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Abstract:	Males sometimes engage in fights over contested resources such as access to mates; in this case, fighting behaviour may be adjusted based on the value they place on the females. Resource value RV can have two components. Males can assess the quality of females, which constitutes an objective assessment of RV. Internal state such as previous mating experience can also influence motivation to fight thus constituting a subjective assessment of RV. If mating opportunities are scarce and available females have a major impact on the lifetime reproductive success of males, then fighting can be fatal; in this situation it is uncertain whether males would adjust fighting behaviour based on RV. We found that both female quality i.e., virginity (objective component of RV) and male mating status (subjective component of RV) influence fighting intensity between males of the entomopathogenic nematode Steinernema longicaudum which engage in lethal fights. Male nematodes were more likely to engage in fighting and fought longer and more frequently in the presence of virgin (high quality) females than in the presence of mated (lower quality) females. Male mating status was also found to influence fighting behaviour; mated males may have superior fighting ability (enhanced resource holding potential RHP), but RV asymmetries between mated and virgin males cannot be excluded. Males were more likely to win when they were resident, but we did not find a significant interaction effect between male mating and residency status.	

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Objective and subjective components of resource value in lethal fights between male entomopathogenic nematodes

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Running title: *female quality and male mating status influence fighting behaviour in male nematodes*

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4 ABSTRACT

5 Males sometimes engage in fights over contested resources such as access to mates; in this 6 case, fighting behaviour may be adjusted based on the value they place on the females. 7 Resource value RV can have two components. Males can assess the quality of females, which constitutes an objective assessment of RV. Internal state such as previous mating experience 8 9 can also influence motivation to fight thus constituting a subjective assessment of RV. If mating opportunities are scarce and available females have a major impact on the lifetime 10 reproductive success of males, then fighting can be fatal; in this situation it is uncertain 11 whether males would adjust fighting behaviour based on RV. We found that both female 12 quality i.e., virginity (objective component of RV) and male mating status (subjective 13 component of RV) influence fighting intensity between males of the entomopathogenic 14 nematode Steinernema longicaudum which engage in lethal fights. Male nematodes were 15 more likely to engage in fighting and fought longer and more frequently in the presence of 16 17 virgin (high quality) females than in the presence of mated (lower quality) females. Male 18 mating status was also found to influence fighting behaviour; mated males were the winners in staged fights between mated and virgin males. Mated males may have superior fighting 19 20 ability (enhanced resource holding potential RHP), but RV asymmetries between mated and virgin males cannot be excluded. Males were more likely to win when they were resident, but 21 22 we did not find a significant interaction effect between male mating and residency status. Key words: entomopathogenic nematodes, female quality, lethal fights, male mating status, 23 resource value. 24

26 INTRODUCTION

Animals frequently engage in contests over resources that impact their fitness, such as mates, 27 territories or nesting sites (Andersson 1994). Amongst the most important factors influencing 28 the extent and outcome of contest behaviour are the resource holding potential of the 29 contestants, the value of the resource and the costs of fighting (Parker 1974; Arnott & Elwood 30 31 2008; Hardy & Briffa 2013). Resource holding potential RHP (fighting ability) of the contestants may depend on their weaponry, energy reserves and body size (Parker 1974; Arnott 32 & Elwood 2009; Hardy & Briffa 2013; Rico-Guevara & Hurme 2019). Resource value RV has 33 two components: objective and subjective; Intrinsic properties of the resource that will yield a 34 certain fitness gain to a successful competitor constitute the objective components of resource 35 36 value, while the value of a given resource may vary for each contestant depending on its own internal state and prior experience (subjective component of resource value) (Enquist & Leimar 37 1987; Arnott & Elwood 2008). These components of resource value may simultaneously 38 39 influence the ultimate value of the resource (Stockermans & Hardy 2013).

Objective and subjective components of resource value have been widely studied in 40 contests between males for mates, where the contested resource is receptive females. When 41 42 females are scarce, males may be more motivated to fight, but this motivation can also depend on female quality which can be based on size (larger females are more fecund), nearness to 43 44 moult or mating status (Crespi 1988; Dick & Elwood 1990; Arnott & Elwood 2008; Keil & Watson 2010; Hoefler et al. 2009; Ancona et al. 2010; Kasumovic et al. 2011). Furthermore, 45 males' internal state, such as their own mating status, can influence subjectively the value that 46 they place on the contested resource (i.e., mates) (Arnott & Elwood 2008). Mated males may 47 become less aggressive and more likely to lose at contests due to the energetic cost of mating 48 (Brown et al. 2006, 2007; Judge et al. 2010). On the other hand, previous mating experience 49 50 has been shown to increase a male's aggression and fighting success in several animal taxa

because of changes in the male's self-assessment of RHP and perception of the resource 51 52 value, that are similar to winner/loser effects (Killian and Allen 2008; Yasuda et al. 2015, Dugatkin & Dugatkin 2011; Zhang et al., 2019). If an animal wins a contest then its 53 perception of fighting ability increases and it is more likely to initiate future fights and win 54 them, whereas losers lower their RHP perception, are less aggressive and more likely to lose 55 (Hsu et al. 2006). Mating status is also related to the residual reproductive value (number of 56 57 future offspring) which is predicted to influence male investment in fighting (Kemp 2006). An unmated male can be more aggressive and more motivated to fight because future mating 58 opportunities are of greater value to him, whereas a mated male may be less willing to engage 59 60 in costly fighting (Kemp 2006).

Fighting is costly, risking injury and using time and energy that could be used in 61 mating or other activities (Maynard Smith & Price 1973; Briffa & Sneddon 2007). As the 62 63 value of the contested resource increases, the cost of fighting can also increase (Enquist & Leimar 1987; Kokko 2013). If the contested resource has a major impact on the lifetime 64 65 reproductive success of the the contenstants, such as limited females, then fights can escalate to serious injury and even death (Enquist & Leimar 1987, 1990; Maynard Smith & Parker 66 1976). Lethal male fighting has been recorded in some vertebrates (e.g., Piper et al. 2008) and 67 68 numerous arthropods, particularly arachnids (DeCarvalho et al. 2004; Sato et al. 2013) and hymenopterans (Anderson et al. 2003; Matthews et al. 2009; Cook & Bean 2006; Hamilton 69 1979; Innocent et al. 2007; Innocent et al. 2011) and lately in entomopathogenic nematodes 70 (Zenner et al. 2014; Kapranas et al. 2016). Whereas contested resource availability, such as 71 72 potential mates, is critical in prompting lethal fights (Enquist & Leimar 1990), it is unclear to what extent the contestants also assess the quality of resource in such species. When mating 73 74 opportunities for males are extremely limited then their motivation to fight would be expected to be independent of resource quality even if the odds of dying in a fight are high (Enquist & 75

Leimar 1990). Amongst parasitoid hymenopterans with lethal male combat, some species
adjust fighting behaviour according to the value of the resource (Liu & Hao 2019), while
others do not (Innocent et al. 2011).

79 Here we investigate the effect of objective and subjective resource value on lethal contests of male entomopathogenic nematodes Steinernema longicaudum Shen and Wang. 80 We are able to manipulate the mating status of the contestant males and also the quality of the 81 82 contested resource by staging dyadic contests in a controlled environment (drops of haemolymph of their insect host). First, we assess how the presence/absence and quality of 83 available females (objective components of resource value) influence fighting behaviour 84 85 between mated males. We predict that the presence of a female, especially a virgin, will increase the frequency and/or severity of fighting in S. longicaudum. Then, we assess whether 86 male mating status (subjective components of resource value) influences the contest outcome. 87 We do not have a directional prediction of how mating influences fighting, but we 88 hypothesize that either mated or virgin males will have a consistent advantage over the other, 89 90 and this will provide insights regarding how male nematodes assess the value of their mates 91 and whether mating status influences fighting ability.

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93 MATERIALS AND METHODS

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95 Nematode Reproductive Biology and Lethal Fighting.

96 Infection by entomopathogenic nematodes including *S. longicaudum* begins when free-97 foraging infective juveniles (IJs) enter an insect host and release their symbiotic bacteria that 98 turn the host insect into a cadaver. In *S. longicaudum*, IJs develop into amphimictic males and 99 females, with a sex ratio that is slightly female biased (Alsaiyah et al. 2009). Two or more 100 generations develop until the cadaver is crowded, prompting the production of IJs that exit in

search of other hosts (Kaya & Gaugler 1993; Dillman & Sternberg 2012). In S. longicaudum 101 it was shown that males are aggressive towards each other and engage in lethal fights (Zenner 102 et al. 2014). During male-male encounters, an aggressor coils its tail end around a victim, with 103 104 the copulatory spicules towards the victim's body. If not gripped close to the tail, the victim may counter-attack, wrapping its tail around the aggressor. Grappling encounters frequently 105 end when the grasped male ceases to move part or all of its body, which may happen within 106 107 minutes (Zenner et al. 2014). Such worms usually die, though partially paralyzed worms 108 occasionally recover movement. The cuticle is sometimes punctured, but paralysis and death occur also without puncture, apparently due to damage to internal organs. 109

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111 Nematode Cultures and Behavioural Assays

112 Steinernema longicaudum (strain CB2B) cultures were routinely maintained using standard 113 procedures by passage through late instar Galleria mellonella (wax moth) larvae (Kaya & Stock 1997) at 27°C. Infective juveniles were stored in tap water at 20°C. We obtained adult 114 115 males for our experiments by placing IJs individually in 25 µl hanging drops of haemolymph 116 from G. mellonella larvae which provides a suitable environment for development to adulthood (Kaya & Stock 1997; Zenner et al. 2014). The hanging drops were placed on the lid 117 118 of a Petri dish which was then inverted over a water-filled Petri dish (6 cm diam.) to prevent 119 desiccation. The IJs in the hanging drops develop to adulthood within 3 days at 27°C. Adult males can be distinguished by their copulatory spicules and their smaller size whereas females 120 are larger in size and bear vulva. We used three days old males and females in our 121 experiments. 122

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124 Experiment 1: Effects of Female Mating Status

All males used in this experiment were mated. We placed each individual naïve male for 16-125 126 20 hours with 2 naïve females to mate. We then removed the females, and successful mating was subsequently confirmed by the production of progeny. We removed two adult males from 127 their hanging drops and placed them immediately, in pairs, back in the drop from which one 128 member of the pair had been taken. Thus, one worm was "resident" and one an "intruder", but 129 130 the identity of each was not tracked, as there were no clear visual differences between them. 131 Worms were assigned to the following treatments: (i) 2 mated males with a mated female (N= 34); (ii) 2 mated males with a virgin female (same age as mated female) (N = 31); (iii) 2 132 mated males without a female (N = 35). A mating attempt was recorded when a male coiled 133 134 around the female at the vulva; insertion of spicules into the vulva was difficult to observe due to the coiling of the male around the female. A fight was recorded once one male coiled 135 tightly around another and then let go. Continuous observations were made for 30 mins during 136 137 which the latency to first fight, the incidence (whether at least one fighting event occurred), number and (for a subsample) duration of fights were recorded. Paralysis and/or death of the 138 males was recorded at the end of the observation and after 24 hours. 139

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141 *Experiment 2: Effects of Male Mating Status*

142 In S. longicaudum, males that have been with a female can be distinguished from those that have not, based on the presence of sperm in the seminal vesicle, visible through the body wall 143 (Ebssa et al., 2008); thus, mating status of individual males in a pair could be recognized. In 144 this experiment, we placed together an adult male that had mated and a virgin one. Mated 145 males were obtained by placing two females in the drop of an individually reared adult male 146 for 24 hours, after which the females were removed to a different drop and observed for later 147 progeny production. We set up pairs of one mated and one virgin adult male either in the drop 148 in which the mated male had been reared and mated (N = 21) or in the drop in which the 149

unmated male had been reared (N = 24). In this case, since mated and unmated males differed in appearance, the identity of "resident" and "intruder" could also be traced. After 24 hours we observed which worms showed signs of paralysis or death. Individual males were identified as the mated or virgin member of the pair by examining for the presence of sperm using a Nikon Optiphot microscope (x40).

156 *Statistics*

Analysis of factors influencing the number of fights, as well copulations observed in our 157 assays and the probability of paralysis/death were explored with generalized linear models as 158 described in Briffa et al. (2013). To test the hypothesis that objective resource value affects 159 fighting behavior, we used a log-linear analysis which is appropriate for small count data, to 160 assess how female presence and mating status (predictor variables) affected the number of 161 162 fights (dependent variable). Post-hoc tests among different treatments whenever applicable were adjusted with the Bonferroni correction. The probability of paralysis and/or death at the 163 end of the 30 min observation period and 24h post observations was explored with logistic 164 analysis. Duration of fighting in different treatments (no females, mated and virgin females) 165 was analysed with a non-parametric Kruskal Wallis test and latency to fighting was analysed 166 using survival regression analysis (Moya-Larano & Wise 2000). In the second experiment, we 167 compared the incidence of paralysis/death in contests staged in drops where the resident male 168 was mated versus drops where the resident was a virgin male, by using a Fisher's exact test. 169 For those pairs in which paralysis or death occurred, we used a logistic analysis by randomly 170 picking a focal male (either mated or virgin) in each replicate and tested the effect of 171 residency and mating status (as factors) and their interaction on the probability of the focal 172 male winning the contest (Briffa et al. 2013). All analysis was performed in SPSS v.21 (IBM, 173 Armonk, NY, USA). 174

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176 **RESULTS**

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178 Female Presence and Quality

Significantly more fights were observed in the presence of virgin females than in the presence 179 of mated females or in the absence of females ($F_{2.97} = 5.79$, P = 0.004, Fig. 1). This was partly 180 explained by the higher incidence of fighting in the presence of a virgin female (74.2%) than 181 with mated or no female (44.1 and 42.9%, respectively). In addition, the latency to fight was 182 shorter in the presence of virgin females than in the presence of mated females or in the absence 183 184 of females (Cox's proportional hazards analysis: Risk = 0.448, 95% CI 0.233-0.860, P = 0.016, Fig. 2), and fights also tended to last longer when there was a virgin female present (Kruskal 185 Wallis *H* = 5.56, DF= 2, *P* = 0.062; Fig. 3). 186

The presence or quality of a female did not influence the incidence of paralysis at the end of the 30 min observation period (G_2 =1.437, P=0.487) when on average 6% of pairs had one male paralysed, nor did it affect the incidence of paralysis and/or death after 24 hours (G_2 =2.839, P= 0.242), where overall 97% of pairs had at least one male either paralysed or dead, mostly represented by dead (90.9% of pairs). Single males routinely suffer 3% mortality within 24 hours (Zenner et al. 2014).

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Moreover, the number of copulations (mating events at vulva) was higher when the female was virgin than mated (mated females: 0.88 ± 0.18 , virgin females: 1.74 ± 0.26 , $F_{1, 63} = 7.322$, P = 0.009).



200 After 24 h, paralysis or death was recorded in all (N = 21) drops where the resident male was mated, but in just 70.8% (17/24) of the drops where the virgin male was resident (Fisher's exact 201 test, P = 0.01). For the 38 pairs where there was paralysis or death after 24 h, we investigated 202 203 whether male mating status and prior residency affected the contest outcome. Mated and resident males were more likely to win (logistic regressions for male mating status: $G_{1,34} =$ 204 75.13, P < 0.001; male residency: $G_{1,34} = 10.313$, P < 0.001) (Figure 4). The effect of male 205 206 mating status was particularly strong: mated males won 92.1% of fights overall. The interaction 207 between male mating status and drop residency was not significant ($G_{1,34} = 0$, P = 0.991).

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210 **DISCUSSION**

211 The presence of a female can lead to escalated male fighting, as it has shown in spiders and parasitoid wasps (Wells 1988; Jackson et al. 2006; Liu et al. 2017). Moreover, the 212 motivation to fight and intensity of fights, and consequently the cost of fighting, is adjusted 213 according to the perceived value of the resource (Parker 1974; Maynard- Smith & Parker 214 1976; Arnott & Elwood 2008; Enquist & Leimar 1987). In our study we found that not only 215 the presence but also the quality of females had an influence on fighting behaviour of male S. 216 217 *longicaudum* nematodes; fighting was more intense in the presence of a virgin female, during our 30 min observation period. In other empirical studies, males similarly exhibited 218 significantly more aggressive behaviours, and fights were more intense in the presence of 219 220 virgin females over mated ones in parasitoid wasps (Liu & Hao 2019), wolf spiders (Hoefler et al. 2008) and Sierra dome spiders (Keil & Watson 2010). The fact that a virgin (but not a 221 mated) female enhances competition between male S. longicaudum suggests either that 222 223 females mate only once, or that there is sperm competition with first male precedence (Birkhead & Moller 1998; Simmons 2001). Sperm precedence has been documented in 224

nematodes including Caenorhabditis elegans (LaMunyon & Ward 1995; Ward & Carrel 225 226 1979). Mated female steinernematids no longer attract males over a distance (Lewis et al. 2002; Hartley 2017), and in our experiment mated females received fewer mating attempts 227 (sperm transfer was not confirmed) than virgins, suggesting at least a reduction in 228 attractiveness. Female steinernematids have at most a short window of availability for mating, 229 with a period of egg-laving succeeded by egg-hatch in utero and subsequent death ("endotokia 230 231 matricida") (Poinar 1990). The exceptionally large macrosperm of S. longicaudum are suggestive of intense sperm competition (LaMunyon & Ward 1999). Because the cost of 232 fighting is high, selection favours individuals that can assess the value of the resource and 233 234 adjust their competitive behaviour accordingly (Parker 1974; Maynard-Smith & Parker 1976; Arnott & Elwood 2008). When mates are limited over time and space and their value is equal 235 236 or higher than the future, then a major part of a male contestant's lifetime reproductive success 237 is at stake and it is predicted that fights can escalate to death (Enquist & Leimar 1987, 1990). In such cases, fights might ensue irrespective of the quality of the mates, since future 238 239 opportunities for reproduction are low, but our results show that male S. longicaudum 240 nematodes assess the quality of the females and adjust their fighting behaviour. We suggest 241 that mated females, due to either once-off mating or intense sperm competition, might simply 242 not represent a valuable resource worth fighting to death for.

Differences in the quantity and quality of food or mates lead to an 'objective' assessment of the value of the contested resource, whereas internal state dictates the subjective assessment of the resource value (Enquist & Leimar 1987; Arnott & Elwood 2008). In dyadic fights between mated *vs* virgin males, the mated males were more competitive, resulting in paralysis and death of the virgin male in over 90% of cases. Mating could enhance fighting skills in nematodes, since similar coiling movements of males around females during copulation are used to lock, paralyse and consequently lead to death of their opponents.

250 Increased fighting skills enhance resource holding potential (Briffa & Lane 2017).

Alternatively, mating in males can result in an increased assessment of fighting ability or an
overestimation of the contested resource value (Killian & Allen 2008; Yasuda et al. 2015).
Mating experience has similar effects to 'winning a fight' experience, which is expected in
turn to raise the subjective value of a female in future contests (Hsu et al. 2006).

255 Contests are frequently asymmetrical; one such asymmetry is between territory owners or residents, and intruders. In such cases, it is usually the resident that wins (Hardy & Briffa 256 2013). Owner-intruder asymmetry is less likely to affect fighting outcome when the ratio of 257 future to current reproduction is low (Enquist & Leimar 1990). The relationship between male 258 259 residency and increased competitive ability that was observed in our experimental conditions 260 might be explained by the fact that the resident males are more adapted to the environment 261 (which is comprised of bacterial/female pheromone odours) of the drop on which they reached adulthood than the intruder males. The extent to which residency effects for male 262 entomopathogenic nematodes occurs in nature is unclear; male entomopathogenic nematodes 263 264 compete in a restricted space, an insect cadaver, in which it might sometimes be difficult to claim residency. However, larger insect cadavers such as wax moth (G. mellonella) may be 265 large enough to allow some degree of compartmentalization or localization of residents, 266 because at the time the invading nematodes are adult, the wax moth cadaver retains internal 267 structural integrity, allowing physical compartmentalization of individuals within it. Regions 268 of the cadaver could also be chemically differentiated, especially if colonized by nematodes 269 270 derived from different lines of free-foraging IJs e.g. those that exited from different natal host species or have different bacterial strains and/or chemical signature. Another source of odours 271 that could differentiate space within a cadaver is the female residents. Female S. longicaudum 272 273 are relatively immobile, tending to coil in situ, while males are active and attracted by female pheromones (Hartley 2017). Thus, it is likely that a "territory" or patch is defined by the area 274

of the cadaver influenced by a female's odour, at least in a cadaver with a relatively low 275 276 population density, and such a patch could be defended by a resident male. The increased probability in Experiment 2 of the resident male winning a fight in drops in which the resident 277 278 was mated might be due to residual pheromones from virgin females increasing the perceived resource value of the drop. Mated S. longicaudum males fight more in the presence of a virgin 279 280 female, as shown in Experiment 1, and female S. longicaudum secrete pheromones that affect 281 males behaviourally and physiologically even when no female is present (Ebssa et al. 2008; Hartley 2017). Female pheromones enhance aggression in male arthropods including crabs, 282 (Smith et al. 1994; Sneddon et al. 2003), crickets (Buena & Walker, 2008) and parasitoids 283 284 (Liu & Hao 2019). However, female pheromones cannot be the explanation for the increased probability of the resident winning in the drops where the resident was virgin, since these 285 drops had not previously held a female. Both mated and virgin males were more likely to win 286 287 when resident, but since mated residents may have experienced pheromone, but virgin residents did not, the lack of an interaction between the effects of male mating status and 288 residency may in fact point to an asymmetrical effect of female pheromone on the aggression 289 290 of mated and virgin males. There are also some observations wherein female presence has a differential effect on fighting intensity of unmated and mated S. longicaudum males 291 292 (Kapranas & Griffin n.d.). Such differential effects of female presence or pheromones may be related to the fact that S. longicaudum that have had no female presence are not immediately 293 ready to mate (Ebssa et al. 2008). 294

Fatal fighting is expected to ensue whenever the value of the resource outbalances the risk of injury, as is the case of entomopathogenic nematode males who have to fight for access to females which are limited in space and time. Our study shows that competitor males adjust their fighting behaviour according to the objective (presence and quality of females) and subjective (male mating experience) components of resource value. Male mating

300	experience could enhance the fighting ability (resource holding potential) but also its
301	influence on fighting ability is confounded by resource value components.

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471 FIGURE LEGENDS

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Figure 1 Fights between two mated males, in the presence or absence of a female within 30 minutes. Bars show average values with asymmetrical, poisson-distributed errors. Bars with accompanied by the same letter are not significantly different (post-hoc multiple comparison tests with Bonferroni adjustment, $\alpha = 0.05$). Figure 2. Latency to first fight between two mated males in the presence or absence of a female within 30 minutes. Figure 3. Time spent fighting by two males in the presence or absence of a female within 30 minutes. Data are represented as box and whisker plots with median (horizontal line), 25–75 percentiles (boxes), ranges for the bottom 25% and the top 25% of the data values (whiskers). Figure 4. Probability of winning of mated and virgin males in pairs of one mated and one virgin adult male set up either in the drop in which mated male had been reared and mated or in the drop in which the unmated male had been reared. Bars show estimated average values with asymmetrical, binomially distributed standard errors











Objective and subjective components of resource value in lethal fights between male entomopathogenic nematodes

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4 ABSTRACT

5 Males sometimes engage in fights over contested resources such as access to mates; in this 6 case, fighting behaviour may be adjusted based on the value they place on the females. 7 Resource value RV can have two components. Males can assess the quality of females, which constitutes an objective assessment of RV. Internal state such as previous mating experience 8 9 can also influence motivation to fight thus constituting a subjective assessment of RV. If mating opportunities are scarce and available females have a major impact on the lifetime 10 reproductive success of males, then fighting can be fatal; in this situation it is uncertain 11 whether males would adjust fighting behaviour based on RV. We found that both female 12 quality i.e., virginity (objective component of RV) and male mating status (subjective 13 component of RV) influence fighting intensity between males of the entomopathogenic 14 nematode Steinernema longicaudum which engage in lethal fights. Male nematodes were 15 more likely to engage in fighting and fought longer and more frequently in the presence of 16 17 virgin (high quality) females than in the presence of mated (lower quality) females. Male 18 mating status was also found to influence fighting behaviour; mated males were the winners in staged fights between mated and virgin males. Mated males may have superior fighting 19 20 ability (enhanced resource holding potential RHP), but RV asymmetries between mated and virgin males cannot be excluded. Males were more likely to win when they were resident, but 21 22 we did not find a significant interaction effect between male mating and residency status. Key words: entomopathogenic nematodes, female quality, lethal fights, male mating status, 23 resource value. 24

25

26 INTRODUCTION

Animals frequently engage in contests over resources that impact their fitness, such as mates, 27 territories or nesting sites (Andersson 1994). Amongst the most important factors influencing 28 the extent and outcome of contest behaviour are the resource holding potential of the 29 contestants, the value of the resource and the costs of fighting (Parker 1974; Arnott & Elwood 30 31 2008; Hardy & Briffa 2013). Resource holding potential RHP (fighting ability) of the contestants may depend on their weaponry, energy reserves and body size (Parker 1974; Arnott 32 & Elwood 2009; Hardy & Briffa 2013; Rico-Guevara & Hurme 2019). Resource value RV has 33 two components: objective and subjective; Intrinsic properties of the resource that will yield a 34 certain fitness gain to a successful competitor constitute the objective components of resource 35 36 value, while the value of a given resource may vary for each contestant depending on its own internal state and prior experience (subjective component of resource value) (Enquist & Leimar 37 1987; Arnott & Elwood 2008). These components of resource value may simultaneously 38 39 influence the ultimate value of the resource (Stockermans & Hardy 2013).

Objective and subjective components of resource value have been widely studied in 40 contests between males for mates, where the contested resource is receptive females. When 41 42 females are scarce, males may be more motivated to fight, but this motivation can also depend on female quality which can be based on size (larger females are more fecund), nearness to 43 44 moult or mating status (Crespi 1988; Dick & Elwood 1990; Arnott & Elwood 2008; Keil & Watson 2010; Hoefler et al. 2009; Ancona et al. 2010; Kasumovic et al. 2011). Furthermore, 45 males' internal state, such as their own mating status, can influence subjectively the value that 46 they place on the contested resource (i.e., mates) (Arnott & Elwood 2008). Mated males may 47 become less aggressive and more likely to lose at contests due to the energetic cost of mating 48 (Brown et al. 2006, 2007; Judge et al. 2010). On the other hand, previous mating experience 49 50 has been shown to increase a male's aggression and fighting success in several animal taxa

because of changes in the male's self-assessment of RHP and perception of the resource 51 52 value, that are similar to winner/loser effects (Killian and Allen 2008; Yasuda et al. 2015, Dugatkin & Dugatkin 2011; Zhang et al., 2019). If an animal wins a contest then its 53 perception of fighting ability increases and it is more likely to initiate future fights and win 54 them, whereas losers lower their RHP perception, are less aggressive and more likely to lose 55 (Hsu et al. 2006). Mating status is also related to the residual reproductive value (number of 56 57 future offspring) which is predicted to influence male investment in fighting (Kemp 2006). An unmated male can be more aggressive and more motivated to fight because future mating 58 opportunities are of greater value to him, whereas a mated male may be less willing to engage 59 60 in costly fighting (Kemp 2006).

Fighting is costly, risking injury and using time and energy that could be used in 61 mating or other activities (Maynard Smith & Price 1973; Briffa & Sneddon 2007). As the 62 63 value of the contested resource increases, the cost of fighting can also increase (Enquist & Leimar 1987; Kokko 2013). If the contested resource has a major impact on the lifetime 64 65 reproductive success of the the contenstants, such as limited females, then fights can escalate to serious injury and even death (Enquist & Leimar 1987, 1990; Maynard Smith & Parker 66 1976). Lethal male fighting has been recorded in some vertebrates (e.g., Piper et al. 2008) and 67 68 numerous arthropods, particularly arachnids (DeCarvalho et al. 2004; Sato et al. 2013) and hymenopterans (Anderson et al. 2003; Matthews et al. 2009; Cook & Bean 2006; Hamilton 69 1979; Innocent et al. 2007; Innocent et al. 2011) and lately in entomopathogenic nematodes 70 (Zenner et al. 2014; Kapranas et al. 2016). Whereas contested resource availability, such as 71 72 potential mates, is critical in prompting lethal fights (Enquist & Leimar 1990), it is unclear to what extent the contestants also assess the quality of resource in such species. When mating 73 74 opportunities for males are extremely limited then their motivation to fight would be expected to be independent of resource quality even if the odds of dying in a fight are high (Enquist & 75

Leimar 1990). Amongst parasitoid hymenopterans with lethal male combat, some species
adjust fighting behaviour according to the value of the resource (Liu & Hao 2019), while
others do not (Innocent et al. 2011).

79 Here we investigate the effect of objective and subjective resource value on lethal contests of male entomopathogenic nematodes Steinernema longicaudum Shen and Wang. 80 We are able to manipulate the mating status of the contestant males and also the quality of the 81 82 contested resource by staging dyadic contests in a controlled environment (drops of haemolymph of their insect host). First, we assess how the presence/absence and quality of 83 available females (objective components of resource value) influence fighting behaviour 84 85 between mated males. We predict that the presence of a female, especially a virgin, will increase the frequency and/or severity of fighting in S. longicaudum. Then, we assess whether 86 male mating status (subjective components of resource value) influences the contest outcome. 87 We do not have a directional prediction of how mating influences fighting, but we 88 hypothesize that either mated or virgin males will have a consistent advantage over the other, 89 90 and this will provide insights regarding how male nematodes assess the value of their mates 91 and whether mating status influences fighting ability.

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93 MATERIALS AND METHODS

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95 Nematode Reproductive Biology and Lethal Fighting.

96 Infection by entomopathogenic nematodes including *S. longicaudum* begins when free-97 foraging infective juveniles (IJs) enter an insect host and release their symbiotic bacteria that 98 turn the host insect into a cadaver. In *S. longicaudum*, IJs develop into amphimictic males and 99 females, with a sex ratio that is slightly female biased (Alsaiyah et al. 2009). Two or more 90 generations develop until the cadaver is crowded, prompting the production of IJs that exit in

search of other hosts (Kaya & Gaugler 1993; Dillman & Sternberg 2012). In S. longicaudum 101 it was shown that males are aggressive towards each other and engage in lethal fights (Zenner 102 et al. 2014). During male-male encounters, an aggressor coils its tail end around a victim, with 103 104 the copulatory spicules towards the victim's body. If not gripped close to the tail, the victim may counter-attack, wrapping its tail around the aggressor. Grappling encounters frequently 105 end when the grasped male ceases to move part or all of its body, which may happen within 106 107 minutes (Zenner et al. 2014). Such worms usually die, though partially paralyzed worms 108 occasionally recover movement. The cuticle is sometimes punctured, but paralysis and death occur also without puncture, apparently due to damage to internal organs. 109

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111 Nematode Cultures and Behavioural Assays

112 Steinernema longicaudum (strain CB2B) cultures were routinely maintained using standard 113 procedures by passage through late instar Galleria mellonella (wax moth) larvae (Kaya & Stock 1997) at 27°C. Infective juveniles were stored in tap water at 20°C. We obtained adult 114 115 males for our experiments by placing IJs individually in 25 µl hanging drops of haemolymph 116 from G. mellonella larvae which provides a suitable environment for development to adulthood (Kaya & Stock 1997; Zenner et al. 2014). The hanging drops were placed on the lid 117 118 of a Petri dish which was then inverted over a water-filled Petri dish (6 cm diam.) to prevent 119 desiccation. The IJs in the hanging drops develop to adulthood within 3 days at 27°C. Adult males can be distinguished by their copulatory spicules and their smaller size whereas females 120 are larger in size and bear vulva. We used three days old males and females in our 121 experiments. 122

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124 Experiment 1: Effects of Female Mating Status

All males used in this experiment were mated. We placed each individual naïve male for 16-125 126 20 hours with 2 naïve females to mate. We then removed the females, and successful mating was subsequently confirmed by the production of progeny. We removed two adult males from 127 their hanging drops and placed them immediately, in pairs, back in the drop from which one 128 member of the pair had been taken. Thus, one worm was "resident" and one an "intruder", but 129 130 the identity of each was not tracked, as there were no clear visual differences between them. 131 Worms were assigned to the following treatments: (i) 2 mated males with a mated female (N= 34); (ii) 2 mated males with a virgin female (same age as mated female) (N = 31); (iii) 2 132 mated males without a female (N = 35). A mating attempt was recorded when a male coiled 133 134 around the female at the vulva; insertion of spicules into the vulva was difficult to observe due to the coiling of the male around the female. A fight was recorded once one male coiled 135 tightly around another and then let go. Continuous observations were made for 30 mins during 136 137 which the latency to first fight, the incidence (whether at least one fighting event occurred), number and (for a subsample) duration of fights were recorded. Paralysis and/or death of the 138 males was recorded at the end of the observation and after 24 hours. 139

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141 *Experiment 2: Effects of Male Mating Status*

142 In S. longicaudum, males that have been with a female can be distinguished from those that have not, based on the presence of sperm in the seminal vesicle, visible through the body wall 143 (Ebssa et al., 2008); thus, mating status of individual males in a pair could be recognized. In 144 this experiment, we placed together an adult male that had mated and a virgin one. Mated 145 males were obtained by placing two females in the drop of an individually reared adult male 146 for 24 hours, after which the females were removed to a different drop and observed for later 147 progeny production. We set up pairs of one mated and one virgin adult male either in the drop 148 in which the mated male had been reared and mated (N = 21) or in the drop in which the 149

unmated male had been reared (N = 24). In this case, since mated and unmated males differed in appearance, the identity of "resident" and "intruder" could also be traced. After 24 hours we observed which worms showed signs of paralysis or death. Individual males were identified as the mated or virgin member of the pair by examining for the presence of sperm using a Nikon Optiphot microscope (x40).

156 *Statistics*

Analysis of factors influencing the number of fights, as well copulations observed in our 157 assays and the probability of paralysis/death were explored with generalized linear models as 158 described in Briffa et al. (2013). To test the hypothesis that objective resource value affects 159 fighting behavior, we used a log-linear analysis which is appropriate for small count data, to 160 assess how female presence and mating status (predictor variables) affected the number of 161 162 fights (dependent variable). Post-hoc tests among different treatments whenever applicable were adjusted with the Bonferroni correction. The probability of paralysis and/or death at the 163 end of the 30 min observation period and 24h post observations was explored with logistic 164 analysis. Duration of fighting in different treatments (no females, mated and virgin females) 165 was analysed with a non-parametric Kruskal Wallis test and latency to fighting was analysed 166 using survival regression analysis (Moya-Larano & Wise 2000). In the second experiment, we 167 compared the incidence of paralysis/death in contests staged in drops where the resident male 168 was mated versus drops where the resident was a virgin male, by using a Fisher's exact test. 169 For those pairs in which paralysis or death occurred, we used a logistic analysis by randomly 170 picking a focal male (either mated or virgin) in each replicate and tested the effect of 171 residency and mating status (as factors) and their interaction on the probability of the focal 172 male winning the contest (Briffa et al. 2013). All analysis was performed in SPSS v.21 (IBM, 173 Armonk, NY, USA). 174

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176 **RESULTS**

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178 Female Presence and Quality

Significantly more fights were observed in the presence of virgin females than in the presence 179 of mated females or in the absence of females ($F_{2.97} = 5.79$, P = 0.004, Fig. 1). This was partly 180 explained by the higher incidence of fighting in the presence of a virgin female (74.2%) than 181 with mated or no female (44.1 and 42.9%, respectively). In addition, the latency to fight was 182 shorter in the presence of virgin females than in the presence of mated females or in the absence 183 184 of females (Cox's proportional hazards analysis: Risk = 0.448, 95% CI 0.233-0.860, P = 0.016, Fig. 2), and fights also tended to last longer when there was a virgin female present (Kruskal 185 Wallis *H* = 5.56, DF= 2, *P* = 0.062; Fig. 3). 186

The presence or quality of a female did not influence the incidence of paralysis at the end of the 30 min observation period (G_2 =1.437, P=0.487) when on average 6% of pairs had one male paralysed, nor did it affect the incidence of paralysis and/or death after 24 hours (G_2 =2.839, P= 0.242), where overall 97% of pairs had at least one male either paralysed or dead, mostly represented by dead (90.9% of pairs). Single males routinely suffer 3% mortality within 24 hours (Zenner et al. 2014).

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Moreover, the number of copulations (mating events at vulva) was higher when the female was virgin than mated (mated females: 0.88 ± 0.18 , virgin females: 1.74 ± 0.26 , $F_{1, 63} = 7.322$, P = 0.009).



200 After 24 h, paralysis or death was recorded in all (N = 21) drops where the resident male was mated, but in just 70.8% (17/24) of the drops where the virgin male was resident (Fisher's exact 201 test, P = 0.01). For the 38 pairs where there was paralysis or death after 24 h, we investigated 202 203 whether male mating status and prior residency affected the contest outcome. Mated and resident males were more likely to win (logistic regressions for male mating status: $G_{1,34} =$ 204 75.13, P < 0.001; male residency: $G_{1,34} = 10.313$, P < 0.001) (Figure 4). The effect of male 205 206 mating status was particularly strong: mated males won 92.1% of fights overall. The interaction 207 between male mating status and drop residency was not significant ($G_{1,34} = 0$, P = 0.991).

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210 **DISCUSSION**

211 The presence of a female can lead to escalated male fighting, as it has shown in spiders and parasitoid wasps (Wells 1988; Jackson et al. 2006; Liu et al. 2017). Moreover, the 212 motivation to fight and intensity of fights, and consequently the cost of fighting, is adjusted 213 according to the perceived value of the resource (Parker 1974; Maynard- Smith & Parker 214 1976; Arnott & Elwood 2008; Enquist & Leimar 1987). In our study we found that not only 215 the presence but also the quality of females had an influence on fighting behaviour of male S. 216 217 *longicaudum* nematodes; fighting was more intense in the presence of a virgin female, during our 30 min observation period. In other empirical studies, males similarly exhibited 218 significantly more aggressive behaviours, and fights were more intense in the presence of 219 220 virgin females over mated ones in parasitoid wasps (Liu & Hao 2019), wolf spiders (Hoefler et al. 2008) and Sierra dome spiders (Keil & Watson 2010). The fact that a virgin (but not a 221 mated) female enhances competition between male S. longicaudum suggests either that 222 223 females mate only once, or that there is sperm competition with first male precedence (Birkhead & Moller 1998; Simmons 2001). Sperm precedence has been documented in 224

nematodes including Caenorhabditis elegans (LaMunyon & Ward 1995; Ward & Carrel 225 226 1979). Mated female steinernematids no longer attract males over a distance (Lewis et al. 2002; Hartley 2017), and in our experiment mated females received fewer mating attempts 227 (sperm transfer was not confirmed) than virgins, suggesting at least a reduction in 228 attractiveness. Female steinernematids have at most a short window of availability for mating, 229 with a period of egg-laving succeeded by egg-hatch in utero and subsequent death ("endotokia 230 231 matricida") (Poinar 1990). The exceptionally large macrosperm of S. longicaudum are suggestive of intense sperm competition (LaMunyon & Ward 1999). Because the cost of 232 fighting is high, selection favours individuals that can assess the value of the resource and 233 234 adjust their competitive behaviour accordingly (Parker 1974; Maynard-Smith & Parker 1976; Arnott & Elwood 2008). When mates are limited over time and space and their value is equal 235 236 or higher than the future, then a major part of a male contestant's lifetime reproductive success 237 is at stake and it is predicted that fights can escalate to death (Enquist & Leimar 1987, 1990). In such cases, fights might ensue irrespective of the quality of the mates, since future 238 239 opportunities for reproduction are low, but our results show that male S. longicaudum 240 nematodes assess the quality of the females and adjust their fighting behaviour. We suggest 241 that mated females, due to either once-off mating or intense sperm competition, might simply 242 not represent a valuable resource worth fighting to death for.

Differences in the quantity and quality of food or mates lead to an 'objective' assessment of the value of the contested resource, whereas internal state dictates the subjective assessment of the resource value (Enquist & Leimar 1987; Arnott & Elwood 2008). In dyadic fights between mated *vs* virgin males, the mated males were more competitive, resulting in paralysis and death of the virgin male in over 90% of cases. Mating could enhance fighting skills in nematodes, since similar coiling movements of males around females during copulation are used to lock, paralyse and consequently lead to death of their opponents.

250 Increased fighting skills enhance resource holding potential (Briffa & Lane 2017).

Alternatively, mating in males can result in an increased assessment of fighting ability or an
overestimation of the contested resource value (Killian & Allen 2008; Yasuda et al. 2015).
Mating experience has similar effects to 'winning a fight' experience, which is expected in
turn to raise the subjective value of a female in future contests (Hsu et al. 2006).

255 Contests are frequently asymmetrical; one such asymmetry is between territory owners or residents, and intruders. In such cases, it is usually the resident that wins (Hardy & Briffa 256 2013). Owner-intruder asymmetry is less likely to affect fighting outcome when the ratio of 257 future to current reproduction is low (Enquist & Leimar 1990). The relationship between male 258 259 residency and increased competitive ability that was observed in our experimental conditions 260 might be explained by the fact that the resident males are more adapted to the environment 261 (which is comprised of bacterial/female pheromone odours) of the drop on which they reached adulthood than the intruder males. The extent to which residency effects for male 262 entomopathogenic nematodes occurs in nature is unclear; male entomopathogenic nematodes 263 264 compete in a restricted space, an insect cadaver, in which it might sometimes be difficult to claim residency. However, larger insect cadavers such as wax moth (G. mellonella) may be 265 large enough to allow some degree of compartmentalization or localization of residents, 266 because at the time the invading nematodes are adult, the wax moth cadaver retains internal 267 structural integrity, allowing physical compartmentalization of individuals within it. Regions 268 of the cadaver could also be chemically differentiated, especially if colonized by nematodes 269 270 derived from different lines of free-foraging IJs e.g. those that exited from different natal host species or have different bacterial strains and/or chemical signature. Another source of odours 271 that could differentiate space within a cadaver is the female residents. Female S. longicaudum 272 273 are relatively immobile, tending to coil in situ, while males are active and attracted by female pheromones (Hartley 2017). Thus, it is likely that a "territory" or patch is defined by the area 274

of the cadaver influenced by a female's odour, at least in a cadaver with a relatively low 275 276 population density, and such a patch could be defended by a resident male. The increased probability in Experiment 2 of the resident male winning a fight in drops in which the resident 277 278 was mated might be due to residual pheromones from virgin females increasing the perceived resource value of the drop. Mated S. longicaudum males fight more in the presence of a virgin 279 280 female, as shown in Experiment 1, and female S. longicaudum secrete pheromones that affect 281 males behaviourally and physiologically even when no female is present (Ebssa et al. 2008; Hartley 2017). Female pheromones enhance aggression in male arthropods including crabs, 282 (Smith et al. 1994; Sneddon et al. 2003), crickets (Buena & Walker, 2008) and parasitoids 283 284 (Liu & Hao 2019). However, female pheromones cannot be the explanation for the increased probability of the resident winning in the drops where the resident was virgin, since these 285 drops had not previously held a female. Both mated and virgin males were more likely to win 286 287 when resident, but since mated residents may have experienced pheromone, but virgin residents did not, the lack of an interaction between the effects of male mating status and 288 residency may in fact point to an asymmetrical effect of female pheromone on the aggression 289 290 of mated and virgin males. There are also some observations wherein female presence has a differential effect on fighting intensity of unmated and mated S. longicaudum males 291 292 (Kapranas & Griffin n.d.). Such differential effects of female presence or pheromones may be related to the fact that S. longicaudum that have had no female presence are not immediately 293 ready to mate (Ebssa et al. 2008). 294

Fatal fighting is expected to ensue whenever the value of the resource outbalances the risk of injury, as is the case of entomopathogenic nematode males who have to fight for access to females which are limited in space and time. Our study shows that competitor males adjust their fighting behaviour according to the objective (presence and quality of females) and subjective (male mating experience) components of resource value. Male mating

300	experience could enhance the fighting ability (resource holding potential) but also its
301	influence on fighting ability is confounded by resource value components.

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471 FIGURE LEGENDS

Figure 1 Fights between two mated males, in the presence or absence of a female within 30 minutes. Bars show average values with asymmetrical, poisson-distributed errors. Bars with accompanied by the same letter are not significantly different (post-hoc multiple comparison tests with Bonferroni adjustment, $\alpha = 0.05$). Figure 2. Latency to first fight between two mated males in the presence or absence of a female within 30 minutes. Figure 3. Time spent fighting by two males in the presence or absence of a female within 30 minutes. Data are represented as box and whisker plots with median (horizontal line), 25-75 percentiles (boxes), ranges for the bottom 25% and the top 25% of the data values (whiskers). Figure 4. Probability of winning of mated and virgin males in pairs of one mated and one virgin adult male set up either in the drop in which mated male had been reared and mated or in the drop in which the unmated male had been reared. Bars show estimated average values with asymmetrical, binomially distributed standard errors











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ANIMAL WELFARE NOTE: Objective and subjective components of resource value in lethal fights between male entomopathogenic nematodes

The experiments reported in this paper were conducted with invertebrates – insects and nematodes - from laboratory cultures. The experiments were conducted in accordance to Science Foundation Ireland's policy concerning use of animals in research (outlined in Directive 2010/63/EU) and is implemented by the Health Products Regulatory Authority (HPRA), the competent authority in Ireland responsible for the protection of animals used for scientific purposes. All experimentation reported in this work abides to the principles of replacement, reduction and refinement as endorsed by the Association for the Study of Animal Behaviour (ASAB).