No evidence that male sexual experience increases mating success in a coercive mating system

Maider Iglesias-Carrasco 1, Rebecca J. Fox*, 1, Alan Vincent, Megan L. Head, Michael D. Jennions

Division of Ecology and Evolution, Research School of Biology, Australian National University, Canberra, ACT 2601, Australia

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Several studies have shown that sexual experience can alter a male’s mating behaviour to increase his future mating success. One explanation is that experienced males are better at courting females and inducing them to mate. Experienced males might also be better at identifying higher quality mates, although fewer studies have tested for this benefit. In both cases, however, these potential benefits of sexual experience might be partially offset by the energetic costs of courting and mating, which tend to reduce a male’s subsequent ability to invest in sexually selected traits, and thereby reduce his future attractiveness and mating success (i.e. hasten the onset of reproductive senescence). Here we used the eastern mosquitofish, Gambusia holbrooki, to test whether sexual experience elevates male mating success. We housed recently matured males either with full mating access to females (experienced males) or in the visual and olfactory presence of females with which they could not mate (naïve males). We then measured the strength of male mate choice for larger, more fecund, females, male mating behaviour (time spent chasing females and the number of copulation attempts) and insemination success. Experienced and naïve males did not differ significantly in their mating behaviour and there was no effect of sexual experience on the likelihood of mating or on the number of sperm inseminated (although experienced males had a tendency to be less successful when performing gonopodial thrusts). Experienced males in two-choice trials were, however, significantly more likely to ‘inspect’ both females and had a significantly stronger preference for larger females. Finally, we measured male immune response and growth to test for any costs of the increased mating effort that is concomitant with greater experience. Experienced males had significantly slower postmaturation growth and a significantly weaker immune response than naïve males.

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In many species a male’s ability to acquire mates depends on sexually selected morphological and behavioural traits that increase his attractiveness or ability to repel rivals (Kuijper, Pen, & Weissing, 2012). These costly sexual traits are often expressed in a context-dependent fashion in response to variation in factors such as food availability (Bonduriansky, 2007), the level of mating competition (Bretman, Gage, & Chapman, 2011), the intensity of sperm competition (Fitzpatrick & Lüpold, 2014; Kelly & Jennions, 2011), and the risk of predation (Kotiaho, 2001). In addition, it is possible that males alter their investment in sexual traits in response to experience gained during sexual encounters. For instance, it has been suggested that as males interact with females they learn to be more effective at seducing or coercing females into mating (e.g. Dukas, 2005; Dukas, Clark, & Abbott, 2006; Pérez-Staples, Martínez-Hernández, & Aluja, 2010). However, relatively few studies are designed to distinguish between learning and other factors (e.g. female preferences for older males or direct effects of male age) that might increase the mating success of more sexually experienced males.

Research on how sexual encounters influence subsequent male reproductive success has tended to focus on female mate choice for more experienced males (Edwardsson, Hunt, Moore, & Moore, 2008; King & Fischer, 2010). These studies often show that females prefer older, potentially more experienced, males (review: Brooks & Kemp, 2001). However, few studies have explored how past sexual experience influences a male’s subsequent sexual behaviour and his ability to acquire mates and successfully copulate, independent of changes in female mating preferences that

* Correspondence: R. Fox, Division of Ecology and Evolution, Australian National University, Canberra, ACT 2601, Australia.
E-mail address: rebecca.fox@anu.edu.au (R. J. Fox).
1 These authors contributed equally to this work.

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depend on his mating history (but see Balaban-Feld & Valone, 2017; Saleem, Ruggles, Abbott, & Carney, 2014).

During sexual encounters with females, males can learn to adjust behaviours such as courtship to increase their likelihood of mating (e.g. Dukas et al., 2006; Pérez-Staples et al., 2010). Similarly, in species where sexual coercion is the dominant male mating tactic, males with previous exposure to females might learn tactics that make them more successful at sneak copulations than sexually naïve males. For example, in eastern mosquitofish, Gambusia holbrooki, juvenile males reared in the absence of females made significantly more copulatory attempts as adults than did those reared with females (Bisazza, Pilastro, Palazzi, & Marin, 1996). However, this study did not quantify the potential benefits of adult male sexual experience due to improvements in mating tactics. In addition to learning how to adjust mating behaviour to increase reproductive success, sexual experience might allow males to better assess mate quality (review: Verzijden et al., 2012). Learning can happen through sexual imprinting early in life (Immelmann, 1975), but it can also occur when mature individuals gain experience with potential mates. For example, male damselflies learn to discriminate between conspecific and heterospecific females based on previous courtship interactions (Svensson, Eroukhmanoff, Kallmön, Runen, & Brodin, 2010). Similarly, in damselfly species with discrete female morphs, males prefer the female phenotype with which they have previously had a successful mating (Finke, Fargeville, & Schultz, 2007). In fruit flies, males that have previously interacted with unreceptive, recently mated females spend less time than inexperienced males courting such females in future encounters (Dukas, 2005). In sum, mating preferences are often plastic, with individuals adjusting their mate choice decisions based on their social environment (e.g. Gasparini, Serena, & Pilastro, 2013; Macario, Croft, Endler, & Darden, 2017; Mautz & Jennions, 2011; review of male mate choice in poeciliid fish: Schlupp, 2018) and past encounters with mates (e.g. Balaban-Feld & Valone, 2017; Reif, Linsenmair, & Heisenberg, 2002).

Of course, sexual experience can be costly for males due to the associated increase in mating effort, such as investment in pre- and postcopulatory sexual traits, or even learning itself (Mery & Kawecki, 2003), which can reduce the availability of resources needed for self-maintenance. Mating behaviours, such as courtship or harassment of females, can impose costs on males that include increased energy expenditure and loss of time for other activities such as foraging or evading predators (Kelly & Godin, 2001; Kolluru & Grether, 2005). Similarly, investment in ejaculates trades off with other fitness-enhancing traits and reduces the resources available for survival, precopulatory sexual traits and general maintenance (Barnes & Partridge, 2004). The energetic costs of mating that are associated with gaining sexual experience might therefore interact with age to elevate male reproductive senescence. Prolonged sexual activity has been related to reduced male longevity, a decline in body mass and weaker immune response in many species (e.g. Bleu, Gamelon, & Saxer, 2016; Foo, Nakagawa, Rhodes, & Simmons, 2017; Olsson, Madsen, & Shine, 1997; Roberts, Buchanan, & Evans, 2004; Rolff & Siva-Jothy, 2002; South, Steiner, & Arnqvist, 2009). Several studies in invertebrates have also demonstrated that there are fitness costs associated with commencing reproduction at an early age (review: Wedell, Gage, & Parker, 2002). Many of these studies are, however, correlational. Experiments that measure the costs of mating for individuals that have experienced differences in mating opportunities over significant portions of their reproductive life span are needed, but few such studies exist.

Here we tested whether sexual experience increases a male’s mating success by improving his ability to obtain copulations, despite any associated energetic costs of interacting with females that otherwise reduces investment in costly sexual traits. The eastern mosquitofish is an excellent model system to explore this question (e.g. Bisazza et al., 1996). Males rarely court, and instead pursue and coerce females into mating. They make frequent attempts to copulate (up to one attempt/min, Wilson, 2005) by approaching females from behind and thrusting their gonopodium (a modified anal fin used to transfer sperm) towards her gonopore. This makes it less likely that changes in male mating success reflect changes in female mating preferences based on a male’s mating history. We maintained male mosquitofish from maturity onward for a prolonged period either with continuous mating access to females (‘experienced’ males) or in the visual and olfactory presence of females but without physical access to prevent mating (‘naïve’ males). We predicted that direct sexual experience would benefit males by (1) improving their ability to approach and inseminate females and (2) increasing their ability to assess which females are more profitable mates (e.g. bigger females which are more fecund, Mautz & Jennions, 2011), but that (3) sexual experience might impose an associated cost since experienced males have invested more energy in sperm replenishment and harassment, potentially reducing their ability to invest in other life history traits such as immune function and growth.

METHODS

Ethical Note

The collection of animals was conducted under a Scientific Licence from the Australian Capital Territory (ACT) Government, granted under Section 21 of the Fisheries Act 2000, licence number FS20174. Collection, housing and experimental work conducted as part of the study followed the ASAB/ABS guidelines for the treatment of animals in behavioural research. Information about individuals’ housing conditions are described below. Housing conditions, handling and experimental monitoring were conducted to maximize the animals’ welfare. All experimental procedures were carried out under approval from ANU Animal Ethics Committee (Approvals A2015/07 and A2018/27) and complied with existing laws regulating the treatment of vertebrates in Australia.

Origin and Maintenance of Animals

In November 2017 we collected immature male (N = 150) and recently matured female (N = 300) mosquitofish from ponds in Australia, where it is classified as an invasive pest species. Fish were collected using dip-nets and transported back to the laboratory within 1 h of collection in groups of 30 individuals in 15-litre containers with portable aeration (size at collection: males 15–20 mm; females 30–45 mm). Males were transferred to individual 1-litre tanks, and recently matured females were housed in single-sex 60-litre tanks (50 individuals per tank) containing artificial plants for enrichment and an under-gravel filter system for aeration. We checked for recovery from capture and transport by confirming that all individuals fed normally within 12 h of being established in tanks. Fish were maintained in these conditions on a 14:10 h light:dark cycle at 27 °C (±1) for at least 2 weeks before the experiment. The actual period depended on the time each male took to reach sexual maturity (range 2–4 weeks). Experimental males were fed ad libitum with Artemia salina nauplii twice daily, and stimulus females were fed ad libitum twice daily with commercial fish flakes in the morning and A. salina nauplii in the afternoon.

Males were checked daily to determine when they reached sexual maturity (i.e. fully formed gonopodium with distal spines). At maturity, males were individually transferred to a 4-litre
aquarium containing gravel and artificial plants, and randomly assigned to one of two social environment treatments: (1) ‘experienced’ males had access to a female with which they were able to mate ($N = 54$); (2) ‘naïve’ males were housed with a female behind a mesh screen to prevent mating, but allow visual contact and the dispersal of chemical cues ($N = 51$). All stimulus females were similar in size (weight $0.55–0.75$ g; ‘large’ females sensu Head, Vega Trejo, Jacomb, & Jennions, 2015) and were replaced weekly to maintain the males’ interest in mating. We assumed that ‘experienced’ males mated regularly with females. This is highly likely because males incessantly attempt to mate (e.g., Wilson, 2005). In this study the artificial plants in the tank provided a partial refuge for the female to mitigate continual male harassment. There was no initial size difference between males in the two treatments ($t_{103} = 0.093$, $P = 0.355$). After 16 weeks (approximating the typical reproductive life span of males in the wild for our study population, Kahn, Kokko, & Jennions, 2013) males were transferred back to individual 1-litre tanks for 5 days to allow for full sperm replenishment (O’Dea, Jennions, & Head, 2014). We then ran behavioural assays to test the effect of the male’s mating experience on: (1) the strength of his mating preference for larger females (on average, males prefer larger females; e.g., Mautz & Jennions, 2011); (2) his propensity to mate when presented with a female (i.e., chasing behaviour); and (3) his actual insemination success. Finally, we measured his immune response with a phytohaemagglutinin (PHA) injection assay, and recorded his standard length (SL) as a measure of growth (because there was no initial size difference between treatments, this absolute size measure is synonymous with growth). Each behavioural test was conducted by a different observer to prevent biased assessment of the relationship between male performance across tests. All data were collected blind to a male’s experience treatment.

**Male Mating Preferences: Two Mate-choice Trials**

To test whether mating experience affected male mating preferences we calculated association time with the larger of two females in two-choice trials. The male was introduced to the middle of an aquarium (49 × 20 cm and 22 cm high) containing two test sections, each 7 × 20 cm and 22 cm high. One section housed a large virgin female (>45 mm SL) and the other a small virgin female (<40 mm SL). The side containing the larger female was randomly assigned for each trial. Each female was initially behind a mesh divider and an opaque screen. Aquarium walls were lined with black plastic to reduce disturbance. Before the trial, the male was placed in a plastic container in the centre of the test aquarium. After 5 min of acclimation we carefully removed the container and the opaque screens and then recorded for 10 min the time the male spent <$5$ cm from the mesh separating him from each female. In some trials ($N = 22$ out of 105 total) males did not associate with one of the females. In these cases, we could not be sure that the male had seen (‘inspected’) both females and therefore that the data represented a preference for one female over the other. We therefore adopted a two-step approach to analysing our male mating preference data (see Statistical Analyses).

**Male Mating Effort**

Males were then transferred to a 2-litre tank containing a stock female (mass: $0.55–0.75$ g) behind a mesh barrier. All females had originally been collected from the wild, but then housed in single-sex aquaria for >160 days to ensure that they were not gravid and were no longer storing sperm from a previous mating (maximum recorded duration of sperm storage is 6 months, Constanz, 1989). The male was given 5 min to acclimate after which the mesh partition was removed, and the pair were free to interact. For 10 min we recorded: (1) time spent chasing or associating with the female (<1 body length and oriented towards her); (2) number of mating attempts (gonopodial thrusts towards her gonopore); and (3) number of successful gonopodial thrusts (contacted the gonopore).

**Male Insemination Success**

Males were then left with the female for another 20 min, after which she was removed from the tank, anaesthetized in an ice slurry and her gonoduct flushed with 3 μl of saline solution (0.9% NaCl) to check for sperm. All females re-attained their swimming equilibrium within 2 min of being returned to their tank and behaved normally and fed within minutes of recovery from anaesthesia. There was no mortality. The sample collected was placed in an Eppendorf tube with another 3 μl of saline solution and vortexed for 30 s to break up any sperm bundles. We then pipetted 3 μl of the vortexed solution onto a 20 micron capillary slide (Leja), placed it under a microscope (100× magnification) and used a CEROS II sperm tracker (Hamilton Thorne Research, Beverly, MA, U.S.A.) to count the sperm present for five subsamples per slide (repeatability: $r = 0.968 ± 0.012$). The mean number of sperm was our measure of the relative quantity of sperm transferred.

**Male Immune Response and Body Size**

Finally, we assessed the cell-mediated immunity of males using a PHA injection assay to measure their inflammatory response. This assay has been successfully used in other fish (e.g., Clotfelter, Ardia, & McGraw, 2007). We first validated this technique for G. holbrooki by comparing the inflammatory response of males injected with either PHA or phosphate-buffered saline (PBS) as a control. There was a clear difference in inflammation between PHA-injected males and PBS-injected control males ($t_{123} = 6.98, P < 0.001$).

We anaesthetized experimental males in an ice slurry then measured the thickness of the body at the posterior end of the dorsal fin with a pressure-sensitive spessimeter (Mitutoyo 547-301; accuracy: 0.01 mm; average of five measurements per fish). We then injected 0.01 mg of PHA dissolved in 0.01 ml of PBS into the right side of the caudal peduncle. Males were returned to their corresponding containers for 24 h, after which we re-measured the thickness of the body at the same point to calculate the difference between pre- and post-injection measures (inflammation). Any tissue swelling provoked by the immune response to the injection of the PHA disappeared after 72 h. All the males behaved normally and fed within half an hour of recovering from the anaesthesia. To control for allometry we recorded male body size (SL, ±0.01 mm) using electronic callipers. At the end of the experiment all focal individuals were returned to their 4-litre treatment tanks and after 21 days were euthanized with an overdose of Aqui-S solution (in compliance with Australian government legislation preventing the release of invasive pest species). Nonfocal individuals were returned to stocking tanks to be used, where possible, in subsequent experiments.

**Statistical Analyses**

We analysed data using separate generalized linear models (GLM), with the choice of error function for each model based on inspection of the distribution of the dependent variable and confirmed via Q-Q plots of model residuals. When testing for the effect of mating experience on a preference for large versus small females, we tested first for the effect of experience on the propensity to inspect both females (GLM, binomial error). For those
trials in which both females were inspected ($N = 83$, consisting of 48 ‘experienced’ and 35 ‘naïve’ males), we then tested for the effect of male mating experience on the proportion of time spent with the large female (quasibinomial error, cbind function, see Booksmythe, Backwell, & Jennions, 2013; Deere, Grether, Sun, & Sinheimer, 2012; Vega-Trejo, O’Dea, Jennions, & Head, 2014; Wacker, Østlund-Nilsson, Forsgren, Newport, & Amundsen, 2016). We ran separate GLMs to test for the effect of male experience on time spent chasing the female (quasi-Poisson error), number of gonopodial thrusts (quasi-Poisson error), the proportion of successful gonopodial thrusts (quasibinomial error, cbind function), whether or not a male transferred sperm (quasibinomial error), the quantity of sperm transferred by the 23 ‘naive’ and 19 ‘experienced’ males that successfully inseminated a female (Gaussian error), immune response (Gaussian error) and growth over the treatment period (i.e. final size; Gaussian error). Sperm quantity and immune responses were log transformed to meet assumptions of normality. In all models male mating experience (naïve or experienced) was treated as a fixed factor.

Since male body size might be an important factor driving the behavioural trials, we initially ran models including male size as covariate. As including this variable did not affect any of the results, for simplicity we report models without the covariate. Body size was, however, included as a covariate in the immune response model as it was correlated with the amount of postinjection inflammation ($r = 0.28$, $P = 0.004$). We excluded one outlier from the immune response model because it had an extreme value that is likely to reflect measurement error or a mistake in data transcription. The mean inflammation value was 0.174 ± 0.114 SD ($N = 105$ males), while the outlier had a value of 0.572 (i.e. >3.5 standard deviations from the mean). However, to ensure transparency we also report model results based on the full data set.

Figure 1. The effect of previous sexual experience on (a) the time spent associating with the larger of two females in two-choice trials beyond that expected due to chance alone (i.e. if males spent half of each trial (300 s) with each female; the ‘extra time’ is therefore the actual time minus 300 s); (b) time spent chasing the female; (c) number of mating attempts; (d) sperm transfer success. Means are shown ±SE.
All analyses were conducted using R 3.4.0 (The R Foundation for Statistical Computing, Vienna, Austria, http://www.r-project.org) with alpha significance set at 0.05.

RESULTS

When males were given a choice between two females, experienced males were significantly more likely to inspect both females ($\chi^2_{104} = 6.035, P = 0.014$). Of those males that inspected both females, experienced males spent significantly more time than naïve males in association with the larger female ($t_{104} = 2.628, P = 0.010$; Fig. 1a). For naïve males, there was no difference in the proportion of time spent with the large and small female ($t_{104} = 0.221, P = 0.826$). However, a male’s previous mating experience did not affect his mating effort. There was no difference between experienced and naïve males in the time spent chasing a female ($t_{103} = 0.135, P = 0.893$; Fig. 1b), in the number of mating attempts ($t_{103} = 1.332, P = 0.186$; Fig. 1c), in the likelihood of transferring sperm ($t_{103} = 1.024, P = 0.308$; Fig. 1d) or, if insemination occurred, the quantity of sperm transferred ($t_{104} = 0.642, P = 0.524$). The proportion of successful gonopodial thrusts made by naïve males, however, was marginally nonsignificantly greater than that of experienced males ($t_{103} = 1.886, P = 0.059$).

The opportunity to mate with females had a negative effect on postmaturation growth, such that experienced males were significantly smaller than naïve males ($t_{104} = 3.857, P < 0.001$; Fig. 2a). Excluding one very large male (>24 mm) in the naïve group did not change this result ($t_{102} = 3.696, P < 0.001$). In addition, experienced males had a significantly weaker immune response than naïve ones ($t_{101} = 2.049, P = 0.043$). However, if we included an outlier in our model there was no longer a significant effect of the experience treatment on male immune response ($t_{102} = 1.647, P = 0.103$; Fig. 2b).

DISCUSSION

By manipulating the access of male G. holbrooki to females over their entire adult life span prior to testing, we explored whether direct sexual experience increased male mating success due to learning how to be more effective at gaining copulations. More specifically, we tested how adult sexual experience affected a male’s ability to mate, independent of female mating preferences for males with different mating histories. This claim is based on the assumption that female choice for more experienced males is unlikely, owing to the coercive mating system. The use of adults in our experiment contrasts with an earlier study in G. holbrooki that explored how exposure to females when males were still juveniles affected their subsequent adult sexual behaviour (Bisazza et al., 1996). We tested whether sexual experience increased a male’s reproductive success in terms of both his ability to gain copulations and his ability to discriminate between potential mates based on their size. We predicted that if interacting with females allows males to learn, then sexually experienced male G. holbrooki would be more likely to achieve coercive copulations and would show stronger preferences for more fecund mates (i.e. larger females). Our prediction about female choice is based on earlier work showing that males are more likely to copulate when they attempt to mate with larger females (Filastro, Giacomello, & Bisazza, 1997). Males should therefore learn that larger females are more profitable, at least in terms of the likelihood of mating (but see Fitzpatrick & Servadio, 2018 who reviewed the evidence that more fecund females are ‘devalued’ due to an increase in shared paternity if males preferentially mate with them). We also sought to confirm that sexually experienced males pay a cost due to their greater past mating effort.

As predicted, experienced males differed from naïve males in their mate preferences, spending significantly more time associating with the larger of two females. Experienced males also grew more slowly and had a weaker immune response than naïve males, confirming that mating effort is costly. However, contrary to our predictions, experienced males did not make significantly more copulation attempts than naïve males, nor were they more likely to inseminate females. Intriguingly, the proportion of successful gonopodial thrusts made by naïve males was marginally nonsignificantly greater than that of experienced males. There is therefore

Figure 2. Effect of previous sexual experience on (a) male body size and (b) immune response (raw data are shown). The red point in (b) represents a male with an extreme immune response value.
no evidence that experienced males with the opportunity to interact with females over their adult life span had an advantage over naïve males because they learned to mate more successfully. Previous studies of learning in relation to mate choice have focused on species recognition (e.g. Svensson et al., 2010), or how social cues influence mating decisions (via mate choice copying: e.g. Schlupp & Ryan, 1996; Widemo, 2006; Witte & Ryan, 2002; review: Schlupp, 2018). Fewer studies have examined the role of learning to discriminate between conspecific mates based on their quality and/ or the likelihood of obtaining a copulation. Mating is generally costly for males due to sperm and energy expenditure, so when the costs of choice are low, as is always the case when two females are simultaneously available (Barry & Kokko, 2010), males are expected to direct their mating efforts towards high-quality females (Bonduriansky, 2001). In our study, experienced males had a stronger preference for associating with larger, more fecund, females and, hence, a stronger ability to discriminate between partners based on their quality under low-cost conditions (but see Fitzpatrick & Servedio, 2018 for the potentially confounding effect of greater shared paternity). There are several potential explanations for this result. First, the difference between experienced and naïve males in time spent in association with larger females might be related to learning about variation in actual female quality (Verzijden et al., 2012). However, this would require males to have had previous access to females of different phenotypes and being able to ascertain the number and quality of offspring they sired, which is implausible. Second, experienced males might have learnt that they are more successful at copulating with larger females as a greater size difference between males and females increases a male’s mating success (see Bisazza & Marin, 1995; Pilastro et al., 1997). Third, weaker discrimination by naïve males could be a strategy to reduce lost mating opportunities at the end of their reproductive life span by minimizing the time spent searching for, or inspecting, females. Similarly, the fact that naïve males were less likely to ‘inspect’ both females could mean that ‘inspection’ is learned.

Contrary to findings in some species (e.g. Dukas, 2005; Edvardsson et al., 2008; Saleem et al., 2014), sexual experience did not improve male G. holbrooki mating ability, when measured as: (1) the number of gonopodial thrusts; (2) the proportion of successful thrusts; (3) the rate of successful sperm transfer; and (4) the amount of sperm transferred. Although mating in G. holbrooki appears to result from male coercion, there is some evidence that the amount of sperm transferred. Although mating in G. holbrooki has often been demonstrated in invertebrates (e.g. Dukas, 2005; Fedorka, Zuk, & Mousseau, 2004; McKean & Nunney, 2001; Rolf & Siva-Jothy, 2002; review: Lwniczak et al., 2007) and in some vertebrates (e.g. birds: Roberts et al., 2004; mammals: Muehlenbein & Bribiescas, 2005).

In G. holbrooki there was no effect of sexual experience on the time males spent chasing a female or on the rate of copulation attempts. This contrasts with other species where past encounters with females affect male mating effort (Balaban-Feld & Valone, 2017; Saleem et al., 2014). For example, in fruit flies, experienced males were less likely to behave aggressively and to use coercion against females than were naïve males (Baxter & Dukas, 2017). Alternatively, naïve males might have behaved more aggressively because they perceived their future reproductive opportunities to be low (Fischer, Perlick, & Galez, 2008) as, despite exposure to females, they had previously failed to mate. For example, Bisazza et al. (1996) raised juvenile male G. holbrooki in the presence or absence of females and found that, upon adulthood, naïve males made significantly more copulatory attempts and more successful copulatory attempts than experienced males, but copulation efficiency was broadly equivalent for naïve and experienced males. In our study, naïve and experienced males did not differ in the number of copulation attempts, but the proportion of successful attempts by naïve males was marginally nonsignificantly greater than that of experienced males (P = 0.059). There are many untestable reasons why the results of our study and that of Bisazza et al. (1996) might differ (e.g. population differences, test conditions, sampling errors). The most obvious difference, however, is whether males were exposed to females to gain experience as juveniles (Bisazza et al., 1996) or adults (our study). Nevertheless, both studies suggest that male G. holbrooki do not significantly improve the effectiveness of their coercive behaviour by having the opportunity to interact with females (i.e. learn).

In support of our prediction that sexually experienced males incur costs from their greater mating effort, experienced male G. holbrooki grew more slowly than naïve males and tended to have a weaker immune response. Our results suggest that continuous harassment of females and sperm replenishment are therefore costly in species with coercive mating tactics. Studies of other poeciliids have similarly shown that mating effort can reduce male growth (Jordan & Brooks, 2010). Our finding of a reduced immune response in experienced males is consistent with the idea that mating effort suppresses immunity (Folstad & Karter, 1992). This has often been demonstrated in invertebrates (e.g. Fedorka, Zuk, & Mousseau, 2004; McKean & Nunney, 2001; Rolf & Siva-Jothy, 2002; review: Lwniczak et al., 2007) and in some vertebrates (e.g. birds: Roberts et al., 2004; mammals: Muehlenbein & Bribiescas, 2005).

Conclusions

Past sexual experience did not confer an advantage to male G. holbrooki by improving their ability to inseminate a female. This suggests that having the opportunity to interact with females did not cause males to learn how to be more effective at completing copulations and transferring sperm. There was, however, a potential benefit of interacting with females if the mate choice experiment is interpreted as evidence that sexually experienced males improved their ability to direct their mating effort towards large, more fecund females. However, this type of male preference might be disadvantageous (or confer no advantage) if males more often copulate with preferred, large females, thereby reducing the mean paternity per male (Fitzpatrick & Servedio, 2018; Servedio, 2007). In addition, we confirmed there are significant costs of mating effort for males over their reproductive life span. Future studies examining the effects of lower immunity and slower growth on male lifetime reproductive success would be useful to further our understanding of life history trade-offs (Brooks & Garratt, 2017).


