, females mated with males that courted first significantly more than with the 414 other males (exact binomial test, p=4.887e-06).

415

416 **Discussion**

417 Preference for wings

We found that females mated more with "wild type" males than "no wing" males 418 419 in female mate choice experiments. In our observations, females tended to mate with the males that courted first. However, because we found no evidence that "wild type" males 420 421 courted first, we conclude that mating more with "wild type" males was due to female 422 preference for wings. Because our observations and previous studies showed that D. 423 guttifera males do not perform courtship with obvious wing motion (Spieth, 1952; Grossfield, 1977; Wen & Li, 2011), mate choice by females does not seem to require male 424 425 wing motion. Instead, females may recognize male wing smell or shape for species 426 identification and use these features to choose mates.

Similarly, males might use the presence of wings to assess the quality of females. 427 Male mate choice has been reported in multiple *Drosophila* species (Bonduriansky, 2001; 428 Byrne & Rice, 2006). Another potential explanation for female wing function is related 429 to females' behaviour during a courtship sequence. Grossfield (1977) observed that 430 431 receptive females spread their wings, but non-receptive females repel males by 432 decamping and kicking males. At the moment of mate acceptance, females spread their wings widely, and this motion is likely to work as a visual signal of acceptance (Spieth, 433 434 1952; Grossfield, 1966). In the male mate choice experiments, we found that 1) males mated more with "wild type" females than "no wing" females, 2) courtship latencies were 435 436 not significantly different, and 3) courtship durations of males towards "no wing" females 437 were longer than those towards "wild type" females. Based on these results, we consider it possible that wing motion of females could have a function to show acceptance and 438 increase initiation of copulation by males, and it could be a potential reason why females 439 without wings had a lower rate of mating than intact females. Also, we found that males 440 441 courted first "wild type" females in 29 experiments and "no wing" females in 18 experiments. Although this difference was not statistically significant (exact binomial test, 442 443 p=0.1439), we cannot rule out the possibility that males could recognize female wing 444 shape or smell and use these features to choose mates.

Previous studies showed that wing damage can influence reproductive success, survival, mortality and flight performance in various insects, such as Odonata (Combes, Crall, & Mukherjee, 2010), Hymenoptera (Cartar, 1992) and Lepidoptera (Jantzen & Eisner, 2008). The present study found that both *D. guttifera* males and females mate more with intact wild-type adults than adults without wings. This suggests that wing damage could influence the preference for mates in *D. guttifera*.

451

452 Preference for conspecific wings

453 In male mate choice experiments, no significant difference was seen between the number of matings of males with "guttifera wing" females and those of males with 454 "melanogaster wing" females. But in female mate choice experiments, females were 455 456 found to mate more frequently with "guttifera wing" males than "melanogaster wing" 457 males. In our observations, females tended to mate with males that were the first to court. However, we found no evidence that "guttifera wing" males were the first to court. 458 Therefore, mating more with "guttifera wing" males would be due to female preference 459 for conspecific wings. Because female mate choice generally is stronger and more 460 common than male mate choice (Bonduriansky, 2001), these results would not be 461 462 surprising.

463

464 Preference for the polka-dotted pattern

In female mate choice experiments, females mated more with "*guttifera* wing" males than "*melanogaster* wing" males. This result suggested two possibilities—females have a preference for the polka-dotted pattern or a preference for other traits of conspecific wings. In further experiments, we did not detect females' preference for the details of the polka-dotted pattern or for darkness of the entire wings. Also, we found no evidence of a difference in males' eagerness to mate.

471 Taking these results together, we did not find evidence of a contribution of the 472 sexually monomorphic polka-dotted pattern on wings to mate choice of either of the sexes 473 in D. guttifera. To our knowledge, this is the first study to examine the contribution of sexually monomorphic ornaments in *Drosophila* to mate choice. Roy and Gleason (2019) 474 also did not detect a contribution of sexually dimorphic black spots on male wings to 475 476 mate choice in three Drosophila species. We cannot rule out, however, the possibility of 477 a weak preference for the polka-dotted pattern, which was not statistically significant in 478 this study, but is strong enough to be a selective pressure to fix the pattern in the 479 population. Also, although the preference was not found in our study in laboratory 480 conditions, we cannot exclude the possibility of the preference in a natural habitat. So far, the contribution of sexually monomorphic ornaments to mate choice is known in king 481 482 penguins (Pincemy et al., 2009; Nolan et al., 2010). In the future, the question of what conditions enable sexually monomorphic ornaments to contribute to mate choice should 483 484 be examined using species of various groups.

485 A contribution to mate choice was not detected in this study, and the function of 486 the polka-dotted pattern on wings of D. guttifera is still unclear. Considering the extremely sophisticated gene regulation that acts to form the polka-dotted pattern 487 488 (Fukutomi & Koshikawa, 2021; Fukutomi, Kondo, Toyoda, Shigenobu, & Koshikawa, 2021), it is difficult to think that the pattern evolved randomly as a neutral trait without 489 adaptive significance. Potential candidates of the function of the polka-dotted 490 pigmentation include thermoregulation, background matching, warning colouration and 491 species recognition. The ecological adaptations of D. guttifera and related species might 492 493 play an important role in the evolution of the pigmentation pattern of the present form. 494 Future research is needed to reveal what adaptive significance the polka-dotted pattern has in *D. guttifera*. 495

496

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- 503
- 504 Conflict of interest
- 505 The authors declare that they have no conflict of interest.

506 **References**

- Andersson, M. (1982). Female choice selects for extreme tail length in a widowbird.
 Nature, 299(5886), 818-820. https://doi.org/10.1038/299818a0
- Avent, T. D., Price, T. A. R., & Wedell, N. (2008). Age-based female preference in the
 fruit fly *Drosophila pseudoobscura*. *Animal Behaviour*, 75(4), 1413-1421.
 https://doi.org/10.1016/j.anbehav.2007.09.015
- Bell, A. M., Hankison, S. J., & Laskowski, K. L. (2009). The repeatability of behaviour:
 a meta-analysis. *Animal Behaviour*, 77(4), 771-783.
 https://doi.org/10.1016/j.anbehav.2008.12.022
- Bonduriansky, R. (2001). The evolution of male mate choice in insects: a synthesis of
 ideas and evidence. *Biological Reviews*, 76(3), 305-339.
 https://doi.org/10.1017/s1464793101005693
- Bonduriansky, R., & Chenoweth, S. F. (2009). Intralocus sexual conflict. *Trends in Ecology & Evolution*, 24(5), 280-288. https://doi.org/10.1016/j.tree.2008.12.005
- Byrne, P. G., & Rice, W. R. (2006). Evidence for adaptive male mate choice in the fruit
 fly *Drosophila melanogaster*. *Proceedings of the Royal Society B: Biological Sciences*, 273(1589), 917-922. https://doi.org/10.1098/rspb.2005.3372
- Campbell, I. (2007). Chi squared and Fisher–Irwin tests of two by two tables with
 small sample recommendations. *Statistics in Medicine*, 26(19), 3661-3675.
 https://doi.org/10.1002/sim.2832
- 526 Cartar, R. V. (1992). Morphological senescence and longevity: an experiment relating
 527 wing wear and life span in foraging wild bumble bees. *Journal of Animal Ecology*,
 528 61(1), 225-231. https://doi.org/10.2307/5525
- 529 Chialvo, C. H. S., White, B. E., Reed, L. K., & Dyer, K. A. (2019). A phylogenetic
 530 examination of host use evolution in the *quinaria* and *testacea* groups of *Drosophila*.
 531 *Molecular Phylogenetics and Evolution*, 130, 233-243.
 532 https://doi.org/10.1016/j.ympev.2018.10.027
- Colinet, H., & Renault, D. (2012). Metabolic effects of CO2 anaesthesia in *Drosophila melanogaster*. *Biology Letters*, 8(6), 1050-1054.
 https://doi.org/10.1098/rsbl.2012.0601
- Combes, S. A., Crall, J. D., & Mukherjee, S. (2010). Dynamics of animal movement in
 an ecological context: dragonfly wing damage reduces flight performance and
 predation success. *Biology Letters*, 6(3), 426-429.
 https://doi.org/10.1098/rsbl.2009.0915
- 540 Dion, W. A., Shittu, M. O., Steenwinkel, T. E., Raja, K. K., Kokate, P. P., & Werner, T.
 541 (2020). The modular expression patterns of three pigmentation genes prefigure

- unique abdominal morphologies seen among three *Drosophila* species. *Gene Expression Patterns*, 119132. https://doi.org/10.1016/j.gep.2020.119132
- 544 Dufour, H. D., Koshikawa, S., & Finet, C. (2020) Temporal flexibility of gene regulatory
 545 network underlies a novel wing pattern in flies. *Proceedings of the National*546 *Academy of Sciences of the United States of America*, 117(21), 11589-11596.
 547 https://doi.org/10.1073/pnas.2002092117
- Edwards, K. A., Doescher, L. T., Kaneshiro, K. Y., & Yamamoto, D. (2007). A database
 of wing diversity in the Hawaiian *Drosophila*. *PLoS One*, 2(5), e487.
 https://doi.org/10.1371/journal.pone.0000487
- Fisher, R. A. (1915). The evolution of sexual preference. *The Eugenics Review*, 7(3), 184192.
- Fukutomi, Y., & Koshikawa, S. (2021). Mechanism of color pattern formation in insects.
 In H. Hashimoto, M. Goda, R. Futahashi, R. Kelsh & T. Akiyama (Eds.), *Pigments, pigment cells, pigment patterns*. Singapore: Springer. (in press)
- Fukutomi, Y., Kondo, S., Toyoda, A., Shigenobu, S., & Koshikawa, S. (2021).
 Transcriptome analysis reveals *wingless* regulates neural development and signaling
 genes in the region of wing pigmentation of a polka-dotted fruit fly. *FEBS Journal*,
 288(1), 99-110. https://doi.org/10.1111/febs.15338
- Fukutomi, Y., Matsumoto, K., Agata, K., Funayama, N., & Koshikawa, S. (2017). Pupal
 development and pigmentation process of a polka-dotted fruit fly, *Drosophila guttifera* (Insecta, Diptera). *Development Genes and Evolution*, 227(3), 171-180.
 https://doi.org/10.1007/s00427-017-0578-3
- Fukutomi, Y., Matsumoto, K., Funayama, N., & Koshikawa, S. (2018). Methods for
 staging pupal periods and measurement of wing pigmentation of *Drosophila guttifera*. JoVE (Journal of Visualized Experiments), (131), e56935.
 https://doi.org/10.3791/56935
- Fuyama, Y. (1979). A visual stimulus in the courtship of *Drosophila suzukii*. *Experientia*,
 35(10), 1327-1328. https://doi.org/10.1007/BF01963987
- Grossfield, J. (1966). The influence of light on the mating behaviour of *Drosophila*. In M.
 R. Wheeler (Ed.), Studies in Genetics III (pp. 147-176). Austin, Texas: The
 University of Texas Publication.
- Grossfield, J. (1977). *Drosophila* courtship: decapitated quinaria group females. *Journal of the New York Entomological Society*, 85(3), 119-126.
- Hegde, S. N., Chethan, B. K., & Krishna, M. S. (2005). Mating success of males with and
 without wing patch in *Drosophila biarmipes*. *Indian Journal of Experimental Biology*, 43(10), 902-909.

- Izumitani, H. F., Kusaka, Y., Koshikawa, S., Toda, M. J., & Katoh, T. (2016).
 Phylogeography of the subgenus *Drosophila* (Diptera: Drosophilidae): evolutionary
 history of faunal divergence between the Old and the New Worlds. *PLoS One*, *11*(7),
 e0160051. https://doi.org/10.1371/journal.pone.0160051
- Jantzen, B., & Eisner, T. (2008). Hindwings are unnecessary for flight but essential for
 execution of normal evasive flight in Lepidoptera. *Proceedings of the National Academy of Sciences of the United States of America*, 105(43), 16636-16640.
 https://doi.org/10.1073/pnas.0807223105
- Kopp, A., & True, J. R. (2002). Evolution of male sexual characters in the oriental *Drosophila melanogaster* species group. *Evolution & Development*, 4(4), 278-291.
 https://doi.org/10.1046/j.1525-142x.2002.02017.x
- Koshikawa, S. (2020). Evolution of wing pigmentation in *Drosophila*: Diversity,
 physiological regulation, and *cis*-regulatory evolution. *Development, Growth & Differentiation*, 62(5), 269-278. https://doi.org/10.1111/dgd.12661
- Koshikawa, S., Fukutomi, Y., & Matsumoto, K. (2017). *Drosophila guttifera* as a model
 system for unraveling color pattern formation. In T. Sekimura & H. F. Nijhout (Eds.), *Diversity and Evolution of Butterfly Wing Patterns* (pp. 287-301). Singapore:
 Springer.
- Koshikawa, S., Giorgianni, M. W., Vaccaro, K., Kassner, V. A., Yoder, J. H., Werner, T.,
 & Carroll, S. B. (2015). Gain of cis-regulatory activities underlies novel domains of
 wingless gene expression in *Drosophila*. *Proceedings of the National Academy of Sciences of the United States of America*, 112(24), 7524-7529.
 https://doi.org/10.1073/pnas.1509022112
- Kraaijeveld, K., Kraaijeveld-Smit, F. J., & Komdeur, J. (2007). The evolution of mutual
 ornamentation. *Animal Behaviour*, 74(4), 657-677.
 https://doi.org/10.1016/j.anbehav.2006.12.027
- Lande, R. (1980). Sexual dimorphism, sexual selection, and adaptation in polygenic
 characters. *Evolution*, 34(2), 292-305. https://doi.org/10.1111/j.15585646.1980.tb04817.x
- Miller, M. E., Marshall, S. A., & Grimaldi, D. A. (2017) A review of the species of *Drosophila* (Diptera: Drosophilidae) and genera of Drosophilidae of Northeastern
 North America. *Canadian Journal of Arthropod Identification*, 31. https://
 doi.org/10.3752/cjai.2017.31
- Nolan, P. M., Dobson, F. S., Nicolaus, M., Karels, T. J., McGraw, K. J., & Jouventin, P.
 (2010). Mutual mate choice for colorful traits in king penguins. *Ethology*, *116*(7),
 635-644. https://doi.org/10.1111/j.1439-0310.2010.01775.x

- Parkash, R., Lambhod, C., & Singh, D. (2013). Ecological significance of wing spot
 dimorphism in *Drosophila biarmipes* (Diptera, Drosophilidae). *Acta Entomologica Sinica*, 56(11), 1267-1274.
- Patterson, J. T. (1943). The Drosophilidae of the Southwest. Austin, Texas: The
 University of Texas Publication.
- Petrie, M., & Halliday, T. (1994). Experimental and natural changes in the peacock's
 (*Pavo cristatus*) train can affect mating success. *Behavioral Ecology and Sociobiology*, *35*(3), 213-217. https://doi.org/10.1007/BF00167962
- Petrie, M., Halliday, T., & Sanders, C. (1991). Peahens prefer peacocks with elaborate
 trains. *Animal Behaviour*, 41(2), 323-331. https://doi.org/10.1016/S00033472(05)80484-1
- Pincemy, G., Dobson, F. S., & Jouventin, P. (2009). Experiments on colour ornaments
 and mate choice in king penguins. *Animal Behaviour*, 78(5), 1247-1253.
 https://doi.org/10.1016/j.anbehav.2009.07.041
- Prud'Homme, B., Gompel, N., Rokas, A., Kassner, V. A., Williams, T. M., Yeh, S. D.,
 True, J. R., & Carroll, S. B. (2006). Repeated morphological evolution through cisregulatory changes in a pleiotropic gene. *Nature*, 440(7087), 1050-1053.
 https://doi.org/10.1038/nature04597
- R Core Team (2018). R: A language and environment for statistical computing. R
 Foundation for Statistical Computing, Vienna, Austria. Retrieved from https://www.R-project.org/
- Roy, P. R., & Gleason, J. M. (2019). Assessing the use of wing ornamentation and visual
 display in female choice sexual selection. *Behavioural Processes*, *158*, 89-96.
 https://doi.org/10.1016/j.beproc.2018.10.010
- Ryan, M. J. (1985). The túngara frog: a study in sexual selection and communication.
 Chicago: University of Chicago Press.
- Shittu, M., Steenwinkel, T., Koshikawa, S., Werner, T. (2020). The making of transgenic *Drosophila guttifera*. *Methods and Protocols*, 3(2), 31.
 https://doi.org/110.3390/mps3020031
- Singh, B. N., & Chatterjee, S. (1987). Greater mating success of *Drosophila biarmipes*males possessing an apical dark black wing patch. *Ethology*, 75(1), 81-83.
 https://doi.org/10.1111/j.1439-0310.1987.tb00643.x
- 646 Spieth, H. T. (1952). Mating behaviour with in genus *Drosophila* (Diptera). New York:
 647 Bulletin of American Museum of Natural History.
- Spieth, H. T. (1966). Courtship behavior of endemic Hawaiian *Drosophila*. Austin, Texas:
 The University of Texas Publication.

- Theis, A., Salzburger, W., & Egger, B. (2012). The function of anal fin egg-spots in the
 cichlid fish *Astatotilapia burtoni*. *PLoS One*, 7(1), e29878.
 https://doi.org/10.1371/journal.pone.0029878
- Verspoor, R. L., Cuss, M., & Price, T. A. R. (2015). Age-based mate choice in the
 monandrous fruit fly *Drosophila subobscura*. *Animal Behaviour*, *102*, 199-207.
 https://doi.org/10.1016/j.anbehav.2015.01.024
- Wen, S. Y., & Li, Y. F. (2011). An evolutionary view on courtship behavior of *Drosophila*:
 from a comparative approach. *Low Temperature Science*, *69*, 87-100.
- Werner, T., Koshikawa, S., Williams, T. M., & Carroll, S. B. (2010). Generation of a novel
 wing colour pattern by the Wingless morphogen. *Nature*, 464(7292), 1143-1148.
 https://doi.org/10.1038/nature08896
- Werner, T., Steenwinkel, T., & Jaenike, J. (2020). The encyclopedia of North American
 Drosophilids Volume 1: Drosophilids of the Midwest and Northeast. Houghton, MI:
 Michigan Technological University.
- Wright, S. (1992). Adjusted p-values for simultaneous inference. *Biometrics*, 48(4), 10051013. https://doi.org/10.2307/2532694
- Zahavi, A. (1975). Mate selection—a selection for a handicap. *Journal of Theoretical Biology*, 53(1), 205-214. https://doi.org/10.1016/0022-5193(75)90111-3

668 Figure legends

669 **Figure 1.**

Flies used in mating experiments. (a) A "wild type" adult. (b) A "no wing" adult, whose wings were removed. (c) A "black wing" adult, whose wings were painted black. (d) A "*guttifera* wing" adult, whose wings were replaced with *D. guttifera* wings. (e) A "*melanogaster* wing" adult, whose wings were replaced with *D. melanogaster* wings. (f) An "incomplete pattern" adult, whose wings were replaced with wings without pigmentation spots around the campaniform sensilla. (g) A "*melanogaster* black wing" adult, whose wings were replaced with *D. melanogaster* black wing"

677

678 **Figure 2.**

Experiments to test the preference for wings. (a) Male preference for females with/without wings. The heights of the bars (sum of gray and black areas) indicate the numbers of the first mating. The black areas represent the numbers of the second mating with the other female. The gray areas represent the number of males which mated only once. White areas represent no mates. (b) Female preference for males with/without wings.

685

686 **Figure 3.**

Experiments to test the preference for conspecific wings. (a) Male preference for females with "*guttifera wing*" or "*melanogaster wing*". The heights of the bars (sum of gray and black areas) indicate the numbers of the first mating. The black areas represent the numbers of the second mating with the other female. The gray areas represent the number of males which mated only once. b) Female preference for males with "*guttifera wing*" or "*melanogaster wing*".

693

694 **Figure 4.**

Experiments to test female preferences for different types of wings. (a) Preference for males with "*guttifera wing*" or "incomplete pattern". (b) Preference for males of "wild type" or with "black wing". (c) Preference for males with "*melanogaster* wing" or "*melanogaster* black wing".

699

700 **Table 1.**

701 General design of mate choice experiments. One adult and two adults of the other sex

702 (Type A and B) were used in seven mate choice experiments. Each item was observed for

two types of adults.









	Туре А	Туре В	Observed items
Male mate choice experiments	wild type	no wing	 Which type of a female mated
			 Which type of a female was courted first
	guttifera wing	<i>melanogaster</i> wing	 Courtship latency and duration
Female mate choice experiments	wild type	no wing	
	guttifera wing	melanogaster wing	◆ Which type of a male mated
	guttifera wing	incomplete pattern	 Which type of a male courted first
	wild type	black wing	 Courtship latency
	melanogaster wing	<i>melanogaster</i> black wing	