Effects of a sex ratio gradient on female mate-copying and choosiness in *Drosophila melanogaster*

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Abstract

In many sexually reproducing species, individuals can gather information about potential mates by observing their mating success. This behavioral pattern, that we call mate-copying, was reported in the fruit fly *Drosophila melanogaster* where females choosing between 2 males of contrasting phenotypes can build a preference for males of the phenotype they previously saw being chosen by a demonstrator female. As sex ratio is known to affect mate choice, our goal was to test whether mate-copying is also affected by encountered sex ratios. Thus, we created a gradient of sex ratio during demonstrations of mate-copying experiments by changing the number of females observing from a central arena 6 simultaneous demonstrations unfolding in 6 peripheral compartments of a hexagonal device. We also tested whether the sex ratio experienced by females during demonstrations affected their choosiness (male courtship duration and double courtship rate) in subsequent mate-choice tests. Experimental male:female sex ratio during demonstrations did not affect mate-copying indices, but positively affected the proportion of both males courting the female during mate-choice tests, as well as male courtship duration, the latter potentially explaining the former relationship. As expected, the sex ratio affected female choosiness positively, and *Drosophila* females seem to have evolved a mate-copying ability independently of sex ratio, and a capacity to adapt their choosiness to male availability. This suggests that, as in many animal species, individuals, especially females, can adapt their mate choice depending on the current sex ratio.

Key words: competition, *Drosophila melanogaster*, experimental protocol, mate-copying, social learning, sex ratio.

Choosing a mate is a major fitness-affecting decision in any sexually reproducing organism. In species where females invest more than males in the production of a single offspring, females are selected to become the choosy sex (Johnstone et al. 1996; Trivers 1972). Furthermore, in males of such species, natural and sexual selection shape traits that are related to male quality, which in turn can be used by females to assess male quality because these traits reveal that they are better fathers providing better or more resources to the female or the offspring (Candolin 2003). Male mating success is thus affected by various parameters revealing their intrinsic quality.
(Weatherhead and Boag 1995), such as their size, ornaments, bright colors, or songs as this was documented in many animal taxa including vertebrates and invertebrates (Partridge and Farquhar 1983; Scarcy 1992; Madsen et al. 1993; Andersson 1994; Aspi and Hoikkala 1995; Bateman et al. 2001; Dreher and Prohl 2014; reviewed in Danchin and Cézilly 2008).

Alternatively but not exclusively, females can get further information about potential mates by observing their mating success (Danchin et al. 2004). We call such observational learning mate-copying. It leads females to either mate preferentially with the specific male they saw being chosen by another female (individual-based copying, Pruett-Jones 1992; Bowers et al. 2012), or with a male showing similar characteristics as the male they saw being chosen by another female (trait-based copying, Bowers et al. 2012). Trait-based mate-copying is particularly interesting because learning to prefer males of a given phenotype rather than a specific male (Witte et al. 2013) may potentially lead to the establishment of persistent local traditions in mate choice (Danchin et al., submitted for publication), which in turn may strongly affect sexual selection differentially across populations, thus setting the stage for speciation.

Mate-copying has been reported in many social and non-social species, including fish (Dugatkin and Godin 1993), birds (White and Galef 1999), humans (Waynforth 2007), and other mammals (Galef et al. 2008), as well as in 1 insect, Drosophila melanogaster (Mery et al. 2009; Loyau et al. 2012; Dagaef et al. 2016; Germain et al. 2016; Danchin et al., submitted for publication; Nöbel et al., submitted for publication). In particular, D. melanogaster females can perform trait-based mate-copying after watching only a single live demonstration of 1 female copulating with a male of a given phenotype and 1 male of another phenotype being rejected (Dagaef et al. 2016; Danchin et al., submitted for publication; Nöbel et al., submitted for publication). More generally, it is now accepted that many animal species from a vast array of taxa can learn from others (i.e., socially learn), particularly in the context of mate choice (Avital and Jablonka 2000; Danchin et al. 2004; Galef and Laland 2005).

Sex ratio is known to affect male–male competition in a mate-choice context in various species (Lawrence 1986; Jirokulk 1999; Weir et al. 2011), probably because it affects the availability of potential partners. For instance, in D. melanogaster, male sexual behavior is influenced by the number of rivals (Bretman et al. 2009), and male sperm depletion starts after just one copulation, so that the number of emerging offspring is divided by more than 3 after 4 consecutive copulations (Demerec and Kaufmann 1941; Lefèvre and Jonsson 1962; Loyau et al. 2012). Thus, females are also expected to adapt their sexual behavior to the sex ratio (i.e., to the number of competitor males), for instance by accepting mates more readily when the male-to-female ratio is low, that is, when the number of potential female competitors is high.

Females can use 2 different sources of information to select a male partner: females might rely on their personal assessment of males’ courtship during the mate-choice test, or only rely on the social information provided by demonstrator females during the demonstration. The latter option is probably more economical in a context of high level of female competition, as mate choice is costly to females (reviewed in Reynolds and Gross 1990; Andersson 1994; Vakirtzis 2011). For instance, females invest time and energy to assess male quality, and male courtship may sometimes be harmful to them (Andersson 1994). Moreover, the risk of losing potential mates to competitors increases with the time spent in assessing males. Under high competition scenario, one could expect females to use the more easily gathered social information only, thus minimizing risks of losing potential mates to competitors. In contrast, under low female competition (i.e., in male-biased sex ratios), this risk is minor, allowing them to spend time in male assessment.

So far, no study has investigated the effect of sex ratio on mate-copying or social learning in general. As in the more general context of mate choice, we can expect females to show contrasting choosiness when being within groups of varying sex ratios (Berglund 1994; Jirokulk 1999; Passos et al. 2014). For example, being under strong competition to access males, females learning within a mixed-sex group mainly composed of females might be much more prone to accept the very first male that courts them, and thus, ignore social information. Contrarily, females learning within a mixed-sex group essentially composed of males can be expected to be much choosier and to take the time to gather more information about the various potential males before selecting one of them.

Here, we studied mate-copying along a gradient of group sex ratio during demonstrations. All mate-copying designs in Drosophila involve a demonstration phase (or simply demonstration) followed by a mate-choice test. We manipulated the population sex ratio during demonstrations by varying the number of observer females (and thus group size) in the central arena of a hexagonal experimental device (see “Materials and Methods” section). By assuming that observer females can assess and remember the group size and sex ratio they experienced during the demonstration, we predicted that along our gradient of increasing sex ratios during demonstrations, observer females would become more and more choosy during the subsequent mate-choice test, leading them to 1) accept copulation slower (i.e., a longer delay between first courtship and copulation initiation), and thus 2) increasing the rate of replicates in which both males courted the female before copulation initiation (“double courtship rate”), a parameter with methodological implications for future mate-copying experiments.

Concerning mate-copying, we could predict at least 3 possible outcomes, according to how we envisage the group size and sex ratio effects. The group size effect can be either positive or negative: first, females might learn better in a group than alone, thus predicting a positive mate-copying to group size relationship in a form of “social learning.” Inversely, in large groups of females (i.e., at female-biased sex ratios) group members may start to disturb and stress each other, thus hampering proper learning and leading to a negative mate-copying to group size relationship in our system. On the other hand, sex ratio could have a negative effect: we could expect that under female-biased sex ratios during demonstration (i.e., in large groups) observer females will adopt the less costly strategy and thus favors social information use over the time-consuming assessment of both males, thus leading to a negative mate-copying to sex-ratio relationship. These potential and contradictory effects of group size and sex ratio on mate-copying, being non-exclusive, might also cancel out each other, leading to no detectable pattern in mate-copying with group size and/or sex ratio. They may also lead to an optimal group size at which mate-copying is more efficient. The mate-copying to group size and sex ratio relationship should thus depend on the relative importance of these various potential effects, so that we did not have clear predictions.

Materials and Methods

Fly maintenance

Wild-type Canton-S flies were raised in 30 ml vials containing 10 ml of a standard corn flour—agar—yeast medium. They were maintained at 25 ± 1°C, 59 ± 5% humidity in a 12 h/12 h light/dark cycle.
Virgin flies were collected daily within 7 h after emergence, sexed without anesthesia, and kept in same-sex groups of 7 in a vial with medium. Experiments were conducted on 3–5 days old flies and fly manipulation was performed by gentle aspiration, using a glass pipette, tubing, and gauze.

**Experimental protocol**

All experiments were conducted under similar conditions as fly maintenance. Air pressure at the airport Toulouse-Blagnac weather station was in the range of 1,004–1,034 hPa and was previously shown to constitute a highly reliable proxy of atmospheric pressure in the experimental room (Dagaeff et al. 2016). Contrasting male phenotypes were created by randomly dusting them with green or pink powders (Mery et al. 2009) 20–30 min prior to use them as demonstrators or potential mates, so that they could clean the excess of dust. Demonstrations were run according to the “speed learning” design (Dagaeff et al. 2016) within a hexagon device (Figure 1A) composed of 1 central arena devoted to female observers (2.7 cm × 1.5 cm, volume = 8.6 cm³) and 6 peripheral compartments devoted to demonstrators (1.5 cm × 1.5 cm × 1.5 cm each) separated from the central arena by a glass partition (0.8 mm thick). The 4 experimental groups differed in the number of observer females: 1, 6, 12, and 24 observer females in the central arena, being able to witness 6 simultaneous demonstrations each ongoing in 1 of the 6 peripheral compartments. Each demonstration involved 1 female apparently choosing the same male color as the other demonstrator females and rejecting the male of the other color. Thus, with a total of 6 demonstrator females and 12 demonstrator males, with 1, 6, 12, and 24 observer females in the central arena, the sex ratio within a hexagon ranged from 1.7 to 1, 0.67, and 0.4 males per female, respectively. Mate-copying tests were run using a device made of a double plastic tube (0.8 cm × 3 cm each) separated by a microscopy cover slide (1.6 cm × 1.6 cm, Figure 1B).

Before the beginning of the demonstration, observer females were placed in the central arena. Demonstrator females were placed individually in small plastic tubes with 2 males of the color chosen for the demonstration of that hexagon. As soon as copulation started, the couple and a male of the opposite color were transferred carefully into a peripheral compartment of the hexagon in which the copulation continued. Demonstrations started when the first demonstrator trio was transferred, and ended when all 6 couples were broken, or as soon as new courtship occurred in 1 peripheral compartment after the end of copulation, despite the fact that some of the other copulations might still be ongoing. Thus, demonstration length varied from 16 to 24 min. Then, all observer females were removed by gentle aspiration and placed all together in a food vial until the mate-choice test. For the condition with 24 observer flies, only 12 randomly chosen observer females were kept together in a vial for the test. Results were obtained from 57, 16, 13, and 14 demonstration blocks (i.e., hexagons) for conditions with 1, 6, 12, and 24 observer flies, respectively.

The mate-choice tests started 45–60 min after the end of the demonstration (for practical reasons, half of the flies were tested after 45 min and the other half after 60 min). Each observer fly was placed in one side of a double plastic tube device, and a pair of males, one of each color, in the other side. The males used in the test phase came from a different vial than those used in the demonstration, and were powdered 20–30 min before the beginning of the test. After 2 min resting, the partition was removed, thus beginning the test. The first wing vibration of a male was recorded as courtship initiation, as well as, the color of the courting male, the time of the beginning of copulation, and the color of the chosen male. In trials in which both males courted the female before she chose, the median time of double courtship was 68 s, and the minimum time of double courtship was 1 s.

After the end of the experiment, flies were transferred into a vial and euthanized in a freezer. As in previous studies (Dagaeff et al. 2016; Danchin et al., submitted for publication; Nobel et al., submitted for publication) mate-choice tests were successful if they led to a copulation and if both males courted the observer female before the initiation of the copulation, as this was the only situation when observer females were visibly in a situation of choice.

**Mate-copying index**

Replicates in which the observer female copulated with the male of the phenotype preferred during the demonstration (copied) were attributed a mate-copying score of 1, and 0 in the opposite case. The mate-copying index (MCI) was calculated as the mean mate-copying score for each treatment, which reveals female preference in the corresponding experimental group. MCIs significantly higher than 0.5 (random choice) reveal mate-copying.
GLMM, Wald

demonstrator female, and delay between demonstration and test:

tial confounding parameters in univariate tests and found no signifi-

cracy (package lme4, Bates et al. 2015). We analyzed all successful

tions, that is, replicates in which both males courted the female

designs, delay between demonstration and test) were found non-significant in univariate tests (GLMM, Wald \( \chi^2 \) test, \( N = 441 \), \( \chi^2 = 1.299, 3.390, \) and 1.738, \( P = 0.255, 0.640, \) and 0.187, respectively). Thus, the starting model included sex ratio, normalized air pressure, time when the first courtship began, and first-courtimg male as well as their interactions as fixed effects.

The log-transformed courtship duration was analyzed in a LMM with logistic regression. We analyzed all trials with detailed times of courtship and copulation initiation (432 trials out of 455). Log-transformation (natural log) was used to achieve a Gaussian distribution of that variable. All potential confounding parameters (time of the test, test chamber ID in the set of 6 tube designs, delay between demonstration and test) were non-significant in univariate tests (LMM, \( N = 432 \), \( F = 1.039, 0.997, \) and 0.228, \( P = 0.309, 0.419, \) and 0.633, respectively). The starting model thus included sex ratio, log-transformed time of first courtship initiation, and first-courtimg male as fixed effects.

**Results**

**Mate-copying along a gradient of sex ratio**

We analyzed mate-copying indices in a GLMM with binary logistic regression. The starting model included the sex ratio as a continuous variable, first-courtimg male, and normalized air pressure, as well as air pressure changes within 6 h before the experiment as fixed effects. None of the interactions were significant (\( P > 0.14 \) in all cases). Sex ratio had no effect on mate-copying (GLMM, Wald \( \chi^2 \) test, \( N = 172 \), \( \chi^2 = 0.052, P = 0.820 \); Figure 2), nor did air pressure changes within 6 h before the experiment (GLMM, Wald \( \chi^2 \) test, \( N = 172 \), \( \chi^2 = 0.25, P = 0.616 \)). The selected model included normalized air pressure (GLMM, Wald \( \chi^2 \) test, \( N = 172 \), \( \chi^2 = 3.81, P = 0.051 \), positive effect) and first courting male (GLMM, Wald \( \chi^2 \) test, \( N = 172 \), \( \chi^2 = 4.13, P = 0.042 \), negative effect). The MCI was higher when the first male courting in the test was the one of the color that was rejected by the demonstrator females.

Thus, we did not find any significant relationship of mate-copying with sex ratio (Figure 2). When analyzing mate-copying indices in each group using binomial tests, we found a significant departure from random choice for the groups with sex ratios of 1.7, 1, and 0.67 males per female, but not in the group with a sex ratio of 0.4 (Figure 2), although the trend was in the same direction with a tendency to choose the male of the color selected during the demonstration.

**Rate of double courtship along a sex ratio gradient**

We measured the rates of both males courting the female ("double courtship rate"), that is, the proportion of trials in which both males courted the female before she initiated mating with one of them. The starting binary logistic regression model included the sex ratio as a continuous variable, first-courtimg male, and normalized air...
pressure, as well as the log-transformed time of first courtship initiation as fixed effects. None of the interactions were significant ($P > 0.69$ in all cases). First-courting male had no effect on the double courtship rate (GLMM, Wald $\chi^2$ test, $N = 441$, $\chi^2 = 1.15$, $P = 0.283$). The selected model included sex ratio (GLMM, Wald $\chi^2$ test, $N = 441$, $\chi^2 = 6.333$, $P = 0.012$, positive effect, Figure 3), normalized air pressure, and log-transformed time of first courtship initiation (GLMM, Wald $\chi^2$ test, $N = 441$, $\chi^2 = 5.818$ and 8.978, $P = 0.016$ and 0.003, positive and negative effects, respectively). As expected, we found that the double courtship rate increased along the sex-ratio gradient (Figure 3). Females thus appeared to be able to assess the sex ratio during demonstrations, remember it, and adapt their behavior during the subsequent mate-choice test accordingly.

**Courtship duration along the sex ratio gradient**

Because copulation initiation in *D. melanogaster* is mainly under female control (Connolly and Cook 1973; Kimura et al. 2015), we tested whether the decrease in the double courtship rate was due to faster acceptance of copulation in groups with lower sex ratios during the demonstration. In a preliminary model analyzing number of males courting as a fixed effect depending on log-transformed courtship duration, we first found that these 2 variables were highly correlated (LMM, $N = 432$, $F = 124.3$, $P < 0.001$, positive effect), which supports our hypothesis: in trials in which both males courted, the latency between first courtship and copulation initiation was the highest. We then analyzed the delay between first courtship initiation and copulation initiation along the sex ratio gradient (Figure 4). In a LMM with logistic regression in which the response variable was the log-transformed courtship duration, including the sex ratio as a continuous variable, first-courting male, and the log of the time of first courtship initiation as fixed effects. None of the interactions were significant ($P > 0.32$ in all cases). The first-courting male index was not associated with courtship duration (LMM, $N = 432$, $F = 0.048$, $P = 0.827$), while sex ratio (Figure 4) and time of first courtship initiation were (LMM, $N = 432$, $F = 7.828$ and 14.19, $P = 0.005$ and <0.001, positive and negative effect, respectively). Thus, as expected, copulation occurred faster in low sex ratio conditions, suggesting that observer females were much quicker in accepting copulation in such situations.

**Discussion**

We investigated sex-ratio and group size effects during mate-choice demonstrations on the observer females’ tendency to copy the mate choice of demonstrator females for a specific male phenotype as well as their choosiness in the subsequent mate-choice tests. We expected that along a gradient of increasing sex ratio during demonstrations, observer females would accept copulation slower, thus increasing the double courtship rate. We had at least 3 contradictory predictions concerning the mate-copying to sex ratio and group size relationships, so that we had no specific expectation about the direction of the effect. The relationship of mate-copying to group size could be either (i) positive, because of some social facilitation in mate-copying or (ii) negative, as a result of decreasing disturbance by other females with decreasing group size, and the relationship of mate-copying to sex ratio could be (iii) negative, as a result of decreasing female competition for males when sex ratio increases. Finally, a combination of these effects could also produce an optimal group size and sex ratio at which social learning is maximized.

As expected, we found that the frequency of both males courting the observer female and the delay between first courtship initiation and copulation initiation increased along the increasing experimental sex-ratio gradient (and decreasing group size). This supports our hypothesis that female choosiness decreases when sex ratio gets more female biased. Interestingly, females copied on average the observed mate-choice decisions for a certain male phenotype regardless of the other treatments, suggesting that females did not learn better from the other treatments, suggesting that females did not learn better in a group. There seemed to be no “social social-learning” in this observational learning paradigm, which is in accordance with the fact that in a form of olfactory learning, flies tested in groups show increased memory retrieval compared with flies tested individually, but training condition (single vs. group) does not affect memory formation (Chabaud et al. 2009). Similarly, in large groups of observer females (i.e., with female-biased sex ratios), we did not detect any evidence for increased disturbance among flies that would have

![Figure 3. Female choosiness measured as double courtship rate. Rates are expressed as the number of trials in which both males courted the female on the total number of trials. OF, observer females. Error bars represent SEM. Sample sizes are provided above the X-axis.](image)

![Figure 4. Courtship duration measured as mean latency between time of first courtship initiation and copulation initiation along the sex-ratio gradient. OF, observer females. Error bars represent SEM. Sample sizes are provided above the X-axis.](image)
hampered proper learning, as mate-copying appeared relatively unchanged with the size of the group of observer females. Finally, we found no evidence that females learning in female biased groups favor social information use over the personal assessment of male’s quality. This might be explained by a lack of quality differences among presented males that led females to rely preferably on social information gathered during demonstrations. Another explanation is that several contradictory effects were ongoing simultaneously and cancelled out each other. Unfortunately, our experimental design did not allow us to disentangle group size from sex ratio effects as they co-varied in our design.

The ability to copy may be an adaptation to the naturally crowded conditions existing on rotten pieces of fruit to which females are attracted both as sources of food and egg laying site (Rodrigues et al. 2015; Keesey et al. 2016). In these aggregations, copulations are common (Danchin E, personal observation) and surrounding females have the opportunity to watch the mate choice of other females, thus setting the stage for mate-copying in natural situations.

As for methodological implications, our results suggest that it is possible to increase the number of observer females in the central arena of the hexagon during demonstrations at least up to 12 without affecting mate-copying efficiency. This could help in designing future mate-copying experiments, for instance by increasing the number of males in the peripheral compartments, and thus, increasing the sex ratio in order to increase double courtship rates without affecting mate-copying scores. This would allow us to extend the proportion of replicates in which females are visibly in a situation of choice during the mate-choice test.

All in all, our results suggest that mate-copying is quite robust to sex ratio conditions as well as group size differences in general. While this might be caused by contradictory effects of varying sex ratios and group sizes cancelling each other out, the high observed robustness in mate-copying may suggest a real importance of social information use in mate choice (see also Danchin et al. 2004; Galey and Laland 2005). In general, selection might have favored the evolution of mate-copying under varying environmental conditions. In particular, the Fisher runaway process (Fisher 1930) predicts that females should conform to the local preference because male descendants of females mating with the locally non-preferred males would inherit the non-preferred trait, thus counter-selecting them for as long as the local preference persists (Danchin et al., submitted for publication). There is thus strong selection for conforming to the majority. In our experiments, females saw all 6 demonstrator males having apparently chosen to copulate with the same male color morph and rejecting the other, revealing a very strong local preference. In such conditions, the Fisher runaway process predicts that they should build a preference for that type of males independently of the other local conditions.

**Female choosiness increased with sex ratio**

From a female’s point of view, a female-biased local sex ratio is concomitant with high competition for mates, and thus, females should accept any encountered mate more readily and more quickly. Following this logic, we found that females were quicker in initiating copulations under female-biased sex ratios. Similarly, double courtship rates, which constitute a positive proxy of female choosiness in *Drosophila*, decreased when the sex ratio shifted from male-to-female bias. However, the rate of double courtship likely depends on both sexes as males can be more or less interested in courting the female (Eastwood and Burnet 1977; Clowney et al. 2015), and females can exhibit varying choosiness (Maklakov and Arnqvist 2009). Nonetheless, because all males used during mate-choice tests were new and naive males, they had no information on the previous demonstration and thus could not have generated the observed pattern. They all saw a young virgin female, which should be attractive *per se* (Tompkins and Hall 1981). Contrastingly, observer females having been under varying sex ratios during the demonstration are more likely to have driven the observed pattern. All observer females of all experimental groups were raised in unisex groups for 3–5 days from hatching to experiment, and thus all of them experienced the same situation of acute lack of potential mates for a long period before experiments. As a consequence, the difference in choosiness observed between these groups should only come from the difference in the sex ratio and group size experienced during the ~20 min of the demonstration. In effect, demonstrations constituted the first time since sexual maturity in which observer females were in the presence of males, and vice versa, which might have made them highly sensitive to the presence of mates, potentially explaining the fact that the short demonstration period was sufficient to elicit different behavior in females according to the sex ratio and group size they experienced just before.

Results on double courtship rates and on courtship to copulation latency were probably not independent because the shorter the courtship, the shorter the time for the second male to court the female. However, globally our results support our hypothesis that varying levels of competition along the sex-ratio gradient affected female choosiness in *Drosophila*. Similarly, it was shown in several species (Berglund 1994; Passos et al. 2014; Pompilio et al. 2016), as well as in a theoretical study (Bleu et al. 2012), that females tend to maximize the chances of mating with a high-quality male in male-biased sex ratios, while minimizing the risk of remaining unmated in female-biased sex ratios. For similar reasons, the positive relationship between sex ratios and rates of both males courting the observer female during the mate-choice test appears adaptive.

**Mechanisms of sex ratio detection**

Concerning the mechanisms by which females detected the local sex ratio, the characteristics of our hexagon device implies that observer females could only perceive sex ratio visually, which is consistent with the fact that flies can recognize visual patterns and sense motion and color (Behnia and Desplan 2015; Liu et al. 2006; reviewed in Guo et al. 2017), and may even be able to visually recognize individual males and behave differently relative to them according to these males’ past experience (Loyau et al. 2012). Our results thus support the idea that *Drosophila* probably use vision in much more diverse and subtle contexts than previously thought (Loyau et al. 2012). More generally, insect cognitive capacities are being discovered as surprisingly sophisticated. For instance, honeybees *Apis mellifera* have been shown to be able to count visually (Chittka and Geiger 1995; Gross et al. 2009; reviewed in Dacke and Srinivasan 2008). The question of the existence of such skills in *Drosophila* remains open. It would be interesting to study how and for how long such specific behavior can last after the demonstration, in order to assess the dynamics of *Drosophila* mating behavior determinants.

**Effect of first-courting male on MCI**

We found the fact that the first courting male was or was not of the same color that was selected during the demonstration significantly affected mate-copying. Among the group of flies that chose after a double courtship, mate-copying indices were higher when the first
courting male had the color that was rejected during the demonstration. A female has 2 options when the first male courted: accept copulation with the courting male or wait for the second male to start courting. A copier observer female would be less likely to wait for her “non-preferred” male to court her if the first male to court her is of the color that she learnt to prefer during the previous demonstration: a situation that most often would lead to a single courtship before copulation. Contrastingly, a non-copier female or a copier female first courting by the male of her non-preferred color would both be more likely to wait for the second male to court her. Because in our analyses of mate-copying indices, it was necessary to discard mate-choice tests in which only one male courted the female before the onset of the copulation to only keep situations in which females were in a real situation of choice between the 2 males (see “Materials and Methods” section), we could thus expect a lower proportion of copiers in the group “First-courting male = 1” (i.e., females that were first courted by their preferred male), than in the group “First-courting male = 0” (i.e., females that either had no preference or that were first courted by the male of their non-preferred phenotype), as we found. Thus, our measured MCI is conservative.

Effect of atmospheric pressure on Drosophila sexual behavior

As in a previous study (Dagaef et al. 2016), we found a significant positive effect of air pressure on the rate of double courtship, and a slight positive effect on mate-copying. Mating behavior was previously found to be correlated with atmospheric pressure in D. pseudoobscura (Ankney 1984) and in D. melanogaster (Austin et al. 2014). Bad weather can mean death for small insects, and it seems adaptive for them to be able to anticipate weather variations, like air pressure changes, to find a shelter and then save energy in bad weather. Interestingly, flies from the same population were found to differ in response to air pressure: under low pressure some individual flies reduced their mating activity, while others increased it (Austin et al. 2014). Such a polymorphism might reveal phenotypic variation in relation to dispersal. Similarly, mate-copying was found to be reduced under low atmospheric pressure (Dagaef et al. 2016). Here, we further found that D. melanogaster females seem to become much less choosy under bad weather forecast (revealed by lower double courtship rates), probably because they act as quiet as possible in such cases.

In conclusion, we provide evidence that sex ratio as a proxy of female–female competition may affect female choosiness in D. melanogaster. We did not find any relationship between sex ratio and mate-copying efficiency, suggesting that mate-copying is fairly robust to variation in this environmental condition. We speculate that this may be partly because our experimental design did not allow us to separate the contradictory effects of sex ratio on the propensity to mate-copy, or because selection favors copying in general independently from sex ratio conditions. In terms of evolution, our findings suggest that females may have acquired the ability to mate-copy independently of group size and sex ratio, without hampering their capacity to adapt their choosiness to the current population sex ratio, which determines the relative availability of male partners.

Author Contributions

M.M. carried out the experiments, performed the analysis, and drafted the manuscript; S.N. contributed in the analysis and writing of the manuscript; ED and G.I. designed the experiment and jointly supervised all steps in the process. All authors gave final approval for the publication.

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