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Fine-scale dynamics of competitive fertilisation in social groups of red junglefowl (Gallus gallus) shed new light on avian sperm competition

Rômulo Carleial^{1†*}, Grant C. McDonald^{1,2†*}, Lewis G. Spurgin³, Eleanor A. Fairfield³, Yunke Wang¹, David S. Richardson³, Tommaso Pizzari^{1*}

¹Department of Zoology, Edward Grey Institute, University of Oxford, Oxford, OX1 3SZ, United Kingdom ²Department of Ecology, University of Veterinary Medicine Budapest, Budapest 1077, Hungary ³School of Biological Sciences, University of East Anglia, Norwich, United Kingdom.

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13 17 Studies of birds have made a fundamental contribution to elucidating sperm competition processes, 18 experimentally demonstrating the role of individual mechanisms in competitive fertilisation. However, the 19 relative importance of these mechanisms and the way in which they interact under natural conditions 20 remain largely unexplored. Here, we conduct a detailed behavioural study of freely-mating replicate 21 groups of red junglefowl, Gallus gallus, to predict the probability that competing males fertilise individual 22 eggs over the course of 10-day trials. Remating frequently with a female and mating last increased a male's 23 probability of fertilisation, but only for eggs ovulated in the last days of a trial. Conversely, older males, 24 and those mating with more polyandrous females, had consistently lower fertilisation success. Similarly, 25 resistance to a male's mating attempts, particularly by younger females, reduced fertilisation probability. 26 After considering these factors, male social status, partner relatedness and the estimated state of a male 27 extragonadal sperm reserves did not predict sperm competition outcomes. These results shed new light on 28 sperm competition dynamics in taxa such as birds, with prolonged female sperm storage and staggered 29 fertilisations. $\overline{30}$

1. Introduction 31

32 Parker (1970) [1] revolutionized sexual selection theory by proposing that when females obtain sperm from 33 multiple males (polyandry), competition among males continues after mating as their ejaculates compete 34 to fertilise a set of ova, a process which became known as sperm competition [2-9]. While Parker's (1970) 35 intuition was inspired by insects, the realization that polyandry is ubiquitous among sexually-reproducing 36 organisms and often intense [10] has catalysed the study of sperm competition across different taxa [11-13]. 37 An increasing number of studies have sought to experimentally isolate the causal effect of individual 38 sperm competition mechanisms on male reproductive success by quantifying the overall share in paternity 39 at a particular point in time, e.g. the end of a reproductive cycle. This research has been instrumental in 40 identifying male- [4,6,7,13-18] and female-driven [19,20] mechanisms underpinning paternity share under 41 controlled conditions.

42 In natural populations however, multiple variables act simultaneously and interact with each 43 other to determine the outcome of sperm competition. Understanding these interactions is particularly 44 challenging for many internally-fertilising taxa where prolonged female sperm storage increases the 45 temporal window for sperm competition [21,22] and generates opportunity for complex patterns of sperm 46 storage and precedence [23]. For example, in the fruit fly, Drosophila melanogaster, multiple mechanisms 47 including passive sperm release from the female storage organs, active displacement of resident sperm by 48 the second-male sperm, and female-controlled sperm ejection, lead to approximately 80% of eggs being 49 fertilised by the second male to copulate with doubly-mated females [24,25]. An understanding of the 50 temporal dynamics of competitive fertilisation in natural settings remains elusive however, largely due to 51 the difficulty in disentangling multiple mechanisms occurring within the female reproductive tract. 52 Addressing this outstanding challenge is fundamental to resolving the operation of sexual selection and

related processes, such as alternative mating strategies, sexual conflict and the maintenance of geneticvariation in natural populations [5,26,27].

55 Historically, birds have represented a key vertebrate model system for the empirical study of 56 sperm competition [12,28-30], and a crucial counterpoint to studies of insects [1,13,24,25,31-34]. For 57 example, the juxtaposition of avian and insect sperm competition dynamics has helped clarify the 58 differential roles of sperm size and numbers in these clades [35]. The importance of birds is in part due to 59 field studies of extra-pair copulation and extra-pair paternity in socially monogamous species [8,12,29,36-60 39] and the wealth of information on the reproductive biology of the domestic fowl, Gallus domesticus, and 61 other domestic galliforms, which have informed the development of mechanistic models of sperm 62 competition [30,40,41]. As in most other species of birds, female fowl typically ovulate one egg on each 63 successive day of the laying cycle, with fertilisation occurring shortly after ovulation [42]. Females store 64 viable sperm in epithelial invaginations at the utero-vaginal junction of the oviduct - the sperm storage 65 tubules (SSTs) - for up to approximately two weeks [40-43]. During this time, sperm are thought to be 66 passively lost from SSTs at a near constant rate, and sperm from competing ejaculates mix randomly 67 within individual SSTs. Passive sperm loss combined with random sperm mixing within SSTs predicts that 68 - all else being equal - the last male to copulate with a female should retain a fertilisation advantage for 69 subsequent eggs, which is proportional to the time elapsed between competing inseminations [30,41]. The 70 extent of this advantage is modulated by a range of factors. For example, shortly before and after 71 oviposition, the fertilisation success of an ejaculate is reduced due to oviduct contractions associated with 72 egg laying (reviewed in [30]). Moreover, studies of domestic populations, which experimentally controlled 73 for insemination order through the simultaneous artificial insemination of two competing ejaculates, have 74 demonstrated that relative differences in either sperm numbers [44] or fertilising efficiency [45,46] explain 75 variation in paternity share between two competitors. Subsequent artificial insemination work in which 76 both sperm number and fertilising efficiency (mobility) were experimentally manipulated has provided 77 evidence of temporal dynamics in a population of domestic fowl: differences in sperm numbers predicted 78 the paternity of eggs produced shortly following insemination, while differences in mobility predicted 79 paternity later in the laying sequence [47]. Collectively, these experimental studies form the basis of our 80 operational understanding of the mechanisms of avian sperm competition, and suggest that the 81 probability that a given male fertilises a particular egg is determined by the number of other competitors 82 inseminating the same female, the time elapsed between these inseminations and ovulation, and the size 83 and fertilising efficiency of the inseminations of this male in relation to those of his competitors, broadly 84 confirming patterns identified by previous work [48]. However, the extent to which this view adequately 85 captures sperm competition dynamics in natural bird populations remains unknown. Some studies of 86 natural populations have shown that the number of sperm reaching an egg and paternity share can change 87 systematically over the course of a laying cycle in some species [49-56]. Other studies however, have failed 88 to confirm these results [54,56], or found less conclusive [57] or more complex patterns [58,59]. A 89 fundamental source of inconsistency is that most studies of free-ranging groups typically lack information 90 about individual mating events, which makes it impossible to assess the level of sperm competition 91 associated with the fertilisation of individual ova. In the current study we aim to address this knowledge 92 gap by combining fine-grained information on socio-sexual interactions and parentage of individual 93 zygotes in replicate flocks of red junglefowl, G. gallus, the main species that has given rise to the domestic 94 fowl.

95 In nature, red junglefowl live in polygynandrous groups, socially structured in sex-specific 96 hierarchies [60]. Recent work indicates that variance in paternity share is the largest source of variance in 97 male reproductive success, indicating the importance of sperm competition and post-copulatory sexual 98 selection in these populations [61,62]. Sperm competition favours males that remate with the same female 99 at a high frequency [61,62], and a similar advantage derived by frequent remating with the same female 100 has been reported for some wild bird species [63] and in other taxa, e.g. some insects [64-66]. Mating in 101 fowl is strongly modulated by the interaction of female and male behaviours. Studies of domestic fowl and 102 red junglefowl indicate that while in small, strongly female-biased groups mating is largely driven by 103 females, in larger groups and groups with a higher proportion of males, the majority of mating attempts is 104 initiated by males [60]. However, differential female behaviour plays an important role in determining the 105 outcome of these male mating attempts. For example, females can influence the success of an insemination 106 by differentially resisting copulation attempts, manipulating male-male competition and copulation 107 interference, and by ejecting semen after insemination [60]. These mechanisms can be modulated by 108 characteristics of individual males and females, and their interactions. For example, male social status 109 mediates competitive access to mates and mating opportunities, and is favoured by female responses 110 before and after mating [60]. On the other hand, male status may be negatively related to sperm fertilising 111 efficiency [46,67], as observed in species with more distinct alternative mating tactics [68,69], suggesting a 112 possible trade-off between male investment in pre- vs. post-copulatory intrasexual competition. Similarly, 113 male age has been linked to lower fertilisation success, as older males transfer fewer sperm than younger 114 males [70,71]. Moreover, sperm of older males tend to swim more slowly, possibly due to lower levels of 115 seminal antioxidants [72]. Female age may also affect the outcome of sperm competition. On the one hand,

116 older females are less polyandrous and avoid males more often than younger females [73], and have a

117 faster rate of sperm loss from the SSTs than younger females [74]. On the other, resistance to males and

118 male coercion are costly [75] and older females might be less able to resist male attempts. Finally, genetic

119 relatedness between a male and a female may influence fertilisation success. Experimental evidence 120 suggests that red junglefowl females favour sperm from unrelated males after mating, possibly in order to 121 reduce the risks of inbreeding depression [76,77], a pattern observed in some other taxa [20]. Despite this 122 experimental work however, we currently lack an understanding of how these mechanisms collectively 123 influence sperm competition dynamics in social groups of freely-interacting birds.

124 Here, we study mechanisms underpinning fowl sperm competition by using detailed mating 125 behaviour and parentage data of freely-mating replicate groups of a captive population of red junglefowl. 126 This dataset has been previously analysed to investigate the relationship between male phenotypes and 127 overall male reproductive success [62], the role of female sociality on the structure of sociosexual networks 128 [73], and changes in female and male phenotype associated with mating trials [75]. This earlier work 129 identified a role for both male and female behaviours in determining variation in individual reproductive 130 success [62,73,75]. The current study consolidates and builds on this earlier work, with a drastically 131 different analytical approach which investigates day-to-day variation in the probability of fertilisation of 132 individual eggs laid over successive days of the mating trial. By doing so, the present study moves beyond 133 previous analyses of variation in overall reproductive success and develops a novel methodological 134 platform to examine the relative importance of different factors dynamically through time. We predicted 135 that males that remate more often and those that copulate last with a female will have a higher probability 136 of fertilising an egg. Based on the passive sperm loss model [30], we predicted that the last male advantage 137 should increase with the time elapsed between competing inseminations. We further predicted that 138 probability of fertilisation declines with male age, the level of female resistance (which we predicted to be 139 weaker in older females), the relatedness of the male, the number of males mating with the female, and the 140 depletion of a male's extragonadal sperm stores through previous copulations. Male status might have a 141 positive effect, e.g. through preferential female sperm retention, or a negative effect, through a possible 142 trade-off with fertilising efficiency.

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2. Methods 144

145 We studied a population of red junglefowl at the John Krebs Field Station of the University of Oxford. We 146 analysed the data collected in 20 mating trials conducted in 2011-2013. Detailed accounts of the protocol of 147 the mating trials have been published elsewhere [62,73,75]. Briefly, in each mating trial one of us (GCM) 148 observed a group (males=10, females=12) for 3h in the morning and 3h in the evening on each of 10 149 consecutive days (day1-10). We recorded all male and female mating related behaviours (males: successful 150 and unsuccessful mating attempts, courtship waltzes; females: mating solicitations, level of resistance to 151 mating attempts [78]), and male agonistic interactions (pecks, fights, aggressive waltzes and avoidances). 152 Eggs (n=982) were collected daily from the second day of the trial (day2) to the day following the end of 153 the trial (day11), artificially incubated for 5-9 days, and parentage of the embryos was assigned 154 molecularly [62].

155 We analysed the effect of multiple predictor variables on the probability of fertilisation of 156 individual eggs by competing males, which were defined as all males that mated successfully with the 157 female up to, and including, the day before the laying of an egg. Thus, copulations on the day an egg was 158 laid were not considered as competing for the fertilisation of that egg, because it is unlikely that sperm 159 from these inseminations could have reached the egg in time for fertilisation [30,40,42]. We considered the 160 following predictor variables (i.e. fixed effects, see electronic supplementary material for fuller discussion): 161 i) number of male competitors, ii) remating rate (the frequency at which a male mates with the same 162 female up until the day prior to egg-laying), iii) mating order, iv) male social status (calculated using Elo 163 scores [79]), v) male age, vi) average female resistance to mating attempts of the focal male (scored 164 following [78]), vii) female age, viii) male-female relatedness (calculated using Wang's estimator [80,81]), 165 ix) an index of the male extragonadal sperm reserves (MSR, Figure S1), and x) time (i.e. the day of the 166 mating trial when the egg was laid).

167

168 (a) Model comparison

169 All analyses were conducted in R v3.6.2 [82], using the "lme4" v1.1-21 package [83]. We built a generalized 170

linear mixed model (GLMM) with binomial distribution and logit function with the fertilisation outcome 171

- of each egg (unsuccessful males=0, successful male=1) as the response variable, predictors (i-ix) and their
- 172 interaction with time (x) as fixed effects (mod13, Tables 1, S1). We used relative values of a predictor
- 173 variable (i.e., the value of each male divided by the mean value of all males competing for that particular
- 174 egg), to standardise a male's value in relation to his direct competitors and to avoid conversion problems
- 175 due to large scaling differences among predictors. Mating order (iii) and relatedness (viii) were not
- 176 standardised because already expressed in relative terms. Collinearity across predictors was calculated 177 using variance inflation factors (VIFs) implemented in the package "car" v3.0-3 [84]. All predictors showed

178 VIFs<3 and were therefore retained [85]. We compared this model (mod13) against simpler models with 179 fewer predictors using the Akaike's information criterion (AIC [86]), such that a difference (ΔAIC) < 2 was 180 interpreted as no difference of fit between models, $2 \le \Delta AIC < 4$ as a difference of fit, $4 \le \Delta AIC < 10$ as a 181 considerable difference of fit, and $\Delta AIC > 10$ as a strong preference for one model over the other [87]. 182 Overall, we defined 44 models a priori (Table S1) using different combinations of predictors, and compared 183 them simultaneously using the package "AICcmodavg" v2.2-2 [88]. All 44 models also included male 184 identity, female identity, female group identity (i.e. the identity of the unique unit of 12 females used in 185 each trial [42]), and mating group identity (i.e., the identity of the unique group of 10 males 12 females 186 used in each trial [42]) as random effects to account for sources of non-independence [42]. We quantified 187 the variance explained by each model as marginal R² (R²GLMM(m)), i.e. the variance explained by fixed 188 effects alone, and conditional R^2 (R^2 GLMM(c)), i.e. the variance explained by the combination of the fixed 189 and random effects [89,90]. Finally, while a male's remating rate (predictor (ii)) reflects his sperm 190 investment in a female, it does not take into consideration that sperm are passively lost from the female SSTs over time. Therefore, we also constructed GLMMs replacing (ii) with an estimated value of a male's 191 192 sperm numbers in the female SSTs at the time of fertilisation (female sperm reserves), which was 193 calculated using the rate of sperm loss described for domestic fowl [91,92] (electronic supplementary 194 material, Figure S2). We compared AIC scores between models using (ii) against models replacing (ii) with 195 female sperm reserves (Table S2).

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197 (b) The effect of individual predictors

We tested the statistical significance of individual predictors in the most complex model (mod13, Tables 1,S1), by performing likelihood ratio tests (LRTs) removing each variable of interest while holding constant the effect of all other predictors. Interaction terms that were not significant were dropped before conclusions were drawn on the significance of the main effects. To further confirm temporal patterns, we explored the effect of individual predictors (i-ix) on fertilisation probability, by running separate GLMMs for each individual day of the trial. Results were qualitatively similar to the original approach (see Table S3, Figure S4).

206 3. Results

207 Models including female sperm reserves had lower AIC scores than equivalent models with remating in 11 208 cases (Δ AIC \geq 6.46, Table S2). In the remaining cases (n=15), mostly when time was included in the model as 209 an interaction term, both models had similar AIC scores (Δ AIC \leq 2.18, Table S2). This suggests that female 210 sperm reserves captures a temporal dimension of sperm competition, and becomes redundant when time 211 is included in the model explicitly. Therefore, we only report models with remating rate below, as this 212 approach does not rely on additional assumptions of patterns of sperm loss from the SSTs.

213

214 (a) Model comparison

Using an AIC approach, the best model predicting the fertilisation probability of individual eggs (mod1, Table 1,S1) included a male's remating rate and its interaction with time (i.e. day of the trial), male mating order and its interaction with time, number of competing males, male age, and the interaction between overall female resistance and female age. Male-female relatedness, his sperm reserves (MSR) and male social status were not included in the best model. The best model was only slightly better than the model without the interaction between female resistance and female age (mod2, Δ AIC=2.86, Table S1), but performed considerably better than all other models (Δ AIC>4, Table S1).

222 The best model explained 14.98% of the variance by fixed effects alone (i.e., R²GLMM(m)), and 223 19.59% when random effects were included (i.e. R²GLMM(c)). When considering R²GLMM(m) for models 224 including only a single predictor, the model with number of competing males explained the most variance 225 (5.5%, mod24, Tables 1,S1) in the probability of fertilisation. The model with female resistance (mod30, 226 Tables 1,S1) explained 2.05%, male age (mod38, Tables 1, S1) explained 1.89%, remating rate (mod33, 227 Tables 1,S1) explained 1.12%, male social status (mod39, Tables 1,S1) explained 0.14%, and mating order 228 (mod41, Tables 1,S1) explained 0.04% of the variance in fertilisation success, respectively. The remaining 229 two predictors, relatedness and MSR, explained each 0.02% (mod43, mod44, Tables 1, S1) of the variance. 230

231 (b) The effect of individual predictors

Results of the model including all main effects and the statistically significant interactions are summarisedin Table 2.

We found a positive and significant interaction between a male's remating rate and time, such that males that copulated more often with the female had a higher probability of fertilisation later in the trial (remating rate*time: χ^2_1 =5.72, p=0.016, Figure 1a). Similarly, males that tended to mate last with a female had a higher probability of fertilising eggs laid later in the trial (mating order*time: χ^2_1 =13.17, p<0.001, Figure 1b), as predicted by the passive sperm loss model. Such last male advantage later in a trial may be caused by increasing variation in the time elapsed between the last insemination by competing males and 240 fertilisation. We explored this hypothesis by testing post-hoc whether the time interval (gap) between the

241 last insemination by the first and last competitor male in the mating order tended to increase over

successive days in the trial. Consistent with expectations the time gap spanning the mating order increased over the course of the trial (Time gap~mating order*time: χ^2_1 =98.85, p<0.001, Figure S5), suggesting that

some males that mate with a female early in a trial are disadvantaged in sperm competition later in the

trial as they fail to remate with the female.

246 Males that experienced more intense female resistance across copulation attempts were 247 significantly less likely to fertilise eggs overall (resistance: χ^2_1 =101.82, p<0.001), and this effect was constant 248 over time (resistance*time: χ^2_1 =1.99, p=0.159), but was dependent on female age. Males were less likely to 249 fertilise when resisted by younger females (resistance*female age: χ^2_1 =4.86, p=0.027, Figure 1c). Competing 250 against more males decreased a male's probability of fertilisation (competitors: χ^2_1 =193.18, p<0.001, Figure 251 1d), but there was no evidence that this effect varied over the course of a trial (competitors*time: $\chi^2_1=0.24$, 252 p=0.628). Similarly, older males had a lower probability of fertilisation overall (male age: χ^2_1 =51.43, 253 p<0.001, Figure 1e), which was consistent throughout the course of a trial (male age*time: $\chi^2_1=0.46$, 254 p=0.496).

255 The state of a focal male's sperm reserves (MSR) did not increase his overall probability of 256 fertilisation (MSR: χ^2_1 =0.42, p=0.515, Figure 1f), and this relationship did not change over time (MSR*time: 257 χ^{2}_{1} =0.26, p=0.614). Similarly, a focal male's relatedness with the female did not predict fertilisation 258 probability (relatedness: χ^2_1 =0.64, p=0.423, Figure 1g), and this pattern did not change over time 259 (relatedness*time: χ^{2}_{1} =0.16, p=0.694). Finally, social status was not associated with a focal male's 260 probability of fertilisation (Elo score: χ^{2}_{1} =0.01, p=0.936, Figure 1h), and this pattern did not change over 261 time (Elo score*time: $\chi^2_1=0.31$, p=0.580). Male identity explained almost all the variance of random effects 262 (Figure S3). 263

265 4. Discussion

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266 Parker's intuition of sperm competition [1] kickstarted a new field of research, which has exploded over recent decades [8-10]. This effort has elucidated the independent role of numerous mechanisms including 267 268 physiological, genetic, behavioural and morphological traits in determining competitive fertilisation. 269 Despite this, we still have a poor understanding of how these mechanisms interact with each other to drive 270 dynamics of sperm competition in natural populations. This is particularly the case for the many taxa, such 271 as birds, with internal fertilisation, prolonged female sperm storage and iterated insemination and 272 fertilisation events, mostly due to the difficulty in linking mating to fertilisation and following temporal 273 patterns of ejaculate utilisation. In this study we combined detailed information on mating behaviour with 274 molecular parentage data from groups of red junglefowl to explain variation in the probability of 275 individual males fertilising individual eggs over successive days. We show that around 20% of the 276 variance is explained by information gleaned from mating behaviour data. The degree of female 277 polyandry has an intuitive negative impact on a male's probability of fertilisation, however the outcome of 278 sperm competition among males is modulated by four other factors.

279 First, a male is more likely to fertilise a female's eggs - especially those ovulated in the last days of 280 a trial - if he mates more frequently with her. Previous work had shown that post-copulatory sexual 281 selection favours males that remate more frequently with their partners in this population [61,62], but 282 there has been no attempt at quantifying temporal effects. Repeated matings with between partners are 283 often observed in birds and the significance of this behaviour has long been debated [93]. Because females 284 can continue to ovulate over successive days following an insemination, and because sperm are lost at a 285 constant rate from the female sperm storage tubules (SSTs), male birds may be selected to top up their 286 sperm representation in a female's SSTs by remating rather than inseminating a single large ejaculate. In 287 addition, we now show that this effect is progressively accentuated over successive days; multiple days of 288 remating are required for males to accumulate sufficiently more sperm than rivals within the female SSTs 289 in order to have an advantage in sperm competition. Positive post-copulatory sexual selection on remating 290 rates has also been demonstrated in some insects [64,66], although in these studies the advantage to 291 remating may be related to mating order, as males that mate more frequently with a female may be more 292 likely to mate last and displace the sperm previously inseminated by rivals. Thus, in polyandrous 293 populations characterised by a degree of sperm competition intensity [61,62], selection may favour males 294 that mate with fewer females, but are able to remate with these females more frequently [94].

Second, controlling for remating rates, we found an interaction between mating order and time;
such that mating last increases the fertilisation probability of males late in a trial. Establishing the effect of
mating order is critical to evaluate different models of sperm competition. A last male advantage is
commonly observed in a number of taxa [13]. In birds, different sperm competition models have been
explored [29,30,41]. First, sperm from different inseminations may stratify within the female's SSTs, with
sperm from later copulations "sitting" on top of sperm from earlier copulations. Second, a new

301 insemination may displace sperm of previous inseminations from the female SSTs. Finally, sperm from 302 earlier inseminations may be lost over time due to passive sperm loss from the SSTs [92,95]. All three 303 models predict some degree of last male advantage in sperm competition, but only the passive sperm loss 304 model predicts that this advantage increases with the time elapsed between competing inseminations 305 [92,95]. Despite early suggestions of sperm stratification in individual SSTs [96], analysis of artificial 306 insemination experiments in domestic fowl indicated that avian sperm competition may conform to 307 passive sperm loss [29,30,41,92,95]. A subsequent careful differential sperm staining experiment in 308 domestic fowl and domestic turkeys, Meleagris gallopavo, found no evidence of sperm stratification [97], 309 with sperm of rival inseminations largely segregated in different SSTs in both species, although some of 310 visual the patterns presented raised the possibility of sperm stratification when rival inseminations mixed 311 within individual SSTs (e.g. figure 2c). The sperm stratification model predicts that the first insemination 312 may experience a renewed increase in fertilisation probability later in the laying sequence, as the sperm 313 from subsequent inseminations are depleted allowing the sperm from the first insemination to leave the 314 SSTs. Our post-hoc analysis showed that the time lag between the last copulations of competing males 315 tended to increase over successive days of the trial, which is consistent with passive sperm loss but less so 316 with sperm stratification. This suggests that most males mate early with a female but only some manage to 317 remate with her later on in the trial. Mating later in the trial provides these males with a fertilisation 318 advantage as the sperm of previous competitors will have been depleted from a female's SSTs through 319 passive loss. Mating last in the first days of the trial does not convey the same advantage, instead our 320 results suggest that, early in the trial, mating first might be advantageous. This conclusion is also 321 supported by our attempts to model sperm loss directly, which yielded considerably better fits than simply 322 modelling remating rates in almost half of the models. Importantly, segregation of rival ejaculates across 323 different SSTs may be a common pattern observed in galliform [97] and passerine birds [98,99]. Random 324 occupation of different individual SSTs by rival ejaculates would likely result in an overall pattern of 325 sperm loss similar to that predicted under conditions of random sperm mixing within individual SSTs [6]. 326 Non-random segregation patterns however, might buffer or accentuate patterns of paternity share 327 predicted by the passive sperm loss model. For example, last male sperm precedence might be favoured if 328 sperm from the last insemination were preferentially stored in the SSTs closer to the influndibulum, which 329 mature later, resulting in a possible fertilising advantage [98]. The lack of remating by many males may 330 occur for two non-mutually exclusive reasons. i) It may take some time for the social hierarchies to 331 influence differential access to mating opportunities, and mating later in the trial may become increasingly 332 difficult for low-ranking males, as dominant males progressively monopolise females [60]. Consistent with 333 this, dominant males were more likely to fertilise eggs on the 8th day of the trial (see Analysing each day of 334 the trial separately in the electronic supplementary material). ii) Mating can be costly for males in these 335 groups [75], and only some males (e.g. younger and in better condition) may be able to maintain high 336 mating rates throughout the trial. These findings highlight the importance of accounting for the temporal 337 dynamics underpinning sperm competition in species with prolonged sperm storage.

338 Third, our results suggest that female resistance may impair the fertilisation probability of 339 successful mating attempts by compromising sperm transfer. Male fowl are on average larger than 340 females, and frequently coerce females into copulating [60], which can undermine female pre-copulatory 341 choice. Female responses to male harassment in fowl range from acquiescence to costly resistance [60]. 342 Previous work in feral domestic fowl has shown that female resistance reduces the probability that a 343 copulation attempt leads to successful sperm transfer [100]. Another possibility is that females are more 344 likely to eject sperm following resisted copulations. Previous studies have demonstrated that female fowl 345 eject sperm differentially [101,102], and the risk of sperm ejection increases over successive matings [101]. 346 Female resistance is common in birds and can be intense in some species [28,103], however its influence on 347 the reproductive outcome has seldom been quantified. In mallards, Anas platyrhynchos, female resistance 348 reduces the probability that a mating attempt leads to successful copulation [104], while experiments in 349 Japanese quails, Coturnix japonica, indicate that males that are more aggressive to females have a lower 350 fertilisation probability [105]. Early models of sperm competition ignored female behaviour. Our results 351 indicate that female behaviour is a key factor in determining the outcome of sperm competition and 352 paternity share, even in species characterised by widespread male sexual coercion. Importantly, we show 353 that the effect of female resistance decreased with female age, suggesting that older females are less 354 effective at dislodging harassing males, or are less able to bias the outcome of sperm competition against 355 their sperm (e.g., through differential sperm ejection [101,102]). Our results suggest that on the one hand, it 356 may be easier for males to coerce older partners, which may be less efficient at resisting, while on the 357 other, mating with older females may result in lower reproductive returns given their reduced fecundity 358 and faster sperm loss rates [74].

Fourth, older males have reduced fertilisation probability for individual eggs, confirming the overall effect of male age on male reproductive success previously reported for these trials [62]. In addition, we now show that this negative effect occurs constantly over the course of a trial, suggesting a consistent detrimental effect of ageing on male ability to compete in sperm competition. These findings are consistent with previous studies of domestic fowl showing that older males have lower sperm quality [72],

364 and fail to transfer sperm more often, or transfer smaller ejaculates, than younger males [70,71]. 365 Interestingly, several studies of extra-pair paternity in natural passerine populations have detected the 366 opposite effect, i.e. older males tend to cuckold younger males and sire more extra-pair offspring [106-109]. 367 This incongruence may reflect differences between fowl and passerine birds, between the polygynandrous 368 and socially monogamous mating systems, or between captive and natural populations. While the effect of 369 phylogenetic and mating system differences are harder to assess, it is likely that high extrinsic mortality 370 rates in natural populations mean that males seldom age beyond their prime, and male age in these 371 conditions may positively covary with experience and genetic quality. Low extrinsic mortality rates in 372 captive populations on the other hand, allow individuals to age beyond their prime, revealing a stronger 373 impact of reproductive ageing. A study of captive house sparrow, Passer domesticus, however, shows that 374 old males (>6 years) can deliver more sperm than young males [110], suggesting that captivity effects may 375 not be sufficient to explain differences in the role of male age in sperm competition between socially 376 monogamous passerines and polygynandrous galliforms. While male reproductive ageing has long been 377 recognised in several populations [111-113], its role in sperm competition dynamics has only recently 378 begun to emerge [114]. Our findings indicate that male age is a key determinant of paternity share in 379 sperm competition in age-structured bird populations and suggest that females could minimise fitness 380 costs associated with mating with older males by promoting sperm competition [112].

381 Finally, our study suggests that factors previously identified as important determinants of sperm 382 competition outcomes may play a more negligible role in freely-mating groups. A male's relatedness to a 383 female, his social status, and the state of his sperm reserves (measured as MSR) were not significant 384 predictors of his fertilisation probability. Individuals are expected to avoid reproducing with close 385 relatives to prevent inbreeding depression [115-117]. Theory predicts that asymmetries in reproductive 386 costs mean that, for intermediate levels of inbreeding depression, females should avoid fertilisation by 387 close relatives more than unrelated males [117]. In line with this, previous experimental studies in fowl 388 have shown that males are as likely to mate with full-sib sisters as with unrelated females [76], whereas 389 females may counteract this by favouring the sperm of unrelated males [76,77], a pattern that has been also 390 found in other taxa [20,118-124, but see 125-127]. While previous studies on post-copulatory inbreeding 391 avoidance have typically used highly controlled experiments with two rival males that differ markedly in 392 their relatedness to the female (e.g., unrelated versus full sibling), studies of natural populations are scarcer 393 (e.g., [123]). It is possible that in the freely interacting groups of the present study, inbreeding risk may be 394 reduced largely at a pre-copulatory stage by mating with less related partners. Additionally, post-395 copulatory inbreeding avoidance may be captured by other mechanisms, such as female resistance, 396 included in the models and/or may have a negligible independent effect on fertilisation probability 397 compared to other factors. Similarly, the lack of an effect of social status on male fertilisation success in the 398 present results also contrasts with previous studies suggesting that dominant male fowl are favoured in 399 post-copulatory sexual selection [61] and that females tend to retain more sperm from dominant males 400 [101,102]. One possibility is that the reproductive advantage associated with dominance in fowl and other 401 species [128,129] could be driven mostly by traits, such as younger male age, lower female resistance and 402 higher remating rates [61,62,66], which were directly included in our models, or pre-copulatory strategies 403 such as mate-guarding [130,131]. Alternatively, the benefits of social dominance may be cancelled out if 404 subordinate males produce ejaculates of higher fertilising efficiency [45,46]. Future studies should 405 disentangle potential positive and negative consequences of status for paternity share variation in 406 polyandrous populations. While our groups were not assembled systematically with respect to male age, 407 social competitive ability, and male-female relatedness, future studies could manipulate group 408 composition experimentally to resolve the causal role these factors in sperm competition, while 409 maintaining a more realistic setting. Exploring additional factors such as the role of sperm fertilising 410 efficiency (e.g. measures of sperm motility [45,46]) will also be important in improving our understanding 411 of sperm competition in bird populations.

412 Birds have provided a model system of vertebrate sperm competition. In studying sperm 413 competition in socially monogamous bird species, Birkhead [48] identified four predictors of the 414 probability of extrapair paternity: (1) the relative timing and success of insemination by competing males, 415 (2) the relative remating rates by competing males, (3) the duration of female sperm storage, and (4) 416 patterns of sperm precedence. Our study has provided evidence that all these factors are important in 417 sperm competition dynamics in polyandrous social flocks. We have provided corroborating evidence of 418 passive sperm loss under more natural conditions, and identified a role for both male and female processes 419 in determining the probability that competing males fertilise individual eggs. Crucially, our results show 420 that the temporal dependency of these effects should be taken into consideration when studying avian 421 sperm competition. Nevertheless, approximately 80% of the variance in probability of fertilisation 422 remained unexplained, demonstrating that much work is still needed in order to understand complex 423 patterns of sperm competition in nature. We hope this paper provides a methodological platform for 424 future studies. While the need for behavioural information to study sperm competition in wild birds has 425 long been recognised [36], this has been notoriously difficult to obtain. Recent advances in tracking 426 technologies however, are beginning to yield behavioural data of unprecedented high-resolution for wild

- 427 animals (e.g., [132]), and we foresee exciting opportunities to understand dynamics of sperm competition
- 428 and their eco-evolutionary significance in natural populations.
- 429

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Table 1. Comparison of selected binomial generalized linear mixed models (GLMMs), predicting the probability male red junglefowl, *Gallus gallus*, fertilise individual eggs, ranked according to AIC. For all models see Table S1.

Model ID	Fixed effects	К	AIC	ΔΑΙΟ	R^{2}_{m}	$R^2_{ m c}$
mod1	RR*T + MO*T + C + A + FR*FA	15	3621.33	0	0.1498	0.1959
mod13	RR*T + MO*T + C*T + A*T + FR*FA + SS*T + MSR*T + R*T	23	3633.24	11.92	0.1460	0.1917
mod24	С	6	3851.02	229.69	0.0550	0.1318
mod30	FR	6	3968.72	347.4	0.0205	0.1355
mod33	RR	6	3997.33	376.03	0.0112	0.1168
mod38	А	6	4016.99	395.66	0.0189	0.0907
mod39	SS	6	4040.27	418.95	0.0014	0.1097
mod41	MO	6	4041.31	419.98	0.0004	0.1089
mod43	R	6	4042.1	420.78	0.0002	0.1107
mod44	MSR	6	4042.24	420.91	0.0002	0.1076

A=Male age; C=Number of competitors; FA=Female age; FR=Female resistance; MO=Mating order; MSR=Male sperm reserves; R=Relatedness; RR=Remating rate, SS=Social status; T=Time; K=Number of parameters; AIC=Akaike information criterion; Δ AIC=with respect to mod1; R^2_m =Marginal R^2 ; R^2_c =Conditional R^2 .

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Table 2. Summary statistics and likelihood ratio tests results of the generalize	d linear
mixed-effects model (GLMM) predicting egg fertilisation success in multiple n	eplicate
groups of red junglefowl, Gallus gallus. Model contains all main effects and stat	tistically
significant interactions (mod5, Table S1).	697

	Egg fertilisation success				
Predictors	Estimate (odds ratio)	Confidence interval	LRT	<i>p</i> 698	
Intercept	30.09	12.19-74.25	-	_	
Remating Rate (RR)	.93	.61-1.42	-	-699	
Time (T)	.82	.7589	-	-	
Mating Order (MO)	.92	.80-1.05	-	700	
Female Resistance (FR)	.12	.0721	-	-700	
Female Age (FA)	.85	.70-1.02	-	-	
Number of Competitors (C)	.65	.6169	193.18	<.001	
Male Age (A)	.44	.3555	51.43	<.001	
Relatedness (R)	.88	.64-1.20	.64	.423	
Social Status (SS)	.99	.73-1.33	.01	.936	
Sperm reserves (MSR)	1.08	.86-1.36	.42	.515	
MO:T	1.08	1.01-1.16	13.17	<.001	
RR:T	1.04	1.02-1.06	5.72	.018	
FR:FA	1.23	1.02-1.49	4.88	.027	
Random Effects					
σ^2	3.29				
τ _{00 MaleID}	0.35				
$\tau_{00 \text{ FemaleID}}$	0				
τ ₀₀ femalegroupID	0				
$\tau_{00 \text{ groupID}}$	0				
Observations	4134				

Notes: LRT=Likelihood ratio test; σ^2 =residual variance; τ_{00} =random intercept variance.

702 Figure and table captions

703 Figure 1. Relationships between different predictors and the probability that competing males fertilise

individual eggs in semi-natural groups of red jungle fowl, *Gallus gallus*. Panels represent the marginal effects

of a predictor based on the generalised linear mixed model (GLMM) in Table 2. a) Relative remating rate and

its interaction with time in days, b) mating order and its interaction with time in days, c) relative female
 resistance and its interaction with female age in years, d) number of competing males, e) relative male age, f)

relative male sperm reserves (MSR), g) relatedness between a male-female pair, and h) relative male Elo score

709 (i.e., social status). Shaded areas represent the 95% confidence intervals. Colour coding in panels a-c

represents three arbitrary values of the interaction term ((a) and (b) time: day2, day5, day9; (c) female age: 1, 5,

711 7years old). Asterisks represent statistical significance (Table 2), such that * = p < 0.05, and *** = p < 0.001.

713 Figures



716 Figure 1

719 Additional Information

720 Ethics

Research was conducted according to United Kingdom home office legislation (Home office licenses 30/2418
and 30/2931) following approval by the Departmental Animal Welfare Ethical Review Body (AWERB).

724 Data Accessibility

725 We will deposit supporting data and R codes in Dryad Digital Repository on acceptance.

726 727 Authors' Contributions

R.C., G.C.M. and T.P. conceived the study. G.C.M. conducted the field work, R.C. analysed the data. L.G. S.,
 E.A.F. and D.S.R. performed molecular analyses for parentage assignment. Y.W. provided unpublished data

E.A.F. and D.S.R. performed molecular analyses for parentage assignment. Y.W. provided unpublished data
for Figure S1. R.C., T.P. and G.C.M. wrote the manuscript, and D.S.R. provided comments. All other authors
gave final approval for publication.

733734 Competing Interests

735 We have no competing interests.

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