Be prudent if it fits you well: male mate choice depends on male size in a golden orb weaver spider

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9 Abstract

 Male preference for high-quality females is expected to evolve when male reproductive potential is restricted. However, when there is competition among males, some models predict the evolution of assortative male mate choice, in which good competitors choose high quality females while poor competitors choose lower quality females to avoid competition. In *Trichonephila clavipes* spiders, males have limited sperm supply and fight for access to females.

2. We tested whether female quality and male size (a proxy of fighting ability) influence male
decisions in *T. clavipes*.

3. We used field experiments in which males could choose among two available females in a
scenario free of competition. Females differed in body size and recent pairing status (whether
the female was accompanied by a male before the experiment).

4. We found that males choose their mates based on both female size and female recent pairing
status. Importantly, male mate choice exhibited plasticity, and varied with male size, as large
males preferred larger females that were recently unpaired, medium-sized males showed no
preference, and small males preferred smaller, recently paired females.

5. Because all females appear to attract males, we predict that variation on male mate choice
attenuates sexual selection on females. Our findings confirm the prediction of variable male
mate choice when there is male-male competition and male reproductive potential is
restricted, a pattern that may be common, but hard to detect.

Key-words: male mate preference, mating tactics, intrasexual competition, fecundity, sperm
competition, *Nephila clavipes*.

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32 Introduction

The Darwin-Bateman paradigm that underlies the traditional sexual selection theory posits that male reproductive potential is virtually unbounded, and male reproductive success is limited only by female availability (reviewed in Parker & Pizzari, 2015). Thus, males are predicted to seek copulations with as many females as possible, irrespective of female quality. This prediction assumes that copulation is virtually without costs for males, which is not necessarily true (Dewsbury, 2005; Tang-Martinez, 2016; Tang-Martinez & Ryder, 2005). Several factors may restrict male reproductive potential, such as sperm limitation (Dewsbury, 1982; Perry, Sirot, &

Wigby, 2013; Wedell, Gage, & Parker, 2002), costly mate search (Kasumovic, Bruce, 40 Herberstein, & Andrade, 2007; Lane, Boutin, Speakman, & Humphries, 2010), provision of 41 paternal care (Berglund, Rosenqvist, & Svensson, 1989), provision of nuptial gifts to females 42 (Cratsley, Rooney, & Lewis, 2003), and sexual cannibalism by females (Andrade, 1996). These 43 factors can limit the number of females a male can copulate with or diminish the benefits of 44 copulating with a high number of females. When male reproductive potential is constrained, if 45 there is variation in female quality (*i.e.*, some females provide more fitness benefits than others 46 to males), one would expect the evolution of male mate choice (Bonduriansky, 2001; Edward & 47 48 Chapman, 2011).

Male mate choice is often based on the number or quality of offspring that a female can 49 produce (Bonduriansky, 2001; Edward & Chapman, 2011). In populations in which male mate 50 choice occurs, males often choose females based on their body size, as larger females typically 51 possess more ova (Arnaud & Haubruge, 1999; Bonduriansky, 2001; Wedell et al., 2002). 52 Moreover, males may also use female social context as a mate choice criterion. For instance, the 53 presence of another male with a female could influence male mating decisions due to two main 54 reasons (Mautz & Jennions, 2011). First, in some mating systems a male accompanying a female 55 56 may aggressively defend her from other males, so that approaching an accompanied female would result in fighting with the resident male, which may be energetically costly (Austad, 1983; 57 Kelly, 2006). Second, even if a male manages to access a recently accompanied female, he will 58 59 probably face sperm competition, which can decrease the benefit of copulating with that female, especially if there is first-male sperm precedence (Bonduriansky, 2001). Therefore, males 60 generally prefer females that are not being guarded or that have not mated recently (e.g., 61 62 Schneider, Lucass, Brandler, & Fromhage, 2011; Schwagmeyer & Parker, 1990).

63 Male mate choice is often associated with restricted male reproductive potential (Bonduriansky, 2001), absence of male-male competition (e.g., Gwynne, 1981) and high female 64 availability (Dougherty & Shuker, 2015). However, male mate choice can occur even when male 65 intra-sexual competition is strong (e.g., Bel-Venner, Dray, Allainé, Menu, & Venner, 2008). 66 Theoretical models predict that the occurrence of male-male competition is one of the 67 mechanisms that can generate variation in male mate choice (Fawcett & Johnstone, 2003; 68 Härdling & Kokko, 2005; Venner, Bernstein, Dray, & Bel-Venner, 2010). By incorporating 69 male-male competition, theoretical models predict that good competitors would prefer high 70 quality females, whereas bad competitors would prefer low quality females (Fawcett & 71 Johnstone, 2003; Härdling & Kokko, 2005; Venner et al., 2010). In this scenario, bad 72 competitors would be performing prudent mate choice (sensu Härdling & Kokko, 2005), in 73 which low-quality males actively choose to mate with low-quality females as a mechanism to 74 avoid competition with other males (e.g., Wada, Arashiro, Takeshita, & Shibata, 2010). 75

Here, we investigate mate choice by males of the golden silk orb-weaver spider 76 Trichonephila clavipes (formerly known as Nephila clavipes, see Kuntner et al., 2018). In T. 77 clavipes, adult males live on female webs and fight against male intruders to monopolize access 78 79 to the female (Christenson & Goist, 1979). Males of this species face high mating costs and severe mate limitation for two main reasons. First, males suffer high mortality risk when 80 traveling between female webs (Vollrath, 1980; Vollrath & Parker, 1992), and second, males are 81 82 strongly sperm limited because spermatogenesis ceases after maturation (Christenson, 1989; Michalik & Rittschof, 2011). Also, T. clavipes males show great variation in body size: within a 83 population, one male can be 20 times heavier than other males (this study). Larger males possess 84 85 higher fighting ability and are more effective in guarding a female (Constant, Valbuena, & Rittschof, 2011). Moreover, adult females in this species vary widely in body size, which is correlated with fecundity (Head, 1995; Honěk, 1993). Females are also polyandrous, so that there is risk of sperm competition (Vollrath, 1980), a factor that should influence males' mating decisions. We expect *T. clavipes* males to perform male mate choice, because their reproductive potential is severely restricted and females vary in quality. Additionally, it is also possible that *T. clavipes* males vary in their choosiness according to body size, because there is male-male competition for access to females and male body size is correlated with their competitive ability.

In this study, we performed field experiments to investigate male mate choice in T. 93 94 *clavipes.* In these experiments, we tested whether female quality and male fighting ability (*i.e.*, size) influence the likelihood that a male would choose a given female. We have two alternative 95 hypotheses. Because all males suffer high mating costs, our first hypothesis is that all males are 96 equally choosy and select high quality females. However, considering the strong competition 97 among males, our second hypothesis is that male mate choice varies according to male 98 competitive ability, so that the most competitive males prefer high-quality females, whereas the 99 100 least competitive males would prefer low-quality females, while males with average competitive ability would likely be non-choosy. 101

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103 Material and methods

104 *Study species*

In our study site, *T. clavipes* is univoltine (*i.e.*, one generation per year) and its reproductive season begins in December and ends in June (P.P. pers. obs.). Females live for three to four months as adults (Christenson & Cohn, 1988), and then take 30 to 40 days to lay their first clutch after their last moult. Adult males live considerably less than females (ca. three weeks after the
last moult) and stop constructing their own webs to search for sexual partners (Brown, 1985). In
our study population, males can visit up to six female webs during their lives (Del Matto and
Santos, pers. obs.). However, as males have a limited sperm supply (Michalik & Rittschof,
2011), it is probable that they can only fertilize a few females. In some cases, males can become
monogynous if they use all their sperm with one female (Christenson & Cohn, 1988).

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115 *Study site and maintenance of study animals*

116 We conducted our experimental trials (see details below) on the webs of female T. clavipes occurring naturally on the gardens surrounding the Zoology Department building in the campus 117 of the University of São Paulo, in São Paulo, Brazil (23.564° S, 46.729° W). We conducted the 118 experiments in two consecutive reproductive seasons (March to May 2017; February and March 119 2018). Early in both seasons (December and January), we collected immature males on the 120 121 gardens and kept these males in the laboratory until sexual maturation to use them in the experimental trials. Males were individually kept in 250 ml plastic cups in controlled conditions 122 (14h:10h light/dark photoperiod, 25° C) on a diet of *Drosophila* flies. We provided three flies 123 and sprayed the male spiders with water every two days. All cups had their tops covered with 124 fine mesh and the inside of the cups was roughened to facilitate climbing of the spiders and silk 125 126 attachment. We recorded the date of the last moult of each male.

127 *Female quality proxies and morphometric measurements*

We use the term "female quality" to refer to the fitness output that a male gains when mating with a female. Thus, female quality can be influenced by both female phenotypic traits and the 130 female's social context. Here, we evaluated female quality using two independent proxies: 131 female body size and recent pairing status (*i.e.*, if she had a male in her web prior to the mate choice experiment). Female body size is positively related to fecundity (Head, 1995; Honěk, 132 133 1993). Consequently, female body size is positively related to female quality. Moreover, because body size does not change after maturation in this species, female body size represents an 134 intrinsic and fixed component of female quality. On the other hand, female pairing status 135 (whether a female is accompanied by a male or not) can vary throughout female adult life, as 136 males come and go from females' webs. We assume that recently paired females were of lower 137 138 quality, because from the point of view of an approaching male, a recently paired female represents two disadvantages: a possible fight with the resident male and a greater risk of sperm 139 competition due to probable recent copulation. As males deposit their own silk on the female 140 web, an approaching male may perceive chemical cues of the presence of a competitor male, 141 even if the competitor is not there anymore (Schneider et al., 2011). 142

To assess female body size, we photographed each female on the day of the beginning of 143 144 each trial (a ruler was placed by the female for scale). We used these photographs to measure female cephalothorax width (mm) using the software ImageJ (US National Institutes of Health, 145 146 Bethesda, MD, http://imagej.nih.gov/ij). We assessed female recent pairing status by recording whether there was at least one male on each female's web immediately before we started each 147 trial. We note that the variable recent pairing status is different from reproductive status. We did 148 149 not collect systematic data about each female's mating history, thus we cannot infer whether 150 unpaired females were virgin. In fact, it is highly likely that all of the females used in our experiments had already copulated at least once before the trials. Hence, what we classified as 151 152 "recently unpaired females" did not have any male on their web on the day of the beginning of trials, but possibly copulated before. To estimate male fighting ability (positively related to male body size; Constant et al., 2011), we measured body mass of focal males that were used in our experiments using a digital scale (to the nearest 0.1 mg) on the day of the beginning of each experimental trial.

157 *Male mate choice experiment*

To investigate male mate choice in T. clavipes, we conducted a field experiment in which we 158 allowed virgin adult males to choose between two females that differed in body size. Focal 159 females could also differ regarding their recent pairing status. Females often build their webs 160 intertwined to other female webs (*i.e.*, aggregated webs; Robinson & Mirick, 1971), and we used 161 these natural aggregations to conduct our trials. For each trial, we selected two females that had 162 163 their webs close to one another and that shared at least one silk-thread. We individually marked 164 these females on the dorsal side of their abdomen with water-based paint (Ziggs' Posterman markers) in order to identify them during the trial. We also removed any males that were on the 165 webs prior to the beginning of the trials, so the males being tested (hereafter, focal males) would 166 167 be in a scenario free of competition. However, because males deposit their own silk on the female web (Schneider et al., 2011), we assume that focal males can detect the cues of previous 168 males in the web. 169

Before we started each trial, we placed the focal male in a plastic container to carry him from the lab to the trial site. We initiated each trial by placing a wooden stick in the container so that the focal male could climb it, leading the male to an intersection of silk-threads connecting the webs of the two females. We evaluated male choice in two moments. We assessed the *initial male choice* by recording to which female the male moved towards within 15 minutes after the focal male reached the intersection of silk-threads (*i.e.*, beginning of the experiment). Our total 176 sample size for the initial choice trials was 73; both focal females had the same recent pairing status in 51 trials, whereas focal females differed in recent pairing status in the remaining 22 177 trials. We only considered that a male made a choice when he moved at least 30 cm towards one 178 of the females. We chose this distance arbitrarily because males would often stop moving and 179 consequently would take a long time to reach a female web, especially in windy days. Our 180 181 methodology is very similar to most investigations on male mate choice in spiders that are conducted in laboratory conditions and mainly evaluate male decisions at a single moment using 182 female silk (e.g., Gaskett, 2007; Rittschof, 2011; Schneider, Zimmer, Gatz, & Sauerland, 2016). 183 184 However, because we conducted the experiment in the field, in which female web construction date was not manipulated, and to account for possible manipulation stress on focal males, we 185 also decided to assess male choice at a posterior moment. Thus, on the following day (16 to 22 186 187 hours after the beginning of the trial), we observed the *late male choice* by noting which female the male was guarding. Our total sample size for the late choice trials was 30; both focal females 188 had the same recent pairing status in 18 trials, whereas focal females differed in recent pairing 189 190 status in the remaining 12 trials. The decrease in sample size (72 to 30 trials) is because we only considered late male choice trials as valid when both experimental females were present on the 191 192 original trial site on their individual web and the webs were still connected to one another. Moreover, we excluded from our analyses trials in which non-experimental males (i.e., other 193 than the focal male) were present with any of the experimental females, as they could influence 194 195 the focal male mating decisions.

196 Statistical analysis of male mate choice experiment

We investigated whether male mate choice in *T. clavipes* depends on femalecephalothorax width, female recent pairing status and male body mass by testing two alternative

199 predictions. We used male body mass as a continuous variable in all of our analyses. However, 200 we use categorical terms of this continuum (*i.e.*, high, medium, and low body mass males) simply as a way to better explain our predictions. Our first prediction is that all males, regardless 201 202 of body mass, due to their high mating costs will pair preferentially with recently unpaired females with wider cephalothorax. Alternatively, our second prediction is that male body mass 203 204 influences male mate choice in a way that high body mass males pair preferentially with recently unpaired females with wider cephalothorax, medium body mass males are unselective regarding 205 female cephalothorax width and recently pairing status, and low body mass males pair 206 207 preferentially with recently paired females with narrower cephalothorax (the opposite choice of heavy males). The reasoning being that as male body mass increases, so does male capacity to 208 monopolize their mates by fighting off intruders, leading to exclusive paternity with a female. 209

We tested these predictions using a modified version of the model proposed by Muniz, 210 Santos, Guimarães, Nakagawa, & Machado (2017), which is a statistical model of comparative 211 mate choice (*i.e.*, it assumes that the choosing individuals perform their decisions by comparing 212 at least two available options). The response variable of the model was the identity of the chosen 213 female, and the predictors included characteristics of focal males and of the females available in 214 215 each trial. Therefore, we built our statistical model based on three main assumptions: (1) that males performed choice by comparing the two available females; (2) that males can assess 216 female size and recent pairing status; and (3) that pairing decisions can be influenced by the traits 217 218 of the available females and male selectivity, whereas male selectivity can be influenced by male traits. We included two female traits in the model: cephalothorax width (continuous) and pairing 219 status (categorical: recently paired or recently unpaired, see details in the "Female quality 220 221 proxies and morphometric measurements" section above). Hence, the preferences of a male are

represented by two selectivity values: S_1 , selectivity for female cephalothorax width; and S_2 , selectivity for female recent pairing status. Positive S_1 values represent preference for females with wider cephalothorax, while negative values represent preference towards females with narrower cephalothorax. Similarly, positive S_2 values represent preference towards previously unpaired females, whereas negative S_2 values represent preference for previously paired females. In both cases, a value of zero represents no male choice.

Given that we hypothesized that male body size could influence their pairing decisions, we assumed that both S_1 and S_2 were functions of male body mass. Additionally, although we had no reasons to expect differences in male choosiness between years, we added the season in which each trial was conducted as a predictor in the models to control for this potential source of variation (parameters *C* and *F* below). Given all that, we calculated S_1 and S_2 values for each male with the following equations:

234
$$S_{1(i)} = A + B * m_{(i)} + C * y_{(i)} (Eq. 1)$$

- $S_{2(i)} = D + E * m_{(i)} + F * y_{(i)}$ (Eq. 2)
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- 237

Where $m_{(i)}$ is the mass of male *i* and $y_{(i)}$ is the season in which the trial was conducted, and *A*, *B*, *C*, *D*, *E*, and *F* are the model parameters (that we needed to fit). Given these selectivity values, the probability $P_{(i,j)}$ that a male *i* will choose female *j*, in a trial with females *j* and *k*, is a function of $S_{I(i)}$, $S_{2(i)}$ and the values of female cephalothorax width *f*, and female recent pairing status *g* of both females. The probability $P_{(i, j)}$ was calculated as follows:

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$$P_{(i,j)} = \frac{\exp(S_{1(i)} * f_{(j)} + S_{2(i)} * g_{(j)})}{\exp(S_{1(i)} * f_{(j)} + S_{2(i)} * g_{(j)}) + \exp(S_{1(i)} * f_{(k)} + S_{2(i)} * g_{(k)})}$$
(Eq. 3)

Whereas the probability that the male will choose female k is calculated as $P_{(i,k)} = 1$ -

247 $P_{(i,j)}$.

The model parameters can be interpreted as follows. On equation 1, the intercept A 248 249 represents general male choice regarding female cephalothorax width, in which positive values 250 represent higher general male choice for females with wider cephalothorax. Whereas, the slope B represents how male choice for wider female cephalothorax increases with male body mass, in 251 which positive values represent greater preference intensity for wider female cephalothorax as 252 male body mass increases. Similarly, in equation 2, the intercept D represents general male 253 254 choice regarding female recent pairing status, in which positive values represent higher 255 preference for recently unpaired females. The slope *E* represents how male choice for recently unpaired females increases with male body mass, in which positive values represent greater 256 257 preference intensity for recently unpaired females as male body mass increases. The relationship 258 between our alternative predictions and the values of these parameters is summarized in Table 1. Our analysis allowed coefficient values that would represent additional scenarios, not included in 259 260 the predictions. For example, it would be possible to find that all males do prefer females with 261 wider cephalothorax and that greater male body mass increases selectivity. However, we did not have a priori theoretical reasons to expect these other possibilities. 262

We implemented the model using the *stan* modelling language (Carpenter et al., 2017) and fit the models by Markov-Chain Monte-Carlo (MCMC) using a Bayesian framework in the software R 3.4.1 (R Core Team, 2017) using the package *rstan* (Stan Development Team, 2018). Prior to model fitting we standardized all continuous predictor variables (male body mass and female cephalothorax width) to zero mean and then divided by two times its standard deviation (following Gelman, 2008). Binary variables (recent pairing status and season) were set as 0 269 (recently paired females and 2017) and 1 (recently not paired females and 2018), respectively. 270 For each model, we ran three MCMC chains with 10,000 iterations each, plus 10,000 burn-in iterations. We adopted *Stan*'s standard uninformative improper priors. For more details on model 271 272 fitting, please see the tutorial in (Muniz et al., 2017). We evaluated convergence of the chains by inspecting R values (all < 1.01) and visually inspecting the chains for each parameter in our 273 model (Supplementary Figures 1 and 2). Because we centred and scaled all predictor variables, 274 coefficients can be interpreted as estimates of effect size. We considered coefficients to be 275 different than zero when 95% credible intervals (95% CI) did not overlap zero. 276

277 **Results**

In all trials, focal males successfully chose one of the two females available within 15 minutes. On average, males took 2.36 ± 3.16 minutes (mean \pm SD) to make this initial decision, depositing their own silk where they went. In the initial male mate choice trials, we found little evidence that male decision was influenced by female cephalothorax width, the previous presence of male partners or the experimental males' own body mass (Figure 1). This means that initial male decisions, upon encountering a female web, did not support any of our predictions.

In the late male choice trials, we found evidence that male pairing decision was influenced by female cephalothorax width, recent pairing status and male body mass (Figures 2 and 3). We found that greater than average male body mass increased the pairing probability with females of wider cephalothorax and that were unpaired before the trial (Figure 2). Additionally, smaller than average male body mass increased the pairing probability with females of narrower cephalothorax and that were paired before the trial (Figure 2). These results support our second prediction that male mate choice is assortative. We found little evidence of 291 differences in male mate choice when trials were conducted on different seasons (Figures 1 and292 2).

293

294 Discussion

In this study, we tested whether female quality and male fighting ability (*i.e.*, body size) would 295 influence male pairing decisions in the golden silk orb-weaver spider *Trichonephila clavipes*. 296 297 Using field experiments, we found little evidence that, on a first moment (*i.e.*, when males are 298 first exposed to silk threads of different female webs), males exert mate choice. However, after a period of a few hours, we found evidence that males express mate choice based on their own 299 fighting ability in an assortative manner. For instance, large males were more likely to guard the 300 301 larger female among those available. Also, when females differed in their recent pairing status, 302 large males preferred to guard the female that had not been paired to other male recently. Small 303 males, on the other hand, were more likely to guard the smaller female, and also, when given an 304 option, preferred a previously paired female to the one that was not recently guarded by other male. Medium-sized males, however, guarded females randomly regarding female size and 305 306 female previous pairing status. Therefore, our results indicate that there is variation in male mate 307 choice with small males showing preference patterns opposite to the large males, while mediumsized males showed no mating preferences. 308

Our experimental design allowed us to investigate whether male mate choice for female traits occurred shortly after males encountered a pair of females or whether males need more time to access females once they encounter them. We found differences in male mate choice along the duration of trials, in which males were initially indifferent to the female traits we evaluated, but later expressed mate choice depending on their own size. Males of many spider 314 species can access female information through silk strands produced by females (reviewed in Gaskett, 2007). For instance, Rittschof (2011) found, with laboratory experiments, that T. 315 *clavipes* males can distinguish females that are closer to oviposition using only cues present in 316 female silk. However, because we conducted our experiments in the field, males might have 317 been subjected to naturally occurring confounding factors that are absent in laboratory 318 experiments. For example, as we did not have information on when females built their webs, it is 319 possible that males chose more recently weaved threads due to their fresher chemical cues. 320 Furthermore, because females occasionally take over webs from other females (P.P. pers. obs.), 321 322 silk components may not be reliable cues to inform the quality of a resident female. Therefore, T. *clavipes* males may need time to evaluate female quality in a natural situation by gathering 323 information on female quality from vibrational cues or from cuticular pheromones (Robinson, 324 1982). This may explain why we found male mate choice only after males approached females 325 and had time to collect more information on their quality. Generally, animals gather information 326 on mate quality using multisensory cues (Rowe, 1999), but several mate choice investigations 327 allow only one sensory input information to focal individuals (e.g., insects: Goubault & Burlaud, 328 2017; fish: Gasparini, Serena, & Pilastro, 2013). Thus, our findings highlight the importance of 329 330 using experimental designs that assess choice in more than one moment and that allow males to receive multisensory female cues in a proper time frame. 331

Traditional mate choice literature predicts that when individuals express mate preference, they should invariably choose high quality mates (Andersson, 1994). Yet, empirical studies that found intra-population variation on mate choice have been accumulating (Jennions & Petrie, 1997). Among several factors that may promote mate choice variation, theoretical models (*e.g.*, Fawcett & Johnstone, 2003; Härdling & Kokko, 2005; Venner et al., 2010) identified that

337 intrasexual competition may be a critical influence on male mate choice variation. Here, we 338 found that T. clavipes males differ in the direction and intensity of their mate choice according to their own size. Only large males behaved according to what is generally predicted by the 339 340 traditional mate choice literature: these males chose larger females that had not been recently paired. Whereas small males preferred smaller females that had been recently paired to other 341 males. Why should these less competitive males choose lower quality females? If males 342 invariably choose to copulate with high quality females, these females would probably mate 343 multiply, and males would probably share the paternity of the eggs. Consequently, the advantage 344 345 of copulating with high quality females over low quality ones would be reduced. Thus, mate choice for higher quality females is only advantageous if the choosing male can effectively guard 346 the female and increase his likelihood of siring most of her offspring. In T. clavipes, guarding 347 efficiency is positively related to male size (Constant et al., 2011). Therefore, the larger the male, 348 the greater is the likelihood that he receives the benefits of mating with a high quality female. 349

Trichonephila clavipes males can guard only one female at a time, which means that 350 351 lower quality females would become available as large males are busy guarding high quality 352 females. Thus, the most profitable tactic for small males may be to choose lower quality females, 353 as a way to avoid competition with other males (*i.e.*, prudent mate choice; Härdling & Kokko, 2005). This rationale is supported by theoretical models that explore the influence of male 354 competitive ability on male mate choice, which usually focus on the extremes of male 355 356 phenotypes using two male categories (e.g., high/low quality in Fawcett & Johnstone, 2003; large/small in Härdling & Kokko, 2005; high/low competitive ability in Venner et al., 2010). Our 357 results on the choice expressed by males of extremes sizes (large/small) match the assortative 358 359 choice predicted by these theoretical studies under certain parameters. However, these models do

360 not clarify what is expected of average competitors. Here, we found that T. clavipes medium-361 sized males are unselective regarding female quality. Following the rationale that competitive ability determines whether males prefer or avoid high-quality females, medium-sized males 362 363 stand in the middle. That is, because medium-sized males are displaced by half of their competitors, they may gain the advantages of pairing with high-quality females sometimes, but 364 not frequently enough to promote preference nor rarely enough to promote avoidance to these 365 females. On the other hand, it would be interesting to explore whether males express consistent 366 preference for a particular female phenotype. This is because our findings that medium-sized 367 368 males are not selective towards female body size could also be explained by individual variation 369 among medium-sized males. Overall, our results match precisely the results of a theoretical study on mutual mate choice in a scenario in which females are weakly sperm limited, female mating 370 costs are low and male mating costs are high (see Figure 2 in Puurtinen & Fromhage, 2017). 371 Surprisingly, Puurtinen & Fromhage (2017) did not consider pre-copulatory aggression explicitly 372 in their model, revealing that our findings may also be justified with other features, such as 373 374 sperm competition.

We found that the smaller the male, the greater is the probability that he chooses females 375 376 that were recently being guarded by another male. It is possible that choosing recently paired females is a mechanism that decreases the risk of being displaced by larger males (in addition to 377 preferring small females). Males using this strategy would copulate mostly with non-virgin 378 379 females, and therefore would share the paternity of the offspring. This choice for recently paired females could indicate that smaller males tend to employ a sneaking tactic, in which the male 380 stays on the periphery of a guarded female's web and attempts to copulate unnoticed by the 381 guarding male (Christenson & Goist, 1979). In this scenario, smaller males would be interpreting 382

383 the cues of male presence as a sign that the female is currently being guarded, and would be choosing to be sneakers rather than guardians. Although we do not know how successful this 384 strategy can be, small males appear to get less attention from guarding males when compared to 385 large males (P.P. pers. obs.). In any case, males are apparently employing different mating tactics 386 depending on their own body size. Here, we only observed their pairing decisions, but males of 387 different fighting abilities may also differ in tactics regarding female search, sperm allocation, 388 and mate guarding tactics. Small males may compensate the lower quality of their partners by 389 being more promiscuous. If that is the case, we would expect that male size is also (i) negatively 390 391 related to movement between female webs, as a strategy to copulate with more females, and (ii) negatively related to investment in each female, in terms of sperm allocation and mate-guarding 392 time. 393

Our results show that males similarly consider female size and female social context 394 when choosing their mates. The combination of these two variables makes it difficult to estimate 395 female quality in certain scenarios (e.g., a recently paired large female may be of similar quality 396 397 to a not recently paired small female). Hence, a simple assortative pairing pattern by body size alone may not necessarily occur in natural conditions. Furthermore, our results indicate that all 398 399 females, regardless of size, are able to attract males to mate, although it seems that they attract more males of a particular fighting ability than randomly expected. Thus, even though male mate 400 choice can generate sexual selection on females and even favour the evolution of female sexual 401 402 ornaments (e.g., Amundsen & Forsgren, 2003), variation in male mate choice may attenuate sexual selection on females, especially if some males perform prudent mate choice. Therefore, 403 scenarios of variable male mate choice may be common, but hard to detect. Detection of 404 405 variation in male mate choice is made difficult by two main reasons: (1) it generates weak or no

406 sexual selection on females and (2) it requires specific experimental and statistical protocols (as 407 the ones we employed here). We argue that variation in male mate choice may be relatively common, as theory predicts that it will evolve in populations in which mating is costly for males 408 409 and there is strong intrasexual competition among males (Fawcett & Johnstone, 2003; Härdling & Kokko, 2005; Venner et al., 2010). These conditions occur in *T. clavipes*, and may also be met 410 by other species in which males cannot monopolize groups of females, so that these males are 411 continuously searching for new mates, or in mating systems in which males can guard a single 412 female at a time. Such non-territorial mating systems are less studied than territorial ones, but are 413 414 the most common among animals (Herberstein, Painting, & Holwell, 2017). Therefore, future studies employing an experimental protocol similar to ours may discover that variation in male 415 mate choice is very common in natural populations. 416

417

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427 Authors' contributions

428	P.P	and E.S.A.S.	conceived	the	idea	of	the	study,	P.P.	collected	the	data,	P.P.	and	D.G.M.

analysed the data, P.P., D.G.M and E.S.A.S. wrote the manuscript.

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431 Data accessibility

432 Data and analysis scripts are available at http://osf.io/6n7kt/.

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434 **References**

Amundsen, T., & Forsgren, E. (2003). Male preference for colourful females affected by male
size in a marine fish. *Behavioral Ecology and Sociobiology*, *54*(1), 55–64.
doi:10.1007/s00265-003-0593-4

438 Andersson, M. (1994). *Sexual selection*. Princeton, New Jersey: Princeton University Press.

Andrade, M. C. B. (1996). Sexual selection for male sacrifice in the Australian redback spider. *Science*, 271(5245), 70–72. doi:10.1126/science.271.5245.70

Arnaud, L., & Haubruge, E. (1999). Mating behaviour and male mate choice in *Tribolium castaneum* (Coleoptera, Tenebrionidae). *Behaviour*, *136*(1), 67–77.
doi:10.1163/156853999500677

Austad, B. Y. S. N. (1983). A game theoretical interpretation of male combat in the bowl and
doily spider (*Frontinella pyramitela*). *Animal Behaviour*, *31*, 59–73. doi:10.1016/S0003-

- Bel-Venner, M. C., Dray, S., Allainé, D., Menu, F., & Venner, S. (2008). Unexpected male
 choosiness for mates in a spider. *Proceedings of the Royal Society B: Biological Sciences*,
 275(1630), 77–82. doi:10.1098/rspb.2007.1278
- Berglund, A., Rosenqvist, G., & Svensson, I. (1989). Reproductive success of females limited by
 males in two pipefish species. *The American Naturalist*, 133(4), 506–516.
 doi:10.1086/284932
- Bonduriansky, R. (2001). The evolution of male mate choice in insects: a synthesis of ideas and
 evidence. *Biological Reviews*, *76*(3), 305–339. doi:10.1017/s1464793101005693
- Brown, S. G. (1985). Mating behavior of the golden-orb-weaving spider, *Nephila clavipes*: II.
- 456 Sperm capacitation, sperm competition, and fecundity. *Journal of Comparative Psychology*,
- 457 99(2), 167–175. doi:10.1037/0735-7036.99.2.167
- Carpenter, B., Gelman, A., Hoffman, M. D., Lee, D., Goodrich, B., Betancourt, M., ... Riddell,
 A. (2017). Stan: a probabilistic programming language. *Journal of Statistical Software*,
 76(1), 1–32. doi:10.18637/jss.v076.i01
- 461 Christenson, T. E. (1989). Sperm depletion in the orb-weaving spider *Nephila clavipes* (Araneae,
 462 Araneidae). *Journal of Arachnology*, *17*(1), 115–118.
- 463 Christenson, T. E., & Cohn, J. (1988). Male advantage for egg fertilization in the golden orb-

464 weaving spider (*Nephila clavipes*). Journal of Comparative Psychology, 102(4), 312–318.
465 doi:10.1037/0735-7036.102.4.312

- Christenson, T. E., & Goist, K. C. (1979). Costs and benefits of male-male competition in the orb
 weaving spider, *Nephila clavipes. Behavioral Ecology and Sociobiology*, 5(1), 87–92.
 doi:10.1007/bf00302697
- Constant, N., Valbuena, D., & Rittschof, C. C. (2011). Male contest investment changes with
 male body size but not female quality in the spider *Nephila clavipes*. *Behavioural Processes*, 87(2), 218–223. doi:10.1016/j.beproc.2011.04.003
- 472 Cratsley, C. K., Rooney, J. A., & Lewis, S. M. (2003). Limits to nuptial gift production by male
 473 fireflies, *Photinus ignitus. Journal of Insect Behavior*, 16(3), 361–370.
 474 doi:10.1023/A:1024876009281
- 475 Dewsbury, D. A. (1982). Ejaculate cost and male choice. *The American Naturalist*, *119*(5), 601–
 476 610. doi:10.2307/2461181
- 477 Dewsbury, D. A. (2005). The Darwin-Bateman paradigm in historical context. *Integrative and*478 *Comparative Biology*, 45(5), 831–837. doi:10.1093/icb/45.5.831
- 479 Dougherty, L. R., & Shuker, D. M. (2015). The effect of experimental design on the
 480 measurement of mate choice: a meta-analysis. *Behavioral Ecology*, 26(2), 311–319.
 481 doi:10.1093/beheco/aru125

- Edward, D. A., & Chapman, T. (2011). The evolution and significance of male mate choice. *Trends in Ecology & Evolution*, 26(12), 647–654. doi:10.1016/j.tree.2011.07.012
- Fawcett, T. W., & Johnstone, R. A. (2003). Mate choice in the face of costly competition. *Behavioral Ecology*, 14(6), 771–779. doi:10.1093/beheco/arg075
- Gaskett, A. C. (2007). Spider sex pheromones: emission, reception, structures, and functions. *Biological Reviews*, 82(1), 27–48. doi:10.1111/j.1469-185X.2006.00002.x
- 488 Gasparini, C., Serena, G., & Pilastro, A. (2013). Do unattractive friends make you look better?
- 489 Context-dependent male mating preferences in the guppy. *Proceedings of the Royal Society*
- 490 *B: Biological Sciences*, 280(1756), 20123072. doi:10.1098/rspb.2012.3072
- Gelman, A. (2008). Scaling regression inputs by dividing by two standard deviations. *Statistics in Medicine*, 27, 2865–2873. doi:10.1002/sim.3107
- 493 Goubault, M., & Burlaud, R. (2017). Do males choose their mates in the lekking moth Achroia
- 494 *grisella*? Influence of female body mass and male reproductive status on male mate choice.
- 495 Insect Science, 25(5), 861–868. doi:10.1111/1744-7917.12471
- Gwynne, D. T. (1981). Sexual difference theory: mormon crickets show role reversal in mate
 choice. *Science*, *213*(4509), 779–780. doi:10.1126/science.213.4509.779
- 498 Härdling, R., & Kokko, H. (2005). The evolution of prudent choice. *Evolutionary Ecology*499 *Research*, 7(5), 697–715. doi:10.1046/j.1461-0248.1999.00075.x

500	Head, G. (1995). Selection on fecundity and variation in the degree of sexual size dimorphism
501	among spider species (class Araneae). Evolution, 49(4), 776–781. doi:10.2307/2410330

- 502 Herberstein, M. E., Painting, C. J., & Holwell, G. I. (2017). Scramble competition polygyny in
- 503 terrestrial arthropods. In M. Naguib, J. Podos, L. W. Simmons, L. Barrett, S. D. Healy, &
- M. Zuk (Eds.), Advances in the study of behavior (Vol. 49, pp. 237–295). Cambridge,
 Massachusetts: Elsevier. doi:10.1016/bs.asb.2017.01.001
- Honěk, A. (1993). Intraspecific variation in body size and fecundity in insects: a general
 relationship. *Oikos*, 66(3), 483–492. doi:10.2307/3544943
- Jennions, M. D., & Petrie, M. (1997). Variation in mate choice and mating preferences: a review
 of causes and consequences. *Biological Reviews*, 72, 283–327. doi:10.1111/j.1469185X.1997.tb00015.x
- Kasumovic, M. M., Bruce, M. J., Herberstein, M. E., & Andrade, M. C. B. (2007). Risky mate
 search and mate preference in the golden orb-web spider (*Nephila plumipes*). *Behavioral Ecology*, 18(1), 189–195. doi:10.1093/beheco/arl072
- Kelly, C. D. (2006). Fighting for harems: assessment strategies during male e male contests in
 the sexually dimorphic Wellington tree weta. *Animal Behaviour*, 72, 727–736.
 doi:10.1016/j.anbehav.2006.02.007
- 517 Kuntner, M., Hamilton, C. A., Cheng, R.-C., Gregorič, M., Lupše, N., Lokovšek, T., ... Bond, J.
 518 E. (2018). Golden orbweavers ignore biological rules: phylogenomic and comparative

- analyses unravel a complex evolution of sexual size dimorphism. *Systematic Biology*.
 doi:10.1093/sysbio/syy082
- Lane, J. E., Boutin, S., Speakman, J. R., & Humphries, M. M. (2010). Energetic costs of male
 reproduction in a scramble competition mating system. *Journal of Animal Ecology*, *79*(1),
 27–34. doi:10.1111/j.1365-2656.2009.01592.x
- Mautz, B. S., & Jennions, M. D. (2011). The effect of competitor presence and relative
 competitive ability on male mate choice. *Behavioral Ecology*, 22(4), 769–775.
 doi:10.1093/beheco/arr048
- Michalik, P., & Rittschof, C. C. (2011). A comparative analysis of the morphology and evolution
 of permanent sperm depletion in spiders. *Plos One*, 6(1), e16014.
 doi:10.1371/journal.pone.0016014
- Muniz, D. G., Santos, E. S. A., Guimarães, P. R., Nakagawa, S., & Machado, G. (2017). A
 multinomial network method for the analysis of mate choice and assortative mating in
 spatially structured populations. *Methods in Ecology and Evolution*, 8(10), 1321–1331.
 doi:10.1111/2041-210X.12798

Parker, G. A., & Pizzari, T. (2015). Sexual selection: the logical imperative. In T. Hoquet (Ed.),
 Current perspectives on sexual selection (pp. 119–164). Amsterdam: Springer Netherlands.
 doi:10.1007/978-94-017-9585-2_7

537 Perry, J. C., Sirot, L., & Wigby, S. (2013). The seminal symphony: how to compose an ejaculate.

Trends in Ecology & Evolution, 28(7), 414–422. doi:10.1016/j.tree.2013.03.005

- Puurtinen, M., & Fromhage, L. (2017). Evolution of male and female choice in polyandrous
 systems. *Proceedings of the Royal Society B: Biological Sciences*, 284, 20162174.
 doi:10.1098/rspb.2016.2174
- 542 R Core Team. (2017). R: a language and environment for statistical computing. Vienna, Austria:
 543 R Foundation for Statistical Computing. Retrieved from https://www.r-project.org/
- Rittschof, C. C. (2011). Mortality risk affects mating decisions in the spider *Nephila clavipes*. *Behavioral Ecology*, 22(2), 350–357. doi:10.1093/beheco/arq222
- Robinson, M. H. (1982). Courtship and mating behavior in spiders. *Annual Review of Entomology*, 27, 1–20. doi:10.1146/annurev.en.27.010182.000245
- 548 Robinson, M. H., & Mirick, H. (1971). The predatory behavior of the golden-web spider *Nephila*
- 549 *clavipes* (Araneae: Araneidae). *Psyche*, 78(3), 123–139. doi:10.1155/1971/57182
- Rowe, C. (1999). Receiver psychology and the evolution of multicomponent signals. *Animal Behaviour*, 58, 921–931. doi:10.1006/anbe.1999.1242
- Schneider, J. M., Lucass, C., Brandler, W., & Fromhage, L. (2011). Spider males adjust mate
 choice but not sperm allocation to cues of a rival. *Ethology*, *117*(11), 970–978.
 doi:10.1111/j.1439-0310.2011.01960.x
- 555 Schneider, J. M., Zimmer, S. M., Gatz, A. L., & Sauerland, K. (2016). Context- and state-

dependent male mate choice in a sexually cannibalistic spider. *Ethology*, *122*(3), 257–266.
doi:10.1111/eth.12466

Schwagmeyer, P. L., & Parker, G. A. (1990). Male mate choice as predicted by sperm
competition in thirteen-lined ground squirrels. *Nature*. doi:10.1038/348062a0

560 Stan Development Team. (2018). RStan: the R interface to Stan. Retrieved from http://mc561 stan.org/

Tang-Martinez, Z. (2016). Rethinking Bateman's principles: challenging persistent myths of
sexually reluctant females and promiscuous males. *The Journal of Sex Research*, 53(4–5),
532–559. doi:10.1080/00224499.2016.1150938

- Tang-Martinez, Z., & Ryder, T. B. (2005). The problem with paradigms: Bateman's worldview
 as a case study. *Integrative and Comparative Biology*, 45(5), 821–830.
 doi:10.1093/icb/45.5.821
- Venner, S., Bernstein, C., Dray, S., & Bel-Venner, M.-C. (2010). Make love not war: when
 should less competitive males choose low-quality but defendable females? *The American Naturalist*, 175(6), 650–661. doi:10.1086/652432
- Vollrath, F. (1980). Male body size and fitness in the web-building spider *Nephila clavipes*. *Zeitschrift Für Tierpsychologie*, 53(1), 61–78. doi:10.1111/j.1439-0310.1980.tb00733.x
- 573 Vollrath, F., & Parker, G. A. (1992). Sexual dimorphism and distorted sex-ratios in spiders.

- 574 *Nature*, *360*(6400), 156–159. doi:10.1038/360156a0
- Wada, S., Arashiro, Y., Takeshita, F., & Shibata, Y. (2010). Male mate choice in hermit crabs:
 prudence by inferior males and simple preference by superior males. *Behavioral Ecology*,
 22(1), 114–119. doi:10.1093/beheco/arq183
- Wedell, N., Gage, M. J. G., & Parker, G. A. (2002). Sperm competition, male prudence and
 sperm-limited females. *Trends in Ecology & Evolution*, *17*(7), 313–320.
 doi:10.1016/S0169-5347(02)02533-8

582 **Figures and tables**

Table 1. Summary of the predictions investigated in this study and their relationship with model parameter values. Columns two, three, and four represent male selectivity regarding female cephalothorax width and female recent pairing status according to male body mass. Positive selectivity represents preference for recently unpaired females with wider cephalothorax, zero selectivity represents no preference and negative selectivity represents preference for recently paired females with narrower cephalothorax. The fifth and sixth columns represent the expected values of model parameters according to each alternative prediction (see text for details).

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Prediction	Selectivity of high body mass males	Selectivity of medium body mass males	Selectivity of small body mass males	Coefficients A and D	Coefficients B and E
1 - all males equally choosy	Positive	Positive	Positive	Positive	Zero
2 - variable male mate choice	Positive	Zero	Negative	Zero	Positive



Figure 1. Results of the multinomial model investigating initial male mate choice (*i.e.*, which female thread the experimental male climbed) in *Trichonephila clavipes*. Points and segments represent mean estimated parameter value and 95% credible interval. See details about what each parameter represents in the *Statistical analysis of male mate choice experiment* section.



Figure 2. Results of the multinomial model to investigate late male mate choice (*i.e.*, which female the focal male was guarding after 16 to 22 hours of the beginning of the experiment) in *Trichonephila clavipes*. Points and segments represent mean estimated parameter value and 95% credible interval. Stars highlight the model parameters for which the 95% credible interval did not overlap zero. See details about what each parameter represents in the *Statistical analysis of male mate choice experiment* section.



Focal female standardized cephalothorax width (2σ)

Figure 3. Pairing probability of a male with a focal female depending on male body mass and 606 focal female cephalothorax width. In this scenario, a male may choose between the focal female 607 and another female of standardized cephalothorax width of zero value (*i.e.*, an average-sized 608 609 female). Moreover, females do not differ in previous pairing status in this scenario. The probability that a high body mass male (solid line) will pair with the focal female increases as 610 focal female cephalothorax width increases. In contrast, the probability that a low body mass 611 612 male (dotted line) will pair with the focal females decreases as focal female cephalothorax width increases. The probability that medium body mass males (dashed line) will pair with the focal 613 female does not depend on focal female cephalothorax width and is always 50% (random). We 614

- used coefficient values in which 95% credible interval did not overlap zero in the late male mate
- 616 choice model to estimate the predicted values displayed in this figure.