Territorial defence in a network: audiences only matter to male fiddler crabs primed for
 confrontation

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Lay Summary: Being part of a social network means that responses to social confrontations are likely to be more complex than they might seem. Indeed, here we find effects of a wider network of conspecifics on an individual's behaviour in male European fiddler crabs. Males became more aggressive toward intruders if their neighbour was watching when they had previously observed an aggressive interaction between their neighbour and a male territory intruder.

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11 Abstract

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Territorial contests often occur in the presence of conspecifics not directly involved in the 13 interaction. Actors may alter their behaviour in the presence of this audience, an 'audience 14 15 effect', and audiences themselves may alter their behaviour as a result of observing an interaction, a 'bystander effect'. Previous work has documented these effects by looking at 16 each in isolation, but to our knowledge, none has investigated their interaction; something 17 that is more likely to represent a realistic scenario for species where individuals aggregate 18 19 spatially. We therefore have a somewhat limited understanding of the extent and direction of 20 these potentially complex indirect social effects on behaviour. Here we examined how audience and bystander effects work in tandem to modify resident male aggressive behaviour 21 towards intruders in European fiddler crabs, Afruca tangeri. We found that male crabs with 22 an audience showed greater aggressive behaviour towards an intruder compared to males 23 without an audience, but only if they had acted as a bystander to an aggressive signalling 24 interaction prior to the intrusion. Indeed, bystanding during aggressive interactions elevated 25

aggressive responses to intruders maximally if there was an audience present. Our results suggest that bystanding had a priming effect on territory-holding males, potentially by providing information on the immediate level of competition in the local neighbourhood, and that same-sex audiences only matter if males have been primed. This study highlights the fundamental importance of considering broader interaction networks in studying real-world dyadic interactions and of including non-vertebrate taxonomic groups in these studies.

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33 Key words: audience effect, bystander effect, invertebrate, fiddler crab, Afruca tangeri, Uca

34 *tangeri*

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Considering communication in the context of a wider network has revealed behaviours and 37 38 effects such as eavesdropping, audience effects and bystander effects, that would not be observable from a dyadic approach (McGregor, 2005). Territorial contests are typically 39 settled through pair-wise interactions within a network of multiple individuals, rather than in 40 (McGregor and Dabelsteen, 1996). Audience effects occur when animals modify their 41 behaviour due to the presence of other individuals not involved in the interaction 42 43 (Zuberbühler, 2008) and these effects have been shown to alter the intensity of agonistic displays in a number of species (Cruz and Oliveira, 2015; dos Santos et al., 2017; 44 Fitzsimmons and Bertram, 2013; Montroy et al., 2016; Setoguchi et al., 2015). In the context 45 46 of territorial defence, the audience effect has been demonstrated to be dependent not only on 47 the sex of the audience, but also on the territorial status and familiarity of individuals in the wider network (Bertucci et al., 2014; Dzieweczynski et al., 2005). Observed differences in 48 49 behaviour in the presence of an audience suggest that individuals can (1) assess attributes of their audience and (2) adjust their behaviour as a strategy to counter costs (or strengthen 50 benefits) that can come with eavesdropping, or more generally, gathering of social 51 information by audiences (Earley and Dugatkin, 2002). In the bystander effect the audience 52 53 members themselves are influenced by observing an interaction (Earley and Dugatkin, 2002; 54 Oliveira et al., 2001; Peake et al., 2006). The observation acts directly on the motivational system (Hirschenhauser and Oliveira, 2006; Oliveira et al., 2001) and prepares individuals for 55 what may happen next in their social environment (Antunes and Oliveira, 2009). Individuals 56 57 can, for example, be primed to augment their levels of aggression in interactions following bystanding. For example, Clotfelter and Paolino (Clotfelter and Paolino, 2003) found 58 increased aggressiveness by Siamese fighting fish, Betta splendens, towards a novel male 59

conspecific after the observation of an aggressive interaction. However, reverse priming, a
reduction in aggressive behaviour, has also been found, for example, in a study of crayfish, *Orconectes rusticus* (Zulandt et al., 2008), which to our knowledge is also the only
invertebrate species in which bystander effects have been investigated.

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Although audience and bystander effects are frequently documented, it is not clear from the 65 literature how bystander and audience effects may interact. This represents a potential key 66 gap in our understanding of the strength or importance of these effects in *in situ* contexts 67 68 where they are likely to happen concurrently, particularly in species where individuals live in aggregated communities. A study in Siamese fighting fish, suggests that simply viewing an 69 70 unfamiliar male prior to a contest with another male can prime males for increased aggression 71 during a fight, irrespective of whether the audience is present or not during the fight itself 72 (Matos et al., 2003). It also highlights the complexity of bystander and audience effects and the need to manipulate these effects in tandem in order to deepen our understanding of the 73 74 significance of dynamics in the social environment for the expression of behaviour. Here we investigate the interplay between audience and bystander effects on the territorial behaviour 75 76 of male European fiddler crabs, Afruca tangeri. Fiddler crabs are very unlikely to be found in isolation (Pope, 2005), and therefore have a high likelihood of engaging in social interactions 77 with multiple receivers within signal range. Males actively defend territories around their 78 79 burrows (Hemmi and Zeil, 2003) with their one greatly enlarged claw, used for the production of highly conspicuous visual agonistic and courtship signals and for fighting 80 (Oliveira and Custódio, 1998; Wolfrath, 1993). Male crabs without a burrow will wander 81 82 through the population and challenge burrow holders in attempts to acquire a burrow (Jordao and Oliveira, 2005). Burrow holding males must therefore invest in an agonistic response in 83 84 order to retain residency (Oliveira and Custódio, 1998). Recent work in another fiddler crab

85 species suggests that same-sex audiences do not matter for expressed levels of aggression (dos Santos et al., 2017), given the ubiquity of males in the immediate social environment 86 (i.e. males are highly likely to have male neighbours), this is perhaps not surprising. We 87 88 propose instead that males will fine-tune their responsiveness to their social environment according to current social information, such as the immediate level of competition in the 89 90 neighbourhood. If males observe a territorial dispute in their neighbour's territory, they should be primed for a possible challenge of their own territory and therefore will be more 91 92 responsive or sensitive to their immediate social environment. In this case, by standing to an 93 aggressive interaction between a neighbouring male and a stranger (intruding male), should augment the aggressive response exhibited by males towards intruders appearing close in 94 95 time. This should be particularly so when the social stimuli are increased by a male audience 96 as this audience can gather information on the interaction outcome (e.g., the competitive 97 ability of his male neighbour) and in the least, acts as an indicator of a more competitive social environment (i.e. more males are present) than when no male audience is present. In 98 99 this investigation, we used an *in situ* experimental manipulation of the social environment that burrow holding male crabs experienced to test such effects during territorial 100 101 confrontations in the home environment.

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103 Methods

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105 The study was carried out in the Parque Natural da Ria Formosa, Portugal (N370927, 106 E073244) from May to July 2012 in order to quantify audience and bystander effects *in situ* 107 in males of a free roaming population of European fiddler crabs. To test focal male crabs at 108 their home burrow an arena made of bamboo and sand-coloured fabric was placed around 109 two neighbouring males and their burrows (Fig. 1). Neighbouring male pairs were selected 110 according to three criteria: (1) they were matched in claw size (estimated by visual comparison); (2) they were exhibiting courtship behaviour (courtship waving) indicating that 111 they were actively defending a burrow; and (3) they were within one meter of one another 112 (measured using a measuring tape stretched in a straight line from the center of one burrow to 113 the other). Stimulus crabs captured from other areas of the mudflat were tethered with 10cm 114 of clear monofilament line to bamboo posts inserted into the substrate within the arena to 115 simulate wandering male crabs (intruders). This method of tethering stimulus crabs is 116 relatively standard and has been used successfully in this and several other fiddler crab 117 118 species to elicit both courtship and territorial behaviour as observed under natural conditions (Detto and Backwell, 2009; Detto et al., 2006; Detto et al., 2010; How et al., 2008; Milner et 119 al., 2010; Pope, 2005; Reaney, 2007)(Booksmythe et al., 2010). Intruders were matched in 120 121 claw size to focal crabs using visual comparison. Focal crabs were exposed to two 5-minute phases: a 'bystanding phase' (neighbour interacts with a simulated intruder) and a subsequent 122 'interaction phase' (focal male interacts with a simulated intruder) in one of four treatments 123 (3 control and 1 experimental): (1) null control - no neighbour-intruder interaction in the 124 'bystanding phase' and no audience in the 'interaction phase' (n=11); (2) audience control -125 no neighbour-intruder interaction in the 'bystanding phase' and an audience in the 126 'interaction phase' (n=10); (3) bystander control – neighbour-intruder interaction in the 127 'bystanding phase' and no audience in the 'interaction phase' (n=11); and (4) bystander and 128 129 audience (experimental) - neighbour-intruder interaction in the 'bystanding phase' and audience in the 'interaction phase' (n=11) (Fig. 1). 130

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In the 'bystanding phase' of treatments (3) and (4) a stimulus male was tethered 40cm from the neighbour's burrow (min 90 cm from the focal male's burrow) and left for 5 minutes after both males (focal and neighbour) had emerged from their burrows. In the 'bystanding phase' 135 of treatments (1) and (2) the arena was approached (and stimulus male placement simulated) and the crabs were then left for 5 minutes after both males had emerged from their burrow. In 136 the interaction phase a stimulus male was tethered 40cm from the focal male's burrow to 137 simulate a wandering male. In treatments (2) and (4) the neighbouring male was allowed to 138 emerge from his burrow and act as an audience while in treatments (1) and (3) the 139 neighbouring male's burrow was blocked to prevent him from emerging during the trial 140 141 period. Phases began when the focal or both males had surfaced (carapace and major claw visible), as applicable, and lasted for five minutes. Following a trial, burrows were marked 142 143 with a small flag and within a semi-lunar tidal cycle these areas were avoided for further testing to ensure that males were not re-used over the course of the study. We quantified 144 aggression as a behavioural state by measuring the duration of time that focal males were 145 146 engaged in aggressive behaviours toward the simulated intruder in the 5-minute interaction phase. The observed aggressive behaviour, following previously published methods (see 147 Burford et al., 2000; Oliveira et al., 1998; Wolfrath, 1993), included both non-contact 148 149 aggression (threat displays) and that involving physical contact (pushing, grappling and tossing) and the intensity of enactment represents the focal male's willingness to escalate the 150 contest (Oliveira et al., 1998; Wolfrath, 1993). We compared the time focal males spent 151 performing aggressive behaviour among treatments with an analysis of variance with 152 treatment as a fixed effect (SPSS v. 22). We included inter-burrow distance between the focal 153 154 and neighbour as a covariate in the model as this varied across focal individuals. Post hoc analyses were carried out on the marginal means using a least significant difference 155 adjustment for multiple comparisons. All behaviours were scored from video recordings 156 157 (Panasonic HDC-SD800 camcorder) of the trials by a single observer (MKM) naïve to the trial condition. 158

160 The study was carried out under permit (ICNF, Portugal) following ethical approval from the 161 first author's home institution. All captured crabs were kept singly in shaded containers filled 162 with sea water and mud prior to testing and released back to their area of capture at the end of 163 a trial. Crabs were marked with non-toxic paint to ensure that they were not used more than 164 once in case of recapture.

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166 **Results**

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168 There was an overall effect of the treatment that focal males experienced on the amount of aggressive behaviour they displayed toward a simulated intruder ($F_{3,38}=11.797$, p<0.0001, 169 Fig. 2). Our post hoc analysis revealed that there was an effect of having a neighbouring male 170 171 audience present during a territorial intrusion on the level of aggression expressed by a burrow-holding male, but only if burrow-holding males had previously acted as bystander 172 (Treatment 4) to a neighbouring aggressive interaction (Table 1; Fig. 2). In short, males in 173 174 Treatment 4 behaved more aggressively than in any other treatment. Males in two of the control treatments (Treatments 1 and 2) did not differ from one another in behaviour, but 175 males in the null control (no bystanding and no audience; Treatment 1), were less aggressive 176 than males in the bystander control (bystanding, no audience) (Treatment 3). 177

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179 Discussion

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181 Gleaning information from the social environment is essential for an individual to respond to 182 this environment appropriately (Danchin et al., 2004; Seppanen et al., 2007; Valone, 2007). 183 Individuals living in systems that are likely to be high in social noise will be under particular 184 pressure to either filter out information to avoid an inappropriate response or to use specific 185 cues to alert them to a situation where an active response may be required (e.g., Fitzsimmons et al., 2008; Naguib et al., 2004). This latter priming effect may be particularly important in 186 predicting that, for example, a territorial challenge is likely to occur. The anticipation of 187 188 being challenged is likely to affect not only how individuals respond to a rival conspecific during a direct interaction, but also how they respond to the presence of other conspecifics in 189 the immediate environment. These individuals that are not part of the interaction, may 190 themselves be gleaning social information (i.e. they are conspecific audiences). In this study 191 we found the first evidence that a neighbouring male audience moderated male European 192 193 fiddler crab behaviour towards an intruder, but only if males had acted as bystander to an aggressive interaction between a male neighbour and a same-sex intruder immediately prior 194 195 to the encounter. This suggests that focal males that had viewed an aggressive interaction 196 between his neighbour and an intruder were somehow primed for heightened responsiveness 197 to having a male audience present during an interaction between himself and a territorial intruder. The observed increase in aggressiveness was likely also a function of a general 198 priming for an aggressive response as a result of the bystanding, since we did detect greater 199 aggressiveness when males without an audience had acted as bystander compared to males in 200 201 our null control (no audience and no bystanding).

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The prevalence of the use of public information (Danchin et al., 2004) suggests that it could be beneficial for individuals to employ strategies to manage the perceptions of unintended receivers. Fighting could communicate the focal male's motivation to defend his territory (Detto et al., 2010) or ability to do so (Peake et al., 2001) to his neighbour. However, we did not see an effect of a neighbouring male audience unless the focal male had previously acted as a bystander to an aggressive interaction between his neighbour and an intruder. Bioregulatory mediators of the adjustment of aggressive behaviour are likely to be affected in 210 males that anticipate a territorial challenge based on cues in their social environment; facilitating an adjustment to an increased competitive environment with the appropriate 211 behavioural response (Antunes and Oliveira, 2009). It could be that the simple presence of a 212 213 neighbour is not enough of a cue of the extent of the competitive environment, given the ubiquity of territorial neighbours, and their familiarity, in a male's visual field (Detto et al., 214 2010; Hemmi and Zeil, 2003; Pope, 2005). Instead, if a territorial challenge has been 215 observed immediately prior to an intrusion, effects on bio-regulatory mechanisms (Oliveira et 216 al., 2001) could drive a response directly or at least provide a cue as to the probability of an 217 218 escalation of aggression that could lead to a territorial takeover (Oliveira, 2009).

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220 The bystander effect detected in this study is consistent with work in vertebrate species 221 demonstrating increased levels of aggression (e.g., Clotfelter and Paolino, 2003), but 222 inconsistent with the reverse priming demonstrated in another crustacean species (Zulandt et al., 2008). In our study, burrow holding males will have invested energy into establishing 223 territory boundaries with their neighbours (Detto et al., 2010) and as such intruders are a 224 potential threat to the territory more generally if they were to take over a neighbour's burrow. 225 Zulandt and colleagues (Zulandt et al., 2008) suggest that the bystander effect could be 226 resource dependent such that when resources are abundant, and consequently their value 227 228 reduced, an individual will avoid escalation if the observation of a fight indicates that there is 229 an increased likelihood of being challenged within their network. In the present study males were at their home burrow and as a consequence were defending a valuable resource; an 230 added ecological realism that may have contributed to the direction of the effect. 231

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Our study demonstrates that there is likely to be a complex interaction between audience andbystander effects in systems where individuals are part of a network of conspecific

235 interactants. The results suggest that selection has enhanced the use of public information to increase the appropriateness of responses to social stimuli. We investigated effects of the 236 presence of a fiddler crab's nearest neighbour, an individual that the focal is likely to have 237 interacted with before and to be familiar with, however, laboratory work on audience effects 238 has demonstrated that attributes of an audience can influence physiological and behavioural 239 effects on individuals, for example, sex and territorial status (Dzieweczynski et al., 2005). 240 Future *in situ* work investigating behavioural reactions when the audience is a *non*-neighbour 241 or a female, would therefore be particularly enlightening. 242

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357 Figure legends

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Figure 1. Overview of experimental arenas and 4 treatments (3 control and 1 experimental) used to investigate bystander and audience effects in male European fiddler crabs (see text for details).

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Figure 2. Average time that male European fiddler crabs spent engaging in aggressive
behaviour toward experimentally introduced male intruders in 4 social treatments (see Fig. 1)
designed to test for bystander and audience effects (Estimated Marginal Mean (EMM) ±SE;
matching letters indicate treatments where p>0.05 for tested differences).

- 368 Table 1. Results of the *post hoc* analysis of differences among treatments in burrow-holding
- 369 male aggressive responses to simulated intrusions by conspecific male fiddler crabs. P-values
- 370 in bold indicate where significant differences in aggressiveness were found.

			95% CI for Difference	
Treatment B	Mean difference	Lower bound	Upper bound	Р
2 - Audience control	-31.04	-82.41	20.33	0.229
3 - Bystander	-53.73	-103.39	-4.07	0.035
4 - Bystander and audience	-143.18	-194.50	-91.85	p<0.001
3 - Bystander	-22.69	-73.91	28.53	0.375
4 - Bystander and audience	-112.14	-163.35	-60.92	p<0.001
4 - Bystander and audience	-89.45	-140.48	-38.42	0.001
	Treatment B 2 - Audience control 3 - Bystander 4 - Bystander and audience 3 - Bystander 4 - Bystander and audience 4 - Bystander and audience	Mean differenceTreatment Bdifference2 - Audience control-31.043 - Bystander-53.734 - Bystander and audience-143.183 - Bystander-22.694 - Bystander and audience-112.14audience-112.14audience-89.45	95% Cl forMeanLowerTreatment Bdifferencebound2 - Audience control-31.04-82.413 - Bystander-53.73-103.394 - Bystander and audience-143.18-194.503 - Bystander-22.69-73.914 - Bystander and audience-112.14-163.354 - Bystander and audience-112.14-163.35	MeanLowerUpperTreatment Bdifferenceboundbound2 - Audience control-31.04-82.4120.333 - Bystander-53.73-103.39-4.074 - Bystander and audience-143.18-194.50-91.853 - Bystander-22.69-73.9128.534 - Bystander and audience-112.14-163.35-60.92audience-89.45-140.48-38.42

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