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<https://doi.org/10.1093/beheco/arac124>

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Original Article

Opposite-sex associations are linked with annual fitness, but sociality is stable over lifetime

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Received 25 July 2022; revised 27 November 2022; editorial decision 30 November 2022; accepted 6 December 2022

Animal sociality, an individual's propensity to associate with others, has fitness consequences through mate choice, for example, directly, by increasing the pool of prospective partners, and indirectly through increased survival, and individuals benefit from both. Annually, fitness consequences are realized through increased mating success and subsequent fecundity. However, it remains unknown whether these consequences translate to lifetime fitness. Here, we quantified social associations and their link to fitness annually and over lifetime, using a multi-generational, genetic pedigree. We used social network analysis to calculate variables representing different aspects of an individual's sociality. Sociality showed high within-individual repeatability. We found that birds with more opposite-sex associates had higher annual fitness than those with fewer, but this did not translate to lifetime fitness. Instead, for lifetime fitness, we found evidence for stabilizing selection on opposite-sex sociality, and sociality in general, suggesting that reported benefits are only short-lived in a wild population, and that selection favors an average sociality.

Key words: de-lifing, genetic pedigree, lifetime reproductive success, long-term fitness, selection, sociality, social behavior, social network analysis

BACKGROUND

Some individuals are consistently more sociable than others, demonstrated by within-individual repeatability of social traits across vertebrate groups (Aplin et al. 2015; Thys et al. 2017; Dimitriadou et al. 2019; Plaza et al. 2019; Beck et al. 2020; Proops et al. 2021; Strickland et al. 2021). This variation in individual sociality is positively linked with fitness in some taxa (Silk 2007; Silk et al. 2009) and is therefore expected to be subject to selection (Krause and Ruxton 2002). Fitness is a relative measure of an individual's genetic contribution to the population in the next generation, and thus, can only be represented comprehensively and precisely by measures of traits spanning an organism's lifetime (Endler 1986; Blankenhorn 2010; Reid et al.

2019; Moiron et al. 2022). A comprehensive definition of fitness is fundamental to understand the evolutionary pressures that shape variation in sociality. In practice, however, many studies must rely on fitness correlates instead (e.g., number of broods, or survival, instead of genetic contribution). For example, in mammal societies, both variations in within- and between-sex affiliations (Archie et al. 2014) have been linked to lifetime fitness correlates, survival, and longevity (Cameron et al. 2019; Silk et al. 2003; Silk et al. 2010; Stanton and Mann 2012, but also see Thompson and Cords 2018). Whereas, in birds, the subject of this study, the use of fitness correlates (eggs laid, chicks fledged, within-year survival, etc.) are frequent over more precise fitness measures (Moiron et al. 2022), that require intensive field work over a long period of time.

Although some benefits are linked with sociality during the breeding period most of these tend to be short-term and contextual (Bebbington et al. 2017; Riehl and Strong 2018). Instead,

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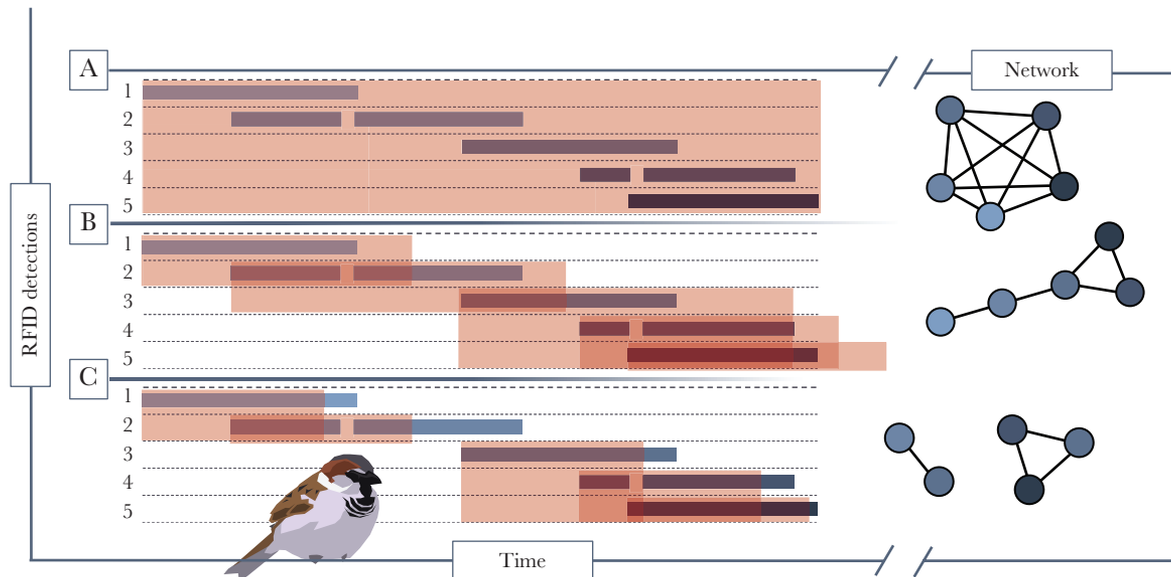


Figure 1.

Three versions of a simulated event (A, B, and C) show the interval over which five individuals (1–5, black/gray bars) spent at a resource over time (t), and the derived social networks from each approach: A = gambit of the group, which links all individuals in a discrete group equally; B = time-window overlap (by Δ_t), which links individuals who overlap at a resource; and C = arrival time (developed for this study), which links individuals who arrive together to a resource. Shaded boxes denote the time period during which individuals are considered to be associated, and overlaps represent an association: A, all individuals within a group; B, where they are physically present at the same time (shaded box), or shortly after they depart to account for birds which were present, but not currently being recorded by the antenna, in that case, over-lapping by Δ_t (shaded over-hanging box, typically a few seconds); or, C, where they arrive within Δ_t of each other, but the subsequent time spent at a feeder is irrelevant. However, note that the function of Δ_t differs between B and C; Where in B, Δ_t functions to detect when birds are in the same place but where one (or more) are not currently being detected by the antenna, in C the function is to link all individuals which arrive together while ignoring those already present at the resource, which has the potential to link two separate groups in A and B. In the case of C, an additional interval (Δ_t) is required to define when birds have left the resource, after which they can be recorded as arriving again.

benefits associated with reproduction are often linked with non-breeding sociality (Firth and Sheldon 2016; Kohn 2017; Maldonado-Chaparro et al. 2018; McDonald et al. 2020; Beck et al. 2020), when group cohesion is stronger (Silk et al. 2014; Kurvers 2020). Sociality may influence fitness in different ways, through benefits to reproductive success or increased survival, and so, the mechanism of selection acting on social traits may also differ. For example, communal foraging between socially associated individuals during the non-breeding period facilitates resource information transfer (Aplin et al. 2012; Firth et al. 2016; Hillemann et al. 2020) and reduces predation risk (Cresswell 1994; Cresswell and Quinn 2011; Sorato et al. 2012), increasing survival. However, these benefits may also incur costs associated with competition for resources and mate choice (Birkhead and Biggins 1987; Le Galliard et al. 2005; Forstmeier et al. 2011; Mayer and Pasinelli 2013; Grant and Grant 2019; but also see Lea et al. 2010). Sociality may also benefit individuals who hold more central social network positions or have access to opposite-sex associates, through enhanced mate choice (McDonald 2007; Oh and Badyaev 2010; Firth et al. 2018; Beck, Farine, and Kempenaers 2021). Although the association between annual fitness correlates and non-breeding sociality has been well described, testing how selection acts on social traits requires lifetime fitness measures, and remains unresolved.

With the recent development of tools to construct and analyze social networks (Wey et al. 2008; Farine and Whitehead 2015), the study of sociality has become popular among behavioral ecologists. Yet, to describe the association between sociality and fitness any potential study must first overcome two problems: 1) A

social association must be clearly defined relative to the behavior of the study system (Figure 1; Psorakis et al. 2012, 2015), and 2) to study the evolution of social behavior, precise measures of individual fitness must be quantified to test for correlation with a social trait—selection. Although annual fitness correlates are widely used, lifetime fitness is more precise and without the stochasticity of annual measures (Dobson et al. 2020; Alif et al. 2022) and thus, can better describe selection pressure acting on a trait (Endler 1986; Blankenhorn 2010; Reid et al. 2019). However, lifetime fitness requires wild animals to be monitored throughout their whole lives, and all breeding attempts, and the fates of their offspring, must be recorded to determine recruitment. All of these require a multi-generational, genetic pedigree (Kruuk 2004; Korsten et al. 2013).

Our study system, an island population of house sparrows *Passer domesticus* (hereafter, sparrow/s) where we monitor all individuals from birth to death, without capture bias (Simons et al. 2015), overcomes both problems. 1) We have sociality data from birds that are electronically registered visiting a feeder. Social centrality measures are repeatable across different timescales and contexts in this and other populations (Plaza et al. 2019). 2) We have lifetime recruitment data available, from a multi-generational genetic pedigree that, because our population is closed, meaning there is no movement of sparrows to or from the island, and that our study covers all sparrows on the island, we can use to compute precise annual and lifetime fitness estimates (Schroeder et al. 2015; Alif et al. 2022).

We tested predictions based on arguments presented above to understand the potential for selection on sociality: 1) We confirmed that the social traits we measured were meaningful by testing for

individual repeatability of sociality; 2) We tested the prediction that non-breeding sociality has fitness benefits, either driven by reproductive success through opposite-sex association or through increased survival through network centrality measures and is subject to selection.

METHODS

Study system

We used data from the Lundy sparrow system, a long-term study based on the island of Lundy (51.11N, 4.40W), ≈ 19 km off North Devon, UK, in the Bristol Channel. The sparrows on Lundy breed in nest boxes sited in groups around the only village on the island. The island is rodent-free and therefore the sparrows have no predators but for the occasional vagrant raptor. House sparrows are a model organism in behavioral ecology and evolution, and much is known about their biology, physiology, and life-history (Andersson 1994; Sánchez-Tójar et al. 2018). House sparrows are socially monogamous, but 25% of broods show they can be genetically promiscuous (Schroeder et al. 2016). On Lundy, they have on average 2–3 broods of 4–5 eggs per breeding season (Westneat et al. 2014). The sex ratio is stable, and the mean lifespan of recruits is three years (Alif et al. 2022). Although sparrows are territorial during the breeding season, during the non-breeding period they form gregarious groups that forage communally for seed and at supplementary bird feeders (Summer-Smith 1963), both of which are available year-round on Lundy.

Most sparrows were first captured, and tissue was sampled in nest boxes at their natal site during the breeding season (April to August) or using mist nets during the post-fledging period (Schroeder et al. 2011; Girndt et al. 2019). Tissue samples were either blood or mouth swabs and were stored in ethanol and refrigerated at 3°C prior to analysis. We genotyped sparrow DNA at <22 microsatellite loci suitable for parentage assignment in sparrows (Dawson et al. 2012). Using the genetic data, we assembled a near-complete genetic pedigree (Schroeder et al. 2015, 2016), which at the time of writing spans 20 years, 2000–2019, and 8379 individuals. We fitted all sparrows with a unique combination of a coded metal ring and three colored leg rings. We also provided each sparrow with a subcutaneous Passive Identification Transponder (PIT tag; TROVANID100: 11.5 \times 2.1 mm and 0.1 g), under the skin of the breast, which we have previously shown has no detrimental effect on subsequent fitness (for details see Schroeder et al. 2011). These tagged individuals were then recorded when they visited a custom-made 19.8 cm \times 19.8 cm Radio Frequency Identification antenna (RFID; DorsetID) mounted on a seed reservoir (for photo see Sánchez-Tójar et al. 2017; Brandl et al. 2019), positioned centrally within our study site. The feeder was open access, and explicitly not limited to a single bird feeding at one time, as is the case at hanging bird feeders (Youngblood 2019; Beck et al. 2020). Our feeder recorded visiting birds every day that the island generators were running (6 am–midnight, seven days a week).

SOCIAL CENTRALITY MEASURES

To quantify the sociality of individual sparrows, we calculated measures of social centrality (hereafter centrality measures) using presence data from the RFID antenna, collected during the non-breeding periods, November–January, of 2015/16 (15 weeks) and

2016/17 (13 weeks, hereafter referred to events). An association between birds observed from these data can reflect either, individuals that choose to maintain some social cohesion, which we consider to be a nonrandom association, or individuals without pre-established social cohesion who coincide in time and space, which we consider random mixing. These data record the presence of birds without distinguishing between these two types of association. Further, dyadic interactions can also be aggressive interactions, and our data cannot exclude these cases. Although in sparrows, dominance hierarchies are typically linear, there is no evidence for a correlation with reproductive fitness (Sánchez-Tójar 2018).

The “gambit of the group” is a common approach used to identify discrete groups among all associating individuals (Whitehead and Dufault 1999; Franks et al. 2010; Figure 1a). However, given the gregarious nature of sparrows and the high activity at our feeder, at which non-discrete groups of sparrows accumulated, the gambit of the group approach overestimated associations between individuals (Figure 1a; also see Ferreira et al. 2020). One solution to this would be to define a nonrandom association where two individuals overlapped by a defined period (Δ) at our bird feeder. However, in our system, again due to the near constant visitations, this resulted in linear network structures, for example, linking the first bird to the second, then the second to the third, and so on (Figure 1b). To account for the social behavior of sparrows, we derived a method to infer nonrandom associations that assumed that nonrandom social associations are established before they attend the feeder (suggested by Summer-Smith 1963). We, therefore, defined an association as two individuals that arrived to feed within 150 s (Δ) of each other. Here an arrival is defined as the (re)appearance of the individual at the feeder after being absent for a period of minimally >300 s (Δ). We defined that $\Delta_i = 150$ s was sufficient to detect and link individuals who arrive together in a group (see Figure 1c), and the resulting data better sampled nonrandom associations between individuals in our system, from watching sparrows in the field and on prerecorded footage, arriving at our feeder (Plaza et al. 2019).

From the resulting association matrices of the two events, 2015/16 and 2016/17, we built a series of weighted, non-directional, social networks (hereafter, network/s), where the vertices represent individual sparrows and interconnecting edges of their associations. First, we built individual networks for each of the 15 weeks in 2015/16, and 13 weeks in 2016/17, to estimate within-individual repeatability in centrality network metrics. This was to validate these centrality measures against the individual repeatability already demonstrated in this system using both RFID and video data (Plaza et al. 2019). Then, we built a network for each event to extract non-breeding sociality. Finally, we also built two bipartite networks from each event (sub-graphs), which only considered association strength between opposite-sex individuals.

From the first non-breeding networks we extracted three centrality measures representing different aspects of sociality for each individual using the “iGraph” R package (Csardi and Nepusz 2006): We selected centrality measures to reflect aspects of an individual’s social preference, following similar studies on sociality (McDonald 2007; Farine and Whitehead 2015; Beck et al. 2021). Degree represents the number of associates and may impact fitness through enhanced mate choice, where individuals position themselves alongside others of lesser quality (Oh and Badyaev 2010), or where same-sex associates benefit reproduction in cooperative breeding species (Bebbington et al. 2017; Riehl and Strong 2018). Strength represents the quality of those relationships and may influence the structure

and behavior of reproductive communities (Firth and Sheldon 2016; Culina et al. 2020). We calculated the strength following (Farine 2013), using the sum of dyadic Simple Ratio Indices (the association probability between a dyad, from 0, never associated, to 1, always associated), which we transformed to give a measure of net association quality (Equation 1, where S denotes strength, $d_{(i)}$ and sum $SRI_{(i)}$ are the degree and SRI of a given individual respectively and $N(V)$ is the number of vertices in a network)

$$S = d_{(i)} \frac{SRI_{(i)}}{N(V)} \quad (1)$$

We used eigenvector centrality (following McDonald 2007, hereafter centrality) to quantify the influence of an individual on all others within the network (Newman et al. 2004; Oh and Badyaev 2010); Finally, we extracted opposite-sex degree from the two bipartite sub-graphs, the number of opposite-sex associations, which we used to represent an individual's pool of potential reproductive partners, as a fourth measure of sociality (following Beck et al. 2021). Given the high density of sparrows visiting the feeder, and the frequency at which those birds were detected we did not threshold our networks (using only a sample of the birds attending an antenna, following Farine and Whitehead 2015) to maintain network structure, although only birds who arrived in a dyad (where degree > 0) were included in our networks.

FITNESS MEASURES

For each of the sparrows that survived to the following breeding period, we used the genetic pedigree to calculate two fitness measures, and for each of these, we calculated one annual measure and one across the lifetime of the sparrow. Both fitness measures are based on the number of recruits, and we have shown that they correlate well with reproductive value, thus representing fitness in an evolutionarily meaningful way (Alif et al. 2022). We defined recruits as offspring that survived and produced genetic offspring themselves.

- 1) *Recruits*. For the number of annual recruits, we summed individual recruits within the breeding year following the social events. We then again summed individual recruits across a lifetime, or up to 2020 as a measure of lifetime recruits. Note that the latter category only contained five sparrows that were still alive at the point of census, and as such, our recruitment data can be considered near complete. We excluded birds that did not survive to breed, and yearlings that had zero recruits.
- 2) *De-lifed fitness*. As a second fitness measure, we used de-lifed fitness (p_{ii} ; Equation 2), which estimates an individual's genetic contribution to the population (Coulson et al. 2006). De-lifed fitness is a retrospective measure of realized fitness, relative to the population each year, calculated by removing (de-lifing) an individual and its offspring from the pedigree and recalculating the resulting change in population growth.

$$p_{ii} = \frac{\xi_{i(i)} - w_{i(i)}}{N_i - 1} \quad (2)$$

Here, p_{ii} is the contribution of individual (i) to population growth during a specific period t . Further, $\xi_{i(i)}$ is a measure of individual performance, here the number of surviving offspring of individual i at the end of the breeding period t . We added a value of one if the individual i itself survived to the next breeding period $t + 1$. The population size at time t is N_t at the beginning of each breeding cycle (here April). To estimate the individual's contribution to

population growth, we use w_t , which represents the ratio of the population size at $t + 1$ to the population size at t . This de-lifed fitness p_{ii} is an annual value per individual, and we calculated it for all birds which produced at least one recruit. We then also summed p_{ii} within individuals as a lifetime de-lifed fitness measure, p_i .

INDIVIDUAL REPEATABILITY IN SOCIALITY

First, we validated our centrality measures by confirming that they were repeatable within-individuals between weeks. We modeled degree, strength, centrality, and opposite-sex degree respectively, as response variables each in four Gaussian linear mixed models against the intercept, and with bird identity modeled as a random effect to compensate for repeat identities between years. We then divided the variance explained by bird identity by the total phenotypic variance of the trait to quantify repeatability (see Nakagawa and Schielzeth 2010). We ran repeatability models using package MCMCglmm default parameters and priors—the models converged robustly and reliably.

SELECTION ON SOCIALITY—ANNUAL AND LIFETIME FITNESS

We quantified the association between centrality measures and fitness. As all four centrality measures are inherently correlated, we modeled each separately to avoid collinearity (Webster et al. 2020). For each centrality measure, we ran two models, with annual recruits and annual de-lifed fitness as the response variables. In the models with annual numbers of recruits as a response, we assumed a Poisson error distribution with a log link function, and in the models explaining de-lifed fitness we assumed a Gaussian error distribution. We mean-centered all centrality measures within each year, eliminating between-year differences, and modeled them as fixed covariates. We also added each sociality variable as a quadratic effect to test for stabilizing or disruptive selection where averages are favored over the extremes (Wolf et al. 2007). Bird identity was modeled as a random effect on the intercept to account for pseudo-replication, and cohort to account for environmental stochasticity. We modeled fixed effects for sex (male, 1 or female, 0) and age (in calendar years) and age as a quadratic effect, to account for variation in fitness as explained by demography (Schroeder et al. 2012). We added sex as an interaction term with age to account for the extra-pair behavior of older males (Gimrdt et al. 2018).

We modeled lifetime recruits and lifetime de-lifed fitness in the same way as the annual ones, but instead of age, we used lifespan or maximum age at the year 2020. Because each bird was only represented once in this dataset, we only modeled the cohort as a random effect.

We used Bayesian Markov Monte-Carlo simulations, using MCMCglmm (Hadfield 2010; R Core Team 2023), to run all models. For all models, we used inverse Wishart priors for random effects, and ran each over 343,000 iterations, with a burn-in of 3000 and a thinning interval of 200. We visually checked the posterior trace plots for all model outputs and ensured that autocorrelation was below 0.1 and effective sample sizes between 1000 and 2000. The fixed effects were considered statistically significant when the 95% credible interval (CI) of its posterior distribution did not span zero.

NULL MODELS AND DOMINANCE INTERACTIONS

We ran a node-permutation null model by shuffling the identities of birds visiting the feeder between existing arrival times in our

association matrices, thereby breaking any link between sociality and fitness, over 1000 randomized permutations (following Farine, 2017). We used these randomized association matrices to construct 1000 new networks and extracted the mode for our four centrality measures. We used these randomized centrality measures to re-run all fitness models. Finally, to exclude the possibility that dominance was interacting with our observed centrality measures, we tested for correlations between the centrality measures and dominance from videos recorded during the same period of our social network events. We represented individual dominance by calculating ELO ratings, based on antagonistic interactions at the bird feeder (for further details see Sánchez-Tójar 2018). We did not include the randomized centrality measures from our null models in these correlations.

RESULTS

The data consisted of 150 individual birds making 410,114 visits to the RFID feeder within our study period (mean = 2734 visits per bird, SD = 8116), across both events. Excluding birds that died prior to the start of our study or those that were ringed after, 160 tagged birds survived in our system in November 2015, and an additional 90 birds were tagged prior to the 2017 breeding period, although not all survived to sampling. After constructing the arrival networks, we identified 3783 associations between 118 PIT-tagged individuals during 2015/2016, and 874 associations between 69 individuals in 2016/2017. These networks contained 66.3% of 122 and 26.3% of 205 breeding birds in 2016 and 2017, respectively. Combined, we had 130 records for annual and lifetime fitness from 102 individuals, 33 were recorded in both years (for summary statistics see Table 1). Degree and opposite-sex degree are closely correlated, implying that those with more opposite-sex associates also tend to have more associates of either sex (Supplementary material Table 1)

INDIVIDUAL REPEATABILITY IN SOCIALITY

We confirmed individual repeatability by week in all four centrality measures between 15 weeks in 2015/16: Degree, $R = 0.29$ (0.15–0.39), Strength, $R = 0.22$ (0.10–0.32), Centrality, 015 (0.03–0.27), and Opposite-sex degree, 0.27 (0.13–0.4); and, 13 weeks in 2016/17: Degree, $R = 0.29$ (0.15–0.39), Strength, $R = 0.22$ (0.10–0.32), Centrality, 015 (0.03–0.27), and Opposite-sex degree, 0.27 (0.13–0.4) (Table 1).

Table 1.

Summary statistics for recruitment and centrality measures for individual house sparrows on Lundy Island during two non-breeding events (November–January 2015/2016 and 2016/2017). Each measure is given as mean, standard deviation, and sample size, including repeatability (R) and 95% CI for centrality measures.

Summary statistics of recruitment

Fitness Variable	Scale	2015/16		2016/17	
		Mean, SD	n (118)	Mean, SD	n (69)
Recruits	Annual	2.65	1.98	0.78	1.07
	Lifetime	5.34	3.76	4.15	3.86

Summary statistics and repeatability of centrality measures

Centrality measure	Mean, SD	R , 95CI	Mean, SD	R , 95CI
Degree	64.11, 32.03	0.29, 0.15–0.39	25.33, 17.37	0.35, 0.16–0.49
Strength	0.06, 0.05	0.22, 0.10–0.32	0.09, 0.13	0.33, 0.16–0.48
Centrality	0.35, 0.30	0.15, 0.03–0.27	0.25, 0.31	0.23, 0.10–0.36
Opposite-sex degree	32.00, 15.86	0.27, 0.13–0.4	12.18, 9.19	0.45, 0.24–0.61

SELECTION ON SOCIALITY—ANNUAL AND LIFETIME FITNESS

Opposite-sex degree had a statistically significant linear relationship with the number of annual recruits and annual de-lifed fitness. Strength and centrality had a negative quadratic association with annual recruitment (Figure 2 and Tables 2 and 3). Age and sex both also predicted annual recruits, with younger individuals and females recruiting more offspring. Age also positively predicted annual de-lifed fitness (Tables 2 and 3).

None of our centrality measures statistically significantly predicted lifetime recruitment (Supplementary Material Table 4), however, all four had a statistically significant negative quadratic relationship with lifetime de-lifed fitness (Figure 2).

NULL MODELS AND DOMINANCE INTERACTIONS

We found no link between fitness and sociality, nor any evidence of selection from our null models (Supplementary Material Figure 3). Likewise, dominance was not strongly correlated with any centrality measure, implying that our method of assigning associations based on arrival time, rather than shared space at a bird feeder, is unlikely to be influenced by dominance (Supplementary Material Table 1).

DISCUSSION

We found evidence for annual fitness benefits of sociality, where individuals with more opposite-sex associates had higher fitness in the breeding period than those with fewer, but this did not translate to lifetime fitness. For lifetime fitness, we found evidence for stabilizing selection on sociality, including opposite-sex degree, suggesting that such benefits are only short-lived, or contextual, in a wild population.

We constructed our networks by linking dyads of birds that arrived together to a bird feeder, but ignored the time that they spent there, to eliminate most random associations. Other studies have adopted similar approaches in high-density and open-feeder systems, or have considered the same implicit problems (Gomes et al. 2021). Ferreira et al. (2020), for example, identified flocks arriving, but then defined associations by spatial proximity at a series of feeder boxes. Further research could optimize our approach for other systems, either by refining the time after which an individual

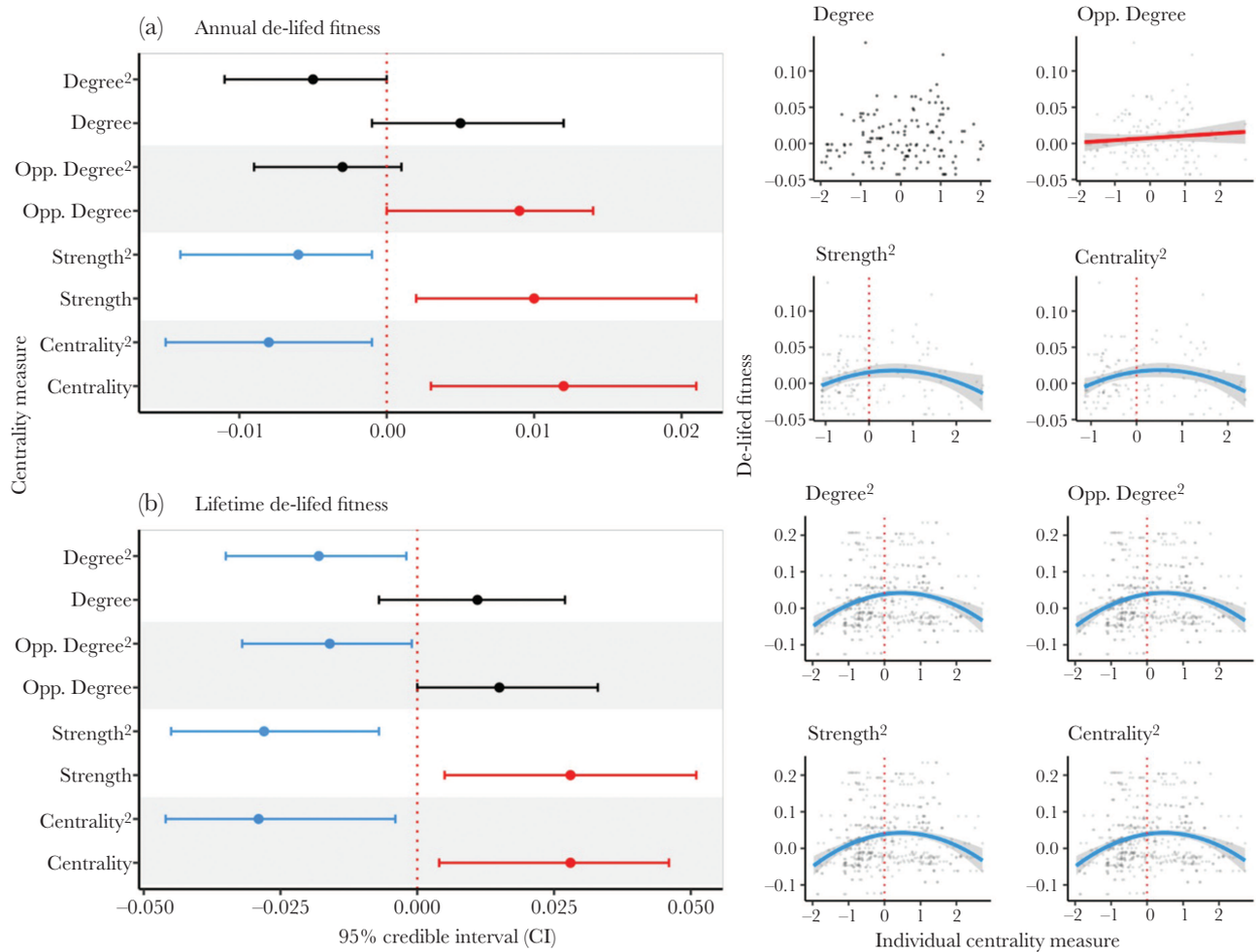


Figure 2.

De-lifed fitness as response variables against Centrality measures from 8 linear mixed models, at two scales, from the Lundy Island house sparrows, derived of 410,114 visits to an RFID baited feeder by 150 individuals: Explanatory variables for Annual de-lifed fitness (A); and Lifetime de-lifed fitness (B), where ² denotes a quadratic function, also shown in the four adjacent panels for A and B, and their 95% credible intervals (CIs). CIs are given as solid bars for each explanatory variable, where a solid point denotes the posterior mode. Black bars denote no effect on the response variable; red denotes a positive and blue, a negative, relationship with the response. In adjacent panels, quadratic functions of each response variable are presented in A and B (on the Y axis: A Centrality, Degree, Opp. Degree, Strength, and B Centrality, Degree, Opp. Degree, Strength). Blue curves represent a negative interaction with fitness measures (on the X axis). Measures with no effect are not shown in the figure (for colour figure refer to online version). We found no link to sex, and age was also subject to stabilizing selection, (given in [Supplementary Material Table 4](#)).

is determined to have left the feeder (Δ_i), or similarly, the time it takes for all members of a group to interact with the feeder upon arrival (Δ_j). Further work may also consider defining associations only where a dyad visits together more often than would be expected by chance but doing so must also consider some method of retaining peripheral associations. Although RFID systems sample sociality well at a feeder, we cannot be sure that sociality traits are maintained in other contexts—future works might consider tracking social behavior across time and space (e.g., see [Ripperger and Carter 2021](#)).

Where previous studies on wild birds have suggested links between aspects of sociality and annual reproductive success (e.g., see [Firth and Sheldon 2016](#); [Kohn 2017](#); [Beck, Farine, and Kempenaers 2020](#)), we were also able to use lifetime measures, which better reflect the genetic contribution of the individual to population growth, and thus, fitness. This also allowed us to also

describe how selection acts upon sociality across the population. We found that sociality had little influence on fitness at the annual scale, apart from for opposite-sex association, which was linked to increased recruitment and de-lifed fitness. Our study corroborates that annual fitness benefits described elsewhere, particularly regarding mate choice ([Oh and Badyaev 2010](#); [Beck et al. 2020, 2021](#)) directly translate into increased annual fitness. At the lifetime scale, our study also provides some insight into the evolution of social behaviors, which we found to be maintained at the population average through stabilizing selection. We are, therefore, to the best of our knowledge, the first study to link sociality with lifetime fitness benefits in a wild bird (but see [Formica et al. 2021](#); [Philon and Blumstein 2022](#)). Our results may also suggest a mechanism for selection on sociality through enhanced mate choice, but the impact on survival was difficult to determine in this study. Sociality is predicted to increase survival through reduced predation risk or

Table 2.

Annual recruitment model outputs from GLMMs for each of our four centrality measures (Degree, Strength, Centrality, and Opposite-sex degree), derived of 410,114 visits to an RFID baited feeder by 150 individuals. Centrality measure of house sparrows on Lundy Island, modeled against annual recruits. We inferred significance where the 95% CI did not span zero, positive effects on the response variable are highlighted in red, and negative in blue.

Annual recruits	Posterior mode 95% credible intervals			
	Degree	Strength	Centrality	Opposite-sex degree
<i>Fixed terms</i>				
(Intercept)	3.49 1.46–6.11	3.69 1.58–6.46	4.19 1.60–6.31	3.78 1.58–6.50
Centrality measure	0.17 -0.01–0.34	0.22 -0.03–0.44	0.26 -0.01–0.45	0.25 0.05–0.41
Centrality measure ²	-0.1 -0.26–0.06	-0.12 -0.32–0.07	-0.16 -0.34–0.08	-0.04 -0.16–0.10
Sex (male)	-0.67 -1.41–0.05	-0.7 -1.46–0	-0.74 -1.48–0	-0.57 -1.47–0
Age	-1.14 -2.03–0.30	-1.15 -2.04–0.28	-1.27 -1.99–0.26	-1.07 -2.00–0.26
Age ²	0.05 -0.07–0.17	0.04 -0.08–0.17	0.06 -0.08–0.16	0.03 -0.07–0.17
Age*Sex (male)	0.21 -0.01–0.48	0.22 -0.02–0.46	0.28 -0.02–0.49	0.21 0–0.49
<i>Random terms</i>				
*Bird ID	0 0–0.13	0 0–0.12	0 0–0.12	0 0–0.13
*Cohort	1.88 0.25–15.69	1.5 0.19–15.57	2.09 0–15.89	2.29 0.11–16.90
Residuals	0 0–0.27	0 0–0.28	0 0–0.29	0 0–0.24

Table 3.

Lifetime recruitment model outputs from GLMMs for each of our four centrality measures (Degree, Strength, Centrality, and Opposite-sex degree), derived of 410,114 visits to an RFID baited feeder by 150 individuals. Centrality measure of house sparrows on Lundy Island, modeled against lifetime recruits. We inferred significance where the 95% CI do not span zero, positive effects on the response variable are highlighted in red, and negative in blue. († Age in lifetime models denotes either lifespan or age in 2020, whichever is greatest)

Lifetime recruits	Posterior mode, 95% credible intervals (lower – upper)			
	Degree	Strength	Centrality	Opposite-sex degree
<i>Fixed term</i>				
(Intercept)	0.85 0–1.93	0.85 0.02–1.95	0.99 -0.06–1.87	0.8 -0.03–1.83
Centrality measure	0.1 -0–0.25	0.1 -0.09–0.3	0.12 -0.07–0.31	0.14 -0.05–0.25
Centrality measure ²	-0.05 -0.19–0.11	-0.01 -0.24–0.11	-0.07 -0.24–0.11	-0.03 -0.16–0.12
Sex (male)	0.33 -0.35–0.99	0.28 -0.36–0.91	0.28 -0.33–0.98	0.48 -0.24–1.03
†Age	0.1 0–0.23	0.11 0–0.24	0.09 0.01–0.25	0.12 0–0.24
†Age*Sex (male)	-0.06 -0.20–0.07	-0.07 -0.19–0.07	-0.07 -0.2–0.06	-0.06 -0.21–0.04
<i>Random effects</i>				
*Cohort	NA 0.07–2.78	0.47 0.07–2.76	0.45 0.06–3	0.39 0.10–2.77
Residuals	NA 0–0.27	0.12 0.01–0.29	0.11 0.02–0.28	0.14 0–0.26

information transfer (Sorato et al. 2012; Hillemann et al 2020), but we found no evidence to suggest that either was selected for, through higher centrality, in our analyses.

Stabilizing selection in this case may be driven by factors such as high mate fidelity or changing sociality with age (Oh and Badyaev

2010; Albery et al. 2022), removing the need to constantly maintain opposite-sex associations over lifetime while maintaining individual fitness. However, those opposite-sex associations may also be beneficial in an extra-pair context from the male perspective (Beck et al. 2020) and require further research.

Our centrality measures were associated with lifetime, but not with annual de-lifed fitness, and only opposite-sex degree was associated with recruitment at the annual scale. We found no relationship between social centrality and dominance in our study using arrival time to define sociality, but aggressive interactions are probably also reduced over the non-breeding period (Summer-Smith 1963). None of our centrality metrics were linked with recruitment at the lifetime scales. Overall, de-lifed fitness better represents fitness as it is a relative measure of the contribution to population growth (Alif et al. 2022). The number of recruits, while intuitively appealing, is not relative, and in good years, more birds may have a higher number of recruits, while in poor years, having one recruit may be an achievement. As such, this measure is not always comparable between years and may explain our results. Further, recruitment is also dependent on parental effects and relationships within the breeding season, which were not quantified here, although they have been suggested elsewhere (Bebbington et al. 2017; Riehl and Strong 2018), whereas de-lifed fitness also captures long-term survival. We found that older males recruited more offspring, likely by virtue of older males siring more extra-pair offspring (Girndt et al. 2018). Likewise, younger birds had lower annual de-lifed fitness, because younger birds had not recruited any offspring in previous years that would contribute to their current de-lifed fitness.

In conclusion, we suggest a link between opposite-sex association and reproductive success at the annual scale, suggesting a mechanism for selection to shape social behavior. At the lifetime scale, we suggest that selection on sociality is stable, suggesting greater fitness for those at the population mean, in a wild population of passerine birds.

SUPPLEMENTARY MATERIAL

Supplementary material can be found at <http://www.beheco.oxfordjournals.org/>

The authors would like to thank the Lundy Landmark trust and the Lundy Field Society for their ongoing support, particularly Dean Jones, Rosie Ellis and Tom Carr. The authors would also thank two anonymous reviewers for their helpful feedback.

FUNDING

This research was supported by the QMEE CDT, funded by NERC grant number NE/P012345/1 (JD), a fellowship from the Volkswagen Foundation (JS), a grant from the German Research Foundation: Deutsche Forschungsgemeinschaft (JS), a European Research Council grant, CIG PCIG12-GA-2012-333096 (JS), and by NERC grant NE/J024597/1 (TB).

DATA AVAILABILITY

Analyses reported in this article can be reproduced using the data provided by Dunning et al. (2022).

Handling Editor: Amanda Ridley

REFERENCES

- Albery GF, Clutton-Brock TH, Morris A, Morris S, Pemberton JM, Nussey DH, Firth JA. 2022. Ageing red deer alter their spatial behaviour and become less social. *Nat Ecol Evol.* 6(8):1231–1238.
- Alif Z, Dunning J, Chik HYJ, Burke T, Schroeder J. 2022. What is the best fitness measure in wild populations? A case study on the power of short-term fitness proxies to predict reproductive value. *PLoS One.* 17(4):e0260905. doi:10.1371/journal.pone.0260905.
- Andersson, M. 1994. Sexual selection. Princeton University Press, New Jersey.
- Aplin LM, Farine DR, Morland-Ferron J, Sheldon BC. 2012. Social networks predict patch discovery in a wild population of songbirds. *Proc R Soc B.* 279(1745):4199–4205. doi:10.1098/rspb.2012.1591.
- Aplin LM, Firth JA, Farine DR, Voelkl B, Crates RA, Culina A, Garroway CJ, Hinde CA, Kidd LR, Psorakis I, et al. 2015. Consistent individual differences in the social phenotypes of wild great tits, *Parus major*. *Anim Behav.* 108:117–127. doi:10.1016/j.anbehav.2015.07.016.
- Archie EA, Tung J, Clark M, Altmann J, Alberts SC. 2014. Social affiliation matters: both same-sex and opposite-sex relationships predict survival in wild female baboons. *Proc R Soc B.* 281(1793):20141261. doi:10.1098/rspb.2014.1261.
- Bebbington K, Kingma SA, Fairfield EA, Dugdale HL, Komdeur J, Spurgin LG, Richardson DS. 2017. Kinship and familiarity mitigate costs of social conflict between *Seychelles warbler* neighbors. *PNAS* 114(43):E9036–E9045. doi:10.1073/pnas.1704350114.
- Beck KB, Farine DR, Kempnaers B. 2020. Winter associations predict social and extra-pair mating patterns in a wild songbird. *Proc R Soc B.* 287(1921):20192606. doi:10.1098/rspb.2019.2606.
- Beck KB, Farine DR, Kempnaers B. 2021. Social network position predicts male mating success in a small passerine. *Behav Ecol.* 32(5):856–864. doi:10.1093/beheco/abab034.
- Beck KB, Valcu M, Kempnaers B. 2020. Analysis of within-individual variation in extrapair paternity in blue tits (*Cyanistes caeruleus*) shows low repeatability and little effect of changes in neighborhood. *Behav Ecol.* 31(6):1303–1315. doi:10.1093/beheco/araa069.
- Birkhead TR, Biggins JD. 1987. Reproductive synchrony and extra-pair copulation in birds. *Ethology.* 74(4):320–334. doi:10.1111/j.1439-0310.1987.tb00942.x.
- Blankenhorn, W. 2010. The quantitative study of sexual and natural selection. In: Kappeler P, editor. *Animal behaviour: evolution and mechanisms*. Germany: Springer.
- Brandl HB, Farine DR, Funghi C, Schuett W, Griffith SC. 2019. Early-life social environment predicts social network position in wild zebra finches. *Proc R Soc B.* 286:20182579–20182579. doi:10.1098/rspb.2018.2579.
- Cameron EZ, Setsaas TH, Linklater WL. 2019. Social bonds between unrelated females increase reproductive success in feral horses. *PNAS.* 106(33):13850–13853. doi:10.1073/pnas.0900639106.
- Coulson T, Benton TG, Lundberg P, Dall SRX, Kendall BE, Gaillard JM. 2006. Estimating individual contributions to population growth: evolutionary fitness in ecological time. *Proc R Soc B.* 273(1586):547–555. doi:10.1098/rspb.2005.3357.
- Cresswell W. 1994. Flocking is an effective anti-predation strategy in redshanks, *Tringa totanus*. *Anim Behav.* 47(2):433–442. doi:10.1006/anbe.1994.1057.
- Cresswell W, Quinn JL. 2011. Predicting the optimal prey group size from predator hunting behaviour. *J Anim Ecol.* 80(2):310–319. doi:10.1111/j.1365-2656.2010.01775.x.
- Csardi G, Nepusz T. 2006. The igraph software package for complex network research. *Inter J Complex Syst.* 1695(5):1–9.
- Culina A, Firth JA, Hinde CA. 2020. Familiarity breeds success: pairs that meet earlier experience increased breeding performance in a wild bird population. *Proc R Soc B* 287(1941):20201554.
- Dawson DA, Horsburgh GJ, Krupa AP, Stewart RK, Skjelseth S, Jensen H, Ball AD, Spurgin LG, Mannarelli ME, Nakagawa S, et al. 2012. Microsatellite resources for *Passeridae* species: A predicted microsatellite map of the house sparrow *Passer domesticus*. *Mol Ecol Res.* 12(3):501–523. doi:10.1111/j.1755-0998.2012.03115.x.
- Dimitriadou S, Croft DP, Darden SK. 2019. Divergence in social traits in *Trinidadian guppies* selectively bred for high and low leadership in a cooperative context. *Sci Rep.* 9(17194). doi:10.1038/s41598-019-53748-4.
- Dobson F, J Murie, V Viblanc. 2020. Fitness estimations for ecological studies: an evaluation in Columbian ground squirrels. *Front Ecol Evol* 8. doi: 10.3389/fevo.2020.00216
- Dunning J, T Burke, AHH Chan, HYJ Chik, TS Evans, J Schroeder. 2022. Opposite-sex associations are linked with annual fitness, but sociality is stable over lifetime. *Behav Ecol.* <https://doi.org/10.5061/dryad.z08kprhd>
- Endler JA. 1986. *Natural selection in the wild*. New Jersey: Princeton University Press.
- Farine DR. 2013. Animal social network inference and permutations for ecologists in R using asnipe. *Meth Ecol Evol.* 4(12):1187–1194. doi:10.1111/2041-210X.12121.

- Farine DR. 2017. A guide to null models for animal social network analysis. *Meth Ecol Evol.* 8(10):1309–1320. doi:10.1111/2041-210X.12772.
- Farine DR, Whitehead H. 2015. Constructing, conducting and interpreting animal social network analysis. *J Anim Ecol.* 84(5):1144–1163. doi:10.1111/1365-2656.12418.
- Ferreira AC, Covas R, Silva LR, Esteves SC, Duarte IF, Fortuna R, Theron F, Doutrelant C, Farine DR. 2020. How to make methodological decisions when inferring social networks. *Ecol Evol.* 10(17):9132–9143. doi:10.1002/ecc3.6568.
- Firth JA, Sheldon BC. 2016. Social carry-over effects underpin trans-seasonally linked structure in a wild bird population. *Ecol Lett.* 19(11):1324–1332. doi:10.1111/ele.12669.
- Firth JA, Sheldon BC, Farine DR. 2016. Pathways of information transmission among wild songbirds follow experimentally imposed changes in social foraging structure. *Biol Lett.* 12(6):20160144. doi:10.1098/rsbl.2016.0144.
- Firth JA, Cole EF, Ioannou CC, Quinn JL, Aplin LM, Culina A, McMahon K, Sheldon BC. 2018. Personality shapes pair bonding in a wild bird social system. *Nat Ecol Evol.* 2(11):1696–1699. doi:10.1038/s41559-018-0670-8.
- Formica V, Donald H, Marti H, Iregebay Z, Brodie III E. 2021. Social network position experiences more variable selection than weaponry in wild subpopulations of forked fungus beetles. *J Anim Ecol.* 90(1):168–182. doi:10.1111/1365-2656.13322.
- Forstmeier W, Martin K, Bolund E, Schielzeth H, Kempenaers B. 2011. Female extrapair mating behavior can evolve via indirect selection on males. *PNAS.* 108(26):10608–10613. doi:10.1073/pnas.1103195108.
- Franks DW, Ruxton GD, James R. 2010. Sampling animal association networks with the gambit of the group. *Behav Ecol Sociobiol.* 64(3):493–503. doi:10.1007/s00265-009-0865-8.
- Girndt A, Chng CWT, Burke T, Schroeder J. 2018. Male age is associated with extra pair paternity, but not with extra-pair mating behaviour. *Sci Rep.* 8:8378. doi:10.1038/s41598-018-26649-1.
- Girndt A, Cockburn G, Sánchez-Tójar A, Hertel M, Burke T, Schroeder J. 2019. Male age and its association with reproductive traits in captive and wild house sparrows. *J Evol Biol.* 32(12):1431–1443. doi:10.1111/jeb.13542.
- Gomes ACR, Boogert NJ, Cardoso GC. 2021. Network structure and the optimization of proximity-based association criteria. *Meth Ecol Evol.* 12(1):88–100. doi:10.1111/2041-210X.13387.
- Grant PR, Grant BR. 2019. Adult sex ratio influences mate choice in Darwin's finches. *PNAS.* 116(25):12372–12382. doi:10.1073/pnas.1903838116.
- Hadfield JD. 2010. MCMC methods for multi-response generalized linear mixed models: the MCMCglmm R package. *J Stat Softw.* 33(2):1–22. doi:10.18637/jss.v033.i02.
- Hillemann F, Cole EF, Sheldon BC, Farine DR. 2020. Information use in foraging flocks of songbirds: no evidence for social transmission of patch quality. *Anim Behav.* 165:35–41. doi:10.1016/j.anbehav.2020.04.024.
- Kohn GM. 2017. Friends give benefits: autumn social familiarity preferences predict reproductive output. *Anim Behav.* 132(201):201–208. doi:10.1016/j.anbehav.2017.08.013.
- Korsten P, Van Overveld T, Adriaensens F, Matthysen E. 2013. Genetic integration of local dispersal and exploratory behaviour in a wild bird. *Nat Commun.* 4:2362. doi:10.1038/ncomms3362.
- Krause JU, Ruxton GD. 2002. *Living in groups: Oxford series in ecology and evolution.* Oxford: Oxford University Press.
- Kruuk LEB. 2004. Estimating genetic parameters in natural populations using the “animal model”. *Proc R Soc B.* 359(1446):873–890. doi:10.1098/rstb.2003.1437.
- Kurvers RHJM, Prox L, Farine DR, Jongeling C, Snijders L. 2020. Season-specific carryover of early life associations in a monogamous bird species. *Anim Behav.* 164:25–37. doi:10.1016/j.anbehav.2020.03.016.
- Le Galliard JF, Fitz PS, Ferrière R, Clobert J. 2005. Sex ratio bias, male aggression, and population collapse in lizards. *PNAS.* 102(50):18231–18236.
- Lea AJ, Blumstein DT, Wey TW, Martin JGA. 2010. Heritable victimization and the benefits of agonistic relationships. *PNAS.* 107(50):21587–21592. doi:10.1073/pnas.1009832107.
- Maldonado-Chaparro AA, Montiglio PO, Forstmeier W, Kempenaers B, Farine DR. 2018. Linking the fine-scale social environment to mating decisions: a future direction for the study of extra-pair paternity. *Biol Rev.* 93(3):1558–1577. doi:10.1111/brv.12408.
- Mayer C, Pasinelli G. 2013. New support for an old hypothesis: density affects extra-pair paternity. *Ecol Evol.* 3(3):694–705. doi:10.1002/ecc3.489.
- McDonald DB. 2007. Predicting fate from early connectivity in a social network. *PNAS* 104(26):10910–10914. doi:10.1073/pnas.0701159104.
- McDonald GC, Engel N, Ratão SS, Székely T, Kosztolányi A. 2020. The impact of social structure on breeding strategies in an island bird. *Sci Rep.* 10(1):13872. doi:10.1038/s41598-020-70595-w.
- Moiron M, Charmanier A, Bouwhuis S. 2022. The quantitative genetics of fitness in a wild seabird. *Evolution.* 76(7). doi:10.1111/evo.14516.
- Nakagawa S, Schielzeth H. 2010. Repeatability for Gaussian and non-Gaussian data: a practical guide for biologists. *Biol Rev.* 85(4):935–956. doi:10.1111/J.1469-185x.2010.00141.X.
- Newman ME, Girvan M. 2004. Finding and evaluating community structure in networks. *Phys Rev E.* 69(2):026113.
- Oh KP, Badyaev AV. 2010. Structure of social networks in a passerine bird: consequences for sexual selection and the evolution of mating strategies. *Am Nat.* 176(3):E80–E89. doi:10.1086/655216.
- Philson CS, Blumstein DT. 2022. Group social structure has limited impact on reproductive success in a wild mammal. *Behav Ecol. arXiv:2202.00102.* doi:10.1093/beheco/arc102.
- Plaza M, Burke T, Cox T, Carroll AF, Girndt A, Halford G, Martin DA, et al. 2019. Social network node-based metrics can function as proxies for animal personality traits. *J Evol Biol.* 33(11):1634–1642. doi:10.32942/OSEIO/YVQ9D.
- Proops L, Troisi CA, Kleinhappel TK, Romero T. 2021. Non-random associations in group housed rats (*Rattus norvegicus*). *Sci Rep.* 11(1):15349. doi:10.1038/s41598-021-94608-4.
- Psorakis I, Voelkl B, Garroway CJ, Radersma R, Aplin LM, Crates RA, Culina A, Farine DR, Firth JA, Hinde CA, et al. 2015. Inferring social structure from temporal data. *Behav Ecol Sociobiol.* 69(857):857–866. doi:10.1007/s00265-015-1906-0.
- Psorakis I, Roberts SJ, Rezek I, Sheldon BC. 2012. Inferring social network structure in ecological systems from spatiotemporal data streams. *J R Soc Interface.* 9(76):3055–3066. doi:10.1098/rsif.2012.0223.
- R Core Team. 2023. R: a language and environment for statistical computing. Vienna, Austria: R Foundation for Statistical Computing. <http://www.R-project.org/>.
- Reid JM, Nietlisbach P, Wolak ME, Keller LF, Arcece P. 2019. Individuals expected genetic contributions to future generations, reproductive value, and short-term metrics of fitness in free-living song sparrows (*Melospiza melodia*). *Ecol Lett.* 3(3):271–285. doi:10.1002/eel3.118.
- Riehl C, Strong MJ. 2018. Stable social relationships between unrelated females increase individual fitness in a cooperative bird. *Proc R Soc B.* 285(20180130):20180130. doi:10.1098/rspb.2018.0130.
- Ripperger SP, Carter GC. 2021. Social foraging in vampire bats is predicted by long-term cooperative relationships. *PLoS Biol.* 19(9):e3001366.
- Sánchez-Tójar A. 2018. The evolution of social dominance in house sparrows. <http://nbn-resolving.de/urn:nbn:de:bsz:352-2-985104uzuvk43>. Accessed June 2022.
- Sánchez-Tójar A, Winney I, Girndt A, Simons MJP, Nakagawa S, Burke T, Schroeder J. 2017. Winter territory prospecting is associated with life-history stage but not activity in a passerine. *J Avian Biol.* 48(3):407–416. doi:10.1111/jav.01055.
- Sánchez-Tójar A, Nakagawa S, Sánchez-Fortún M, Martin DA, Ramani S, Girndt A, Bókony V, Kempenaers B, Liker A, Westneat DF, et al. 2018. Meta-analysis challenges a textbook example of status signalling and demonstrates publication bias. *ELife* 7:e37385. doi:10.7554/eLife.37385.
- Schroeder J, Cleasby IR, Nakagawa S, Ockendon N, Burke T. 2011. No evidence for adverse effects on fitness of fitting passive integrated transponders (PITs) in wild house sparrows *Passer domesticus*. *J Avian Biol.* 42(3):271–275. doi:10.1111/j.1600-048X.2010.05271.x.
- Schroeder J, Nakagawa S, Cleasby IR, Burke T. 2012. Passerine birds breeding under chronic noise experience reduced fitness. *PLoS One.* 7(7):e39200. doi:10.1371/journal.pone.0039200.
- Schroeder J, Nakagawa S, Rees M, Mannarelli ME, Burke T. 2015. Reduced fitness in progeny from old parents in a natural population. *PNAS* 112(13):4021–4025. doi:10.1073/pnas.1422715112.
- Schroeder J, Hsu YH, Winney I, Simons M, Nakagawa S, Burke T. 2016. Predictably philandering females prompt poor paternal provisioning. *Am Nat.* 180(2):219–230. doi:10.1086/687243.
- Silk JB. 2007. The adaptive value of sociality in mammalian groups. *Philos Trans R Soc Biol Sci.* 362(1480):539–559. doi:10.1098/rstb.2006.1994.

- Silk JB, Beehner JC, Bergman TJ, Crockford C, Engh AL, Moscovice LR, Wittig RM, Seyfarth RM, Cheney DL. 2010. Strong and consistent social bonds enhance the longevity of female baboons. *Curr Biol.* 20(15):1359–1361. doi:10.1016/j.cub.2010.05.067.
- Silk JB, Beehner JC, Bergman TJ, Crockford C, Engh AL, Moscovice LR, Wittig RM, Seyfarth RM, Cheney DL. 2009. The benefits of social capital: close social bonds among female baboons enhance offspring survival. *Proc R Soc B.* 276(1670):3099–3104. doi:10.1098/rspb.2009.0681.
- Silk JB, Alberts SC, Altmann J. 2003. Social bonds of female baboons enhance infant survival. *Science* 302(5648):1231–1234. doi:10.1126/science.1088474.
- Silk MJ, Croft DP, Tregenza T, Bearhop S. 2014. The importance of fission-fusion social group dynamics in birds. *Ibis.* 156(4):701–715. doi:10.1111/ibi.12191.
- Simons MJP, Winney I, Nakagawa S, Burke T, Schroeder J. 2015. Limited catching bias in a wild population of birds with near-complete census information. *Ecol Evol.* 5(16):3500–3506. doi:10.1002/ece3.1623.
- Sorato E, Gullet PR, Griffith SC, Russell AF. 2012. Effects of predation risk on foraging behaviour and group size: adaptations in a social cooperative species. *Anim Behav.* 84(4):823–834.
- Stanton MA, Mann J. 2012. Early social networks predict survival in wild bottlenose dolphins. *PLoS One.* 7(10):e47508E47508. doi:10.1371/journal.pone.0047508.
- Strickland K, Mitchell DJ, Delmé C, Frère CH. 2021. Repeatability and heritability of social reaction norms in a wild agamid lizard. *Evolution* 75(8):1953–1965. doi:10.1111/evo.14298.
- Summer-Smith, D. 1963. *The house sparrow*. UK: Collins.
- Thompson NA, Cords M. 2018. Stronger social bonds do not always predict greater longevity in a gregarious primate. *Ecol. Evol.* 8(3):1604–1614. doi:10.1002/ece3.3781.
- Thys B, Eens M, Aerts S, Delory A, Iserbyt A, Pinxten R. 2017. Exploration and sociability in a highly gregarious bird are repeatable across seasons and in the long-term but are unrelated. *Anim Behav.* 123:339–348. doi:10.1016/j.anbehav.2016.11.014.
- Webber QMR, Schneider DC, Vander Wal E. 2020. Is less more? A commentary on the practice of “metric hacking” in animal social network analysis. *Anim Behav.* 168:109–120. doi:10.1016/j.anbehav.2020.08.011.
- Westneat DF, Bókony V, Burke T, Chastel O, Jensen H, Kvalnes T, Lendvai Z, Liker A, Mock D, Schroeder J, et al. 2014. Multiple aspects of plasticity in clutch size vary among populations of a globally distributed songbird. *J Anim Ecol.* 83(4):876–887. doi:10.1111/1365-2656.12191.
- Wey T, Blumstein DT, Shen W, Jordán F. 2008. Social network analysis of animal behaviour: a promising tool for the study of sociality. *Anim Behav.* 75(2):333–344. doi:10.1016/j.anbehav.2007.06.020.
- Whitehead H, Dufault S. 1999. Techniques for analyzing vertebrate social structure using identified individuals: review and recommendations. *Adv Study Behav.* 28(33):74. doi:10.1016/S0065-3454(08)60215-6.
- Wolf M, van Doorn GS, Leimar O, Weissing FJ. 2007. Life-history trade-offs favour the evolution of animal personalities. *Nature.* 447(7144):581–584.
- Youngblood M. 2019. A Raspberry Pi-based, RFID-equipped birdfeeder for the remote monitoring of wild bird populations. *Ring. Migr.* 34(1):25–32. doi:10.1080/03078698.2019.1759908.