# Cooperative nest building in wild jackdaw pairs 

Luca G. Hahn* © , Rebecca Hooper® ${ }^{*}$, Guillam E. McIvor® ${ }^{*}$, Alex Thornton* ${ }^{*}$<br>University of Exeter, Centre for Ecology and Conservation, Penryn Campus, Cornwall, UK

## A R T I C L E I N F O

## Article history:

Received 15 December 2020
Initial acceptance 2 February 2021
Final acceptance 16 April 2021
Available online 6 July 2021
MS. number: 20-00897R

## Keywords:

animal architecture
bird nest
construction behaviour
cooperation
Corvus monedula
division of labour
jackdaw
monogamy


#### Abstract

Animals create diverse structures, both individually and cooperatively, using materials from their environment. One striking example is the nests birds build for reproduction, which protect the offspring from external stressors such as predators and temperature, promoting reproductive success. To construct a nest successfully, birds need to make various decisions, for example regarding the nest material and their time budgets. Research has focused mainly on species where one sex is primarily responsible for building the nest. In contrast, the cooperative strategies of monogamous species in which both sexes contribute to nest building are poorly understood. Here we investigated the role of both sexes in nest building and fitness correlates of behaviour in wild, monogamous jackdaw pairs, Corvus monedula. We show that both partners contributed to nest building and behaved similarly, with females and males present in the nestbox for a comparable duration and transporting material to the nest equally often. However, while females spent more time constructing the nest, males tended to invest more time in vigilance, potentially as a means of coping with competition for nest cavities. These findings suggest a moderate degree of division of labour, which may facilitate cooperation. Moreover, some aspects of behaviour were related to proxies of reproductive success (lay date and egg volume). Females that contributed relatively more to bringing material laid earlier clutches and pairs that spent less time together in the nestbox had larger eggs. Thus, selection pressures may act on how nest-building pairs spend their time and cooperatively divide the labour. We conclude that cooperative nest building in birds could be associated with monogamy and obligate biparental care and provides a vital but relatively untapped context through which to study the evolution of cooperation.


© 2021 The Association for the Study of Animal Behaviour. Published by Elsevier Ltd. All rights reserved.

Across the animal kingdom, species build structures for various purposes relevant for fitness. Such animal architecture (Hansell, 2005,2007 ) is used in diverse contexts, such as creating a protective shelter (Rosell, Bozser, Collen, \& Parker, 2005), reproduction and parental care (Deeming \& Reynolds, 2015), capture of prey (Hunt, 1996), and communication and signalling (Borgia, 1995). A striking example is bird nests built for reproduction (Collias, 1964; Collias \& Collias, 1984; Hansell, 2000; Healy, Walsh, \& Hansell, 2008), which influence fitness by protecting the offspring, for example from predators through camouflage (Bailey, Muth, Morgan, Meddle, \& Healy, 2015) and from environmental stressors, such as temperature (Campbell, Hurley, \& Griffith, 2018; Edwards, Shoot, Martin, Sherry, \& Healy, 2020). Additionally, nests can function as an intraspecific signal of investment in reproduction (Massoni, Reboreda, López, \& Florencia Aldatz, 2012; Soler, Møller, \& Soler, 1998) and to attract mates (Metz, Klump, \&

[^0]Friedl, 2009). While nest-building behaviour has long been assumed to be genetically predetermined (Nickell, 1958), recent evidence highlights an important role for learning (Bailey, Morgan, Bertin, Meddle, \& Healy, 2014; Breen, Guillette, \& Healy, 2016; Walsh, Hansell, Borello, \& Healy, 2013). For example, male zebra finches, Taeniopygia guttata, adjust their preferred material based on their success in a past breeding attempt (Muth \& Healy, 2011). However, research has focused on species in which single individuals (often males) predominantly build the nest: in zebra finches, for instance, studies have focused on males, which are responsible for bringing the nest material (Zann, 1996). While both partners may then contribute to arranging the material in the nest, their cooperative interactions at this stage have not been investigated in detail. There has been some work describing contributions to nest building in cooperative breeders like sociable weavers, Philetairus socius, and white-browed sparrow-weavers, Plocepasser mahali (Collias \& Collias, 1978; Leighton, 2014), but cooperative nest building by monogamous mates remains largely unexplored. This is particularly surprising given that monogamy and biparental care are common in the majority of bird species (Cockburn, 2006;

Orians, 1969). There is therefore a need to investigate whether and how monogamous birds cooperate during nest building. This will allow us to comprehensively understand the costs and benefits of cooperation between partners during this key stage of the breeding cycle, and, more broadly, will allow a deeper insight into the cooperative behaviours underlying animal architecture.

Effective cooperation between mates can be vital for fitness, particularly in species with obligate biparental care (Griffith, 2019). However, the interests of both sexes do not align exactly, generating sexual conflict (Chapman, Arnqvist, Bangham, \& Rowe, 2003; Harrison, Barta, Cuthill, \& Székely, 2009). Research has concentrated largely on how conflicts between mates are resolved when provisioning offspring (Hinde \& Kilner, 2007; Iserbyt, Farrell, Eens, \& Müller, 2015; Johnstone et al., 2014), making monogamous birds central study systems to understand the evolution of cooperative strategies. For instance, theoretical and empirical studies suggest that forms of conditional cooperation, such as turn taking (whereby each partner invests following a contribution by the other) may serve to reduce conflicts of interest and stabilize cooperation between mates (Johnstone et al., 2014; Johnstone \& Savage, 2019). Given that monogamous birds have long served as important model systems for understanding the evolution of cooperation, and that mates in some species are known to build the nest together (Birkhead, 2010; Massoni et al., 2012), it is striking that cooperative nest-building strategies have rarely been examined explicitly. Establishing the role of the two sexes during cooperative nest building is crucial to our understanding of both cooperative strategies and animal architecture.

In birds, the degree of cooperation between the sexes during nest building could be linked to the mating system. For instance, in various polygynous weaver species (Ploceidae) males build nests alone to attract females (Bailey et al., 2016), whereas in monogamous weavers mated pairs build their nest cooperatively (Habig, 2020). Furthermore, two largely genetically monogamous species, Eurasian magpies, Pica pica (Parrot, 1995) and rufous horneros, Furnarius rufus (Diniz, MacEdo, \& Webster, 2019), also build their nest cooperatively (Birkhead, 2010; Massoni et al., 2012). However, fine-scale behaviours and time budgets have not been explored, so cooperative nest building and its fitness consequences remain poorly understood. The degree to which partners cooperate is likely to depend on how much their interests align. In species showing obligate biparental care, mates should invest (relatively equally) in their offspring, because a lack of investment by either parent is likely to lead to failure of the reproductive attempt (Cockburn, 2006; Remeš, Freckleton, Tökölyi, Liker, \& Székely, 2015). Moreover, one could expect greater degrees of cooperation in species with low rates of extrapair fertilization (Lv et al., 2019) and high paternity certainty (Disciullo, Thompson, \& Sakaluk, 2019) as these conditions create highly interdependent fitness outcomes. The success of a clutch could be impacted by how bird pairs cooperate during nest building because cooperation may influence nest quality and because this process is energetically and temporally costly (Collias, 1964; Mainwaring \& Hartley, 2013). The energetic costs of nest building could vary between sexes due to differences in morphology, physiology, energetic demands and available information. Consequently, while both mates may behave similarly, sex-based differences in the costs associated with certain activities could promote task specialization, as shown by evolutionary individual-based simulations of individuals providing two types of parental care (e.g. feeding young and defending them against predators) associated with a sex-based asymmetry regarding the costs (Barta, Székely, Liker, \& Harrison, 2014). This could be important in the context of nest building as well; for example, male magpies and female rufous horneros bring relatively more material to the nest than the opposite sex. Investigating the roles of the
sexes, the level of cooperation, whether cooperation is repeatable within pairs and the fitness consequences during nest building is also vital to further understand how individuals cope with the informational demands of decision-making processes while tracking another individual's behaviour (Emery, Seed, Von Bayern, \& Clayton, 2007). Tracking each other's behaviour could favour greater levels of behavioural synchrony, which could also be related to behavioural compatibility between partners, potentially resulting in more effective cooperation and greater reproductive success (Spoon, Millam, \& Owings, 2006).

Jackdaws, Corvus monedula, provide a particularly suitable study system to investigate cooperation during nest building. They are a highly social, colony-breeding corvid that forms long-term pair bonds (Lorenz, 1931; Wechsler, 1989). Pairs produce one clutch per year, with both sexes providing care to altricial chicks (Henderson \& Hart, 1993). Moreover, unlike most socially monogamous bird species, jackdaws are highly genetically monogamous, so the reproductive success of partners is more interdependent than in species where extrapair offspring are common (Gill, van Schaik, von Bayern, \& Gahr, 2020). In jackdaws, both sexes participate in building nests within cavities, which consist of a platform (made of sticks and twigs) and a cup with soft material (grasses, moss, mud and animal hair, for example). Tightly linked fitness outcomes may generate selection pressure for cooperation between partners throughout the breeding season, including during the nestbuilding stage.

This study had two main objectives. (1) We first quantified the behaviours and time budgets of pairs. We hypothesized that cooperation and division of labour between partners during nest building should evolve where both individuals derive symmetrical fitness benefits from producing a suitable nest. First, we predicted females and males should behave similarly by investing in the nest directly (e.g. by bringing nest material) and indirectly (e.g. through vigilance) (Prediction 1). Second, however, we predicted that the time invested in these behaviours may not be symmetrical between the sexes given morphological, physiological and informational differences (Prediction 2). (2) Our second objective was to examine the ultimate function of behaviours during nest building by investigating three different fitness proxies: relative lay date, clutch size and egg volume. Laying earlier clutches can be advantageous and is often linked to reproductive success in birds (Perrins, 1965, 1970; Mclvor, Goumas, Alothyqi, Troscianko \& Thornton, n.d.), for example because earlier layers face less competition in finding food for their young. Larger eggs could potentially provide the embryo with more resources, aiding its development and increasing the probability of surviving (Krist, 2011). We hypothesized that how much birds invest in their nest and how they share the workload could be associated with reproductive success, with pairs that invest more overall and divide the labour (so that males contribute at least equally) being favoured. While females should invest substantially in the nest because they may be better informed about their requirements for incubation, males should contribute equally because this may allow females to invest more resources in the clutch, potentially maximizing reproductive success. Furthermore, investment may determine the time taken to build the nest, which is a limiting factor for laying the first egg, as well as nest quality, which in turn could enhance embryo development and survival. First, we predicted that how pairs allocate their time and energy between different activities could impact their fitness. More specifically, pairs that invest more in the nest relative to other activities, such as vigilance and nest defence, should lay earlier, and have larger clutches and eggs (Prediction 3). Moreover, pairs that show greater total investment in the nest should lay earlier clutches (Prediction 4). We predicted this because pairs that invest more total time in nest building are likely to complete their nest faster,
and having a complete nest is a crucial limiting factor for clutch initiation. Pairs in which males invest at least equally in the nest as well as in nest site defence should lay earlier and produce larger clutches and eggs (Prediction 5). If the optimal solution was for both individuals to invest equally, one might expect a quadratic relationship between relative contributions of females compared to the overall investment and fitness proxies. Finally, we predicted that jackdaws behaving more synchronously by spending more time together in the nestbox should lay earlier and have larger clutches and eggs (Prediction 6). As selection on nest-building behaviours may depend on the degree to which they constitute repeatable traits, we also investigated the repeatability of behaviour over time.

## METHODS

## Ethical Note

This study was conducted with approval from the University of Exeter Research Ethics Committee (eCORN002970), following the ASAB/ABS Guidelines for the Treatment of Animals in Behavioural Research. Jackdaws had previously been colour ringed for individual identification by qualified ringers licensed by the British Trust for Ornithology. The sex of each individual was confirmed through molecular sexing of blood samples (Griffiths, Double, Orr, \& Dawson, 1998) under a U.K. Home Office licence (project licence 30/3261). Morphometrics of individuals, such as wing length, tarsus length and body mass, were measured when they were caught for ringing (see Greggor, Spencer, Clayton, \& Thornton, 2017 for details). We used the exact age if birds had been ringed as nestlings, juveniles or first years. For birds that had been first captured as adults (as determined by plumage characteristics), we assumed they were at least 2 years old when first captured. This meant that the minimum possible age for birds caught as adults when the colonies were established in 2013 was 7 years in 2018 and 8 years in 2019, but some birds may have been older than this.

## Data Collection

As cavity nesters, jackdaws accept nestboxes and typically return to the same nest site across years, allowing researchers to monitor behaviour by fitting boxes with video cameras. We recorded nest-building behaviour of free-living jackdaws using CCTV cameras with integrated microphones (380TVL CMOS camera, Handykam, Redruth, U.K.) concealed inside the roof space of nestboxes during the breeding seasons of 2018 and 2019 at three breeding colonies in Cornwall, U.K.: X $\left(50^{\circ} 10^{\prime} 23^{\prime \prime} \mathrm{N} ; 5^{\circ} 7^{\prime} 12^{\prime \prime} \mathrm{W}\right), \mathrm{Y}$ ( $50^{\circ} 11^{\prime} 26^{\prime \prime} \mathrm{N}, 5^{\circ} 10^{\prime} 51^{\prime \prime} \mathrm{W}$ ) and $\mathrm{Z}\left(50^{\circ} 11^{\prime} 56^{\prime \prime} \mathrm{N}, 5^{\circ} 10^{\prime} 9^{\prime \prime} \mathrm{W}\right)$. All boxes were of identical dimensions and made from EKOply (Second Life Products, Swansea, U.K.), a recycled plastic plywood alternative. Each box had a dedicated camera system that was installed at least 1 day prior to a recording being made and was then left in place thereafter. The cables from the cameras ran from the back of the box to ground level allowing us to set up a portable video recorder (JXD 990, JXD Co., Shenzhen, China) on the morning of filming without disturbing the nest itself. Nestboxes were filmed slightly but significantly closer to the lay date in 2019 (mean $\pm$ $\mathrm{SD}=7.69 \pm 5.36$ days) than in 2018 ( $10.83 \pm 6.41$ days; linear model, LM: $\left.\beta \pm \mathrm{SE}=-3.136 \pm 1.551, t_{59}=-2.022, P=0.048\right)$. The criterion we employed to film a nestbox was that there had to be at least one layer of nest material, although the precise timing varied somewhat according to requirements for data collection for other studies (Hooper, 2019; Cuff \& Quicray, 2019). All observations were conducted in the morning (start time 0630-0930 hours) to minimize the confounding effect of changing behavioural patterns
throughout the day. In total, we recorded 183.04 h of video data ( $N=62$ videos; one video $=$ one observation; mean video length $\pm$ SD $2.95 \pm 1.07 \mathrm{~h}$, range $1-5 \mathrm{~h}$ ) from 40 distinct, breeding jackdaw pairs across 40 different nestboxes ( $N=5$ videos in five boxes at colony $\mathrm{X}, 27$ videos in 15 boxes at Y and 30 videos in 20 boxes at Z). In 2018, we conducted 35 observations of 29 pairs (six pairs were observed twice), and in 2019 we conducted 27 observations of 25 pairs (two pairs were observed twice; Appendix Table A1). Of the 25 pairs that we filmed in 2019, 14 pairs had previously been studied in 2018. We recorded a minimum of one video at each nest during the middle of the nest-building phase in April (24 pairs were observed only once in our study, either in 2018 or in 2019). We filmed eight pairs once each year, and four pairs twice in one year and once in the other year. Two pairs were observed twice within one year but not in the other year and one pair was observed twice in each year. In two instances, ownership of a box changed between years because one pair was replaced by another pair in 2018 (box Z28) and one pair switched boxes (Z33 to Z45 in 2019). Boxes were checked weekly from mid-March for building activity, and daily from early April to record the exact date of clutch initiation. Checks were then performed daily until no new eggs were recorded on 3 consecutive days, at which point the clutch was considered complete. The eggs were numbered on the day they were laid using a nontoxic marker pen, and in addition to recording the clutch size we also photographed the eggs 8 days after clutch initiation, allowing us to calculate their volume using the method devised by Troscianko (2014). In all observations, jackdaws built a nest, and all but one pair (box Z28, 2018), which was displaced by another pair, laid eggs.

## Video Analysis

We analysed videos in a randomized order with regard to 'year' and 'study site', using the software BORIS version 7.5.1 (Friard \& Gamba, 2016). Relevant behaviours were recorded as either 'point events' or 'states' (to quantify the number or duration of events, respectively; see ethogram in Table 1) and the identity and sex of each individual was determined from its unique colour ring combination. In a minority of cases, rings were not visible in the video during a bird's visit to the nestbox, so the individual's sex was recorded as 'unknown'. When the sex was relevant for analyses, we excluded data from unknown focal individuals. If vocalizations occurred when both members of a pair were in the nestbox, we used fine-scale body movement (e.g. of beak or thorax) to establish which individual was vocalizing. We analysed different types of vocalizations with distinct acoustic qualities separately. We analysed 'chatter', a distinctive sequence of repeated high-pitch vocalizations, separately from other calls (hereafter called 'calls'). These 'chatter' vocalizations are highly distinctive and easy to distinguish by ear from other vocalizations without the need for acoustic software.

## Statistical Analyses

Data were analysed in $R$ version 4.0.2 ( R Core Team, 2020). For all behavioural variables ( $N=10$ ), we extracted raw total durations (states) and quantities (point events) per observation for each pair (in total) and for both sexes separately. To standardize variables, we divided raw data (durations or counts) by the observation length (s), and multiplied state events by 100 (percentage of time) and point events by 3600 (rate per hour). For each standardized behavioural variable, we calculated means and SDs across pairs.

Table 1
Ethogram of behaviours recorded

| Behaviour | Type | Description |
| :--- | :--- | :--- |
| Call | PE | Bird makes a call |
| Remove nest material | PE | Removing objects when leaving the nestbox |
| Build nest | SE | Moving and adjusting nest material using the beak |
| Form cup | SE | Lying down and adjusting soft material to form the cup using legs |
| In box/Visits | $\mathrm{SE} / \mathrm{PE}$ | Bird visits the nestbox (PE) and spends time in it (SE) |
| Chatter | SE | Distinctive soft, high-pitch vocalizations |
| Bring nest material | PE | Carrying objects (grass, twigs, etc.) when entering the nestbox |
| Vigilance | SE | Peeking outside the nestbox |
| Food sharing ( $\delta^{\prime}$ ) | PE | Male visits nestbox and shares food with his mate |
| Modify nest material | SE | Changing the structure of nest material, e.g. by cutting twigs |

Type denotes whether a behaviour was a point event (PE) or a state event (SE) with a duration. Begging calls by females are included in the event 'call'.

## General procedure: mixed models and repeatability analysis

Mixed models. We analysed data with (generalized) linear mixed models (LMMs and GLMMs), using the packages lme4 (for LMMs; Bates, Mächler, Bolker, \& Walker, 2015) and glmmTMB (for GLMMs; Brooks et al., 2017).

To formulate models, we selected the dependent variable of interest (a behaviour or a fitness proxy) and one or more biologically meaningful independent variable(s). We subsequently tested model assumptions such as normality of residuals, homoscedasticity, error structure, dispersion, zero inflation and influential data points (Cook's distance), using diagnostic plots and tests implemented in R (LMMs) or in the package DHARMa (GLMMs; Hartig, 2019). To infer estimates and $P$ values, we used Wald tests in the package car (Fox \& Weisberg, 2011). All models included the variables 'year' (fixed effect), 'pair ID' (random effect) and 'study site' (random effect) to account for temporal and spatial variation as well as pseudoreplication. We had no specific a priori predictions as to the effects of the birds' age, but as it could potentially influence behaviour and reproductive success, we initially included 'age' (years) in analyses as an additional fixed effect. If age did not appear to play an important role, we removed the variable from analyses to avoid over-parametrization and maximize statistical power. Observation level random effects (Harrison, 2014) and Con-way-Maxwell-Poisson error structures accounted for zero inflation and underdispersion, respectively.

Repeatability analysis. We calculated the repeatability of behaviours and fitness proxies in pairs for which repeated measures were available ( $N=28$ observations of fitness proxies for 14 pairs that were observed once per year and $N=38$ observations of behaviour for 16 pairs observed more than once within and/or across years; Appendix Table A1), using the package rptR (Stoffel, Nakagawa, \& Schielzeth, 2017). When quantifying repeatability of state events, we used Box-Cox transformations (Sakia, 1992), using the package MASS (Venables \& Ripley, 2002), to meet assumptions of Gaussian data.

## Sex differences in behaviour (Predictions 1 and 2)

We could identify birds for $76.71 \pm 30.48 \%$ t of the time spent in the box. To quantify sex differences in behaviour, we used a subset of behavioural data where the identity of the focal individual(s) was known ( $N=62$ videos of 40 pairs; two cases were removed in analyses including vocal communication because the microphones failed to record: box Y02, 2018 and box Z19, 2018). We investigated the time both sexes invested in 'vigilance', 'nest building', 'being in the nestbox', 'forming the cup' and 'chatter', using separate LMMs, with the standardized response variables log transformed to meet model assumptions (West, Welch, \& Galecki, 2014). We also examined whether either sex invested more time in 'vigilance' or 'nest building' as response variables given that these were the most frequent behaviours in the nestbox. Models examining 'vigilance' and 'nest building' as response terms also contained the 'number of days the video had been recorded before the lay date' (covariate) to account for the potential influence the date may have on behaviours, and to disentangle variation caused by the year and lay date. For instance, birds may reduce their effort closer to the lay date when the nest should be completed. On the other hand, males could increase their vigilance closer to the lay date to guard the female during her fertile period. 'Modification of material' was too rare to permit formal statistical analysis. We also conducted separate GLMMs on rates of 'material brought', 'material removed', 'visits to the nestbox' and 'calls' (rounded to rate per hour and treated as count data) fitted as a response term. In these analyses, 'sex' was the main predictor variable of interest, but we also modelled an interaction between sex and age of each bird to examine whether sex differences may be age dependent and to include age as a covariate potentially affecting behaviour.

## Nest building and reproductive success (Predictions 3-6)

Dependent variables: reproductive parameters. To examine fitness correlates of behaviours, we separately analysed three different proxies for reproductive success (Table 2). First, we used the relative lay date of the first egg compared to the date the first clutch was initiated per site. As colonial breeders, jackdaws breed within a relatively short period, and early layers may benefit from lower

Table 2
Reproductive parameters examined

| Reproductive parameter | Definition | Error structure | $N$ | Mean $\pm$ SD | Range |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Lay date | Day first egg laid ( $1=$ first egg per site $)$ | COM-Poisson | 61; 39 | $5.19 \pm 2.92$ | 1-14 |
| Clutch size | Total number of eggs laid | COM-Poisson | 61; 39 | $4.43 \pm 0.87$ | 2-6 |
| Egg volume | Egg volume of first egg ( $\mathrm{cm}^{3}$ ) | Gaussian | 60; 38 | $11.42 \pm 0.85$ | 9.75-13.19 |
| Egg volume | Egg volume of third egg ( $\mathrm{cm}^{3}$ ) | Gaussian | 58; 36 | $11.23 \pm 1.08$ | 8.62-13.35 |

COM-Poisson: Conway-Maxwell-Poisson. ' $N$ ' denotes the sample size (number of videos and pairs, respectively) used in the models (unless stated otherwise). To calculate mean, SD and the range (minimum-maximum) of parameters, the sample size was smaller than indicated here because seven pairs observed twice in 1 year were only considered once here.
competition with other colony members. A second proxy of reproductive success was the clutch size. Third, we examined the volume of the first and the third egg. Jackdaws lay an egg per day until the clutch is complete, and they show hatching asynchrony, with the first egg being the one most likely to survive. The second egg has a relatively high probability of survivng as well, whereas the survival rate of the third egg is approximately 0.5 . We did not analyse later eggs as these rarely survive (McIvor, Goumas, Alothyqi, Troscianko \& Thornton, n.d.). One pair (box Z28, 2019) was excluded from these analyses because it was displaced by another pair during nest construction so could not produce a clutch. When analysing egg volume, we removed one pair (box Y21, 2019), which had been parasitized by a conspecific female. Two pairs were removed when analysing the third egg volume as they only laid two eggs (box Y16, 2019; box Z15, 2019).

Behavioural predictors. We defined four 'behavioural concepts' to be used separately as independent variables that may relate to measures of reproductive success. For each of the first three concepts we calculated a distinct principal component analysis (PCA) to summarize (scaled) behavioural variables to be included in models while minimizing model complexity (Budaev, 2010; Morton \& Altschul, 2019) and to account for multicollinearity among variables (Graham, 2003). When performing a PCA, we calculated a correlation matrix including the variables of interest, applied the Kaiser-Meyer-Olkin (KMO) measure (threshold 0.5) to test for sufficient correlation among them (Budaev, 2010), and conducted a 'parallel analysis', which is a tool to determine the number of principal components to be considered objectively (Morton \& Altschul, 2019), using the package psych (Revelle, 2018). According to the 'parallel analysis', a component is included if its eigenvalue is greater than the 95th percentile of a distribution of eigenvalues that were generated from randomized data (Horn, 1965). We constructed alternative models in cases where we analysed distinct predictors each of which reflected a specific hypothesis (for more details see below). To select a model, we employed Akaike's information criterion (AICc to account for small sample sizes; Harrison et al., 2018) in the package bbmle (Bolker \& R Core Team, 2017). A model with the lowest AICc had to differ by at least 2 AICc units to be selected. In instances where only one predictor variable corresponded to one of the behavioural concepts, we did not use model selection and constructed a single model per fitness variable instead. When we detected a significant relationship between behavioural predictors and a fitness proxy, we performed models again with a subset of observations for which data on female body condition were available in that particular year to control for this variable (covariate). We then compared two models with and without female body condition using likelihood ratio tests. Body condition was quantified using the residuals of a regression examining the relationship between a measure of body size $\left(\mathrm{PC} 1_{\text {Body }}\right.$, derived from a PCA containing the variables tarsus and wing length) as independent and body mass as dependent variables. In all models investigating fitness correlates of behaviour, we also included the 'number of days the video was recorded before the lay date' ('day' henceforth, covariate), because this could have influenced the birds' behaviour. Moreover, we fitted 'female age' (covariate) to account for breeding experience (female and male age was significantly correlated: $\rho=0.744, t_{59}=8.560, P<0.001$ ). Another covariate was 'food sharing' by males because this cooperative behaviour did not relate directly to nest building but could affect reproductive success. We outline each concept, and the analytical methods used to examine it, below.
(1) Overall activity levels and intensity of behaviours (Prediction 3). To test Prediction 3 that pairs that invest more in the nest
relative to other activities (such as vigilance) should lay earlier and have larger clutches and eggs, we constructed a PCA of nine behavioural variables ('PCA All'; Table 3, Appendix Table A2). The variable 'food sharing' (rate per hour) was left out of the PCA due to the KMO threshold ( 0.43 ) but included in the models as a covariate as levels of food sharing by the male could influence the female's ability to invest in the nest and the clutch. Based on the results of a 'parallel analysis', we retained two principal components ( $\mathrm{PC} 1_{\text {All }}$ and $\mathrm{PC} 2_{\text {All }}$ ), which were used as explanatory terms in analyses. All behavioural variables loaded negatively onto the first principal component, $\mathrm{PC} 1_{\text {All }}$, which could therefore be interpreted as reflecting the overall 'intensity' of behaviours. The four nestbuilding behaviours ('modify' was too rare for formal statistical analyses and was therefore removed from the PCA) loaded negatively onto the second principal component, PC2 All, whereas the other behaviours (vigilance, vocalizations, time in the box) loaded positively onto PC2 all. These opposite loadings suggest a trade-off, such that pairs may have invested relatively more time in either the nest or in vigilance and vocalizing. Therefore, we hypothesized that (1) all behaviours ( $\mathrm{PC} 1_{\text {All }}$ ), (2) a relative investment in nest building compared to other behaviours ( $\mathrm{PC} 2_{\text {All }}$ ) or (3) both ( $\mathrm{PC} 1_{\text {All }}$ and PC2 $2_{\text {All }}$ ) could be used as predictors of reproductive success. We formulated three corresponding models and two further models which contained (4) only 'year' and 'day' and (5) only an 'intercept'. Subsequently, we compared these models using AICc.
(2) Direct investment in the nest ('effort'; Prediction 4). To examine Prediction 4, we analysed the relationship between a PCA comprising variables related directly to nest building and reproductive success ('PC1 ${ }_{\text {Effort }}$; Table 3, Appendix Table A3). All four variables loaded negatively onto ' $\mathrm{PC} 1_{\text {Effort' }}$ ', suggesting it could be interpreted as a measure of total nest-building effort. Following the 'parallel analysis' (Morton \& Altschul, 2019), we did not consider ' $\mathrm{PC} 2_{\text {Effort }}$ ' further and constructed only one model per fitness proxy, including ' $\mathrm{PC} 1_{\text {Effort }}$ ' as an independent variable instead of comparing alternative models.
(3) Relative investment by females ('division of labour', 'DoL'; Prediction 5). We conducted a third PCA to examine 'DoL', that is, whether the relative proportion of female contribution to nest building and vigilance (compared to the sum of female and male effort) was linked to reproductive success ('PCA ${ }_{\text {DoL }}$ '; Table 3, Appendix Table A4). In this analysis, the sample size was smaller ( $N=47$ observations), because we discarded observations when a proportion could not be calculated (neither sex of a pair showed one of the behaviours). Based on the 'parallel analysis', two principal components were retained (PC1 DoL and PC2 DoL $)$ PC1 DoL suggested that females contributed either more through nest building or vigilance due to opposite loadings, with positive values indicating relatively more investment in the nest and negative values indicating relatively more investment in vigilance (Appendix Table A4). As $\mathrm{PC} 2_{\text {DoL }}$ was strongly dominated by the variable 'relative proportion of material brought by females', which had a loading of 0.99 (Appendix Table A4), we used this variable (as opposed to the principal component) as a predictor variable in our models. We ran separate analyses with each fitness proxy as a response term. For each fitness proxy we constructed alternative models with each of the following explanatory terms: (1) 'PC1 DoL', (2) the 'relative proportion of material brought by females', (3) only 'year' and 'day' or (4) 'intercept' only. For predictors (1) and (2) we also modelled a quadratic effect which could indicate that equal contributions by both sexes are related to greater reproductive success.
(4) Time spent together in the nestbox ('synchrony'; Prediction 6). To test Prediction 6 that the level of 'synchrony' would be linked to reproductive success, we used the 'proportion of time individuals spent together in the nestbox' as an independent variable. To

Table 3
Behavioural concepts used to examine correlates of reproductive success

| Behavioural concept | Definition | Predictor variables in models (standardized by observation length) |
| :---: | :---: | :---: |
| All behaviours ( PCA $_{\text {All }}$; $N=59 ; 39)$ | Intensity of behaviours: nest building, vigilance, time in the box, vocalization | $\mathrm{PC} 1_{\text {All }}$ and $\mathrm{PC} 2_{\text {All }}$ (from a PCA including 'nest building', 'material brought', 'material removed', 'forming the cup', 'vigilance', 'chatter', 'calls', 'together in the nestbox', 'box occupied') |
| Effort ( PCA $\left._{\text {Effort }} ; N=61 ; 39\right)$ | Direct investment in nest-building activities | PC1 1 Effort ('nest building', 'material brought', 'material removed', 'forming the cup') |
| Division of labour ( $\mathrm{PCA}_{\text {DoL }} ; N=47 ; 34$ ) | Relative contribution by females to cumulative time investment in nest building and vigilance ('division of labour') | PC1 $1_{\text {DoL }}$ (relative proportion of 'nest building', 'material brought', and 'vigilance' by females) and 'material brought' (the dominant variable within PC2 $2_{\text {DoL }}$ ) |
| Synchrony ( $N=61$; 39) | Visits and time in the box matched by both birds | 'Together in the nestbox' |

When examining the behavioural concepts 'all behaviours' and 'effort', we used the PCs listed as explanatory variables in the models; for the concept 'division of labour', we fitted models with PC1 $1_{\text {DoL }}$ and with the separate variable 'material brought' (as this variable dominated PC2 ${ }_{\text {Dol }}$, with a loading of 0.99 ). The two sample sizes denote the number of observations and the number of pairs, respectively. In the $\mathrm{PCA}_{\text {All }}$, the variables 'food sharing' ( $\mathrm{KMO}=0.46$ ) and 'modify' (rare behaviour) were left out. For more details on the PCA see Appendix Tables A2-A4.
examine its relationship with fitness measures, we constructed one model per fitness proxy with 'synchrony' being the only independent variable.

## RESULTS

## Behaviours and Sex Differences

## Sex differences (Predictions 1 and 2)

On average, jackdaw pairs occupied their nestbox for $29.09 \pm 19.64 \%$ of the observations and spent $23.17 \pm 24.58 \%$ of that time together (Appendix Table A5). Pairs invested $18.12 \pm 16.43 \mathrm{~min}$ (cumulatively) in building the nest with their beak and $3.23 \pm 3.30 \mathrm{~min}$ forming the cup using their legs (Appendix Table A5). Birds transported $18.89 \pm 20.99$ nest material items to their nestbox during the period of observation (Appendix Table A6). Jackdaws also modified nest material, for example by trimming twigs, but this comprised a small proportion of observation time ( $6.52 \pm 20.75$ s or $0.05 \pm 0.13 \%$ of the observations). The behavioural repertoire of both sexes was broadly similar (Fig. 1,

Appendix Table A7), but they also differed in some behaviours (Fig. 2, Appendix Table A7). Specifically, females spent 1.5 times more time building the nest than males (Fig. 2a, Appendix Table A7). We also found that birds spent more time building in 2018 than in 2019. This difference was not because videos were recorded closer to the lay date in 2019 (LMM: days before lay date: $\beta \pm \mathrm{SE}=0.000 \pm 0.007, X^{2}{ }_{1}=0.004,95 \%$ confidence interval, CI [$0.01,0.01], P=0.952$ ).

Males spent on average 1.4 times more time being vigilant than females, but this difference was not significant (Fig. 2b, Appendix Table A7). Moreover, males spent more time being vigilant than they spent building, while females spent similar times in both behaviours (Appendix Table A7). Males did not increase vigilance when the observation was recorded closer to the lay date (LMM: sex*days before lay date: $\beta \pm \mathrm{SE}=0.021 \pm 0.023, X^{2}{ }_{1}=0.887,95 \%$ CI [-0.01, 0.03], $P=0.346$ ). As with nest building, we found that investment in vigilance varied across years, with levels of vigilance being lower in 2019 than 2018. This difference could not be attributed to differences in the timing of observations across years, as there was no effect of 'days before lay date' in the model (LMM:


Figure 1. Mean relative duration (state events) and frequency of events (point events) by sex ( $N=62$ observations; $N=60$ for vocalizations). The horizontal line marks the proportion of 0.5 , meaning both sexes showed a behaviour equally long or often, respectively. Asterisks indicate a significant sex difference in behaviour based on the model output (calls, nest building): $* P<0.05 ; * * P<0.01$.


Figure 2. Sex differences in three behavioural variables ( $N=62$ observations; $N=60$ for calls). Horizontal lines in the boxes indicate the median, green asterisks indicate the mean. Upper and lower ends of the boxes reflect the 0.25 and the 0.75 quartiles, respectively. Horizontal lines connecting points represent distinct pairs. Asterisks indicate a significant difference: $\mathrm{NS}=0.06 ; * P<0.05 ; * P<0.01$. (a) The time spent building the nest as a percentage of the observation length. (b) The time spent being vigilant as a percentage of the observation length. (c) The number of calls per hour made by both sexes.
days before lay date: $\beta \pm \mathrm{SE}=0.004 \pm 0.009, X^{2}{ }_{1}=0.242,95 \% \mathrm{CI}[-$ $0.01,0.02], P=0.623$ ).

Females called 1.9 times more frequently than males (Fig. 2c, Appendix Table A7), even after removing female begging calls (GLMM: sex: $\beta \pm \mathrm{SE}=-0.690 \pm 0.228, X^{2}{ }_{1}=9.120, P=0.003$ ). There was weak evidence that older birds brought more nesting material (GLMM: age: $\beta \pm \mathrm{SE}=0.257 \pm 0.124, X^{2}{ }_{1}=4.135,95 \% \mathrm{CI}$ [ $0.013,0.500$ ], $P=0.042$; Appendix Fig. A1), but this relationship was not maintained when the four youngest individuals, which were 2 years old, were removed (GLMM: age: $\beta \pm \mathrm{SE}=0.177 \pm 0.127, \quad X^{2}{ }_{1}=1.645, \quad 95 \%$ CI $[-\quad 0.073,0.426]$, $P=0.200$ ). Aside from this there was no evidence for any age effects or sex by age interactions on any aspect of building behaviour (Appendix Table A8).

## Repeatability and variation across pairs and years

There was considerable variation in behaviours between pairs (Fig. 2, Appendix Tables A5, A6), but when inside the nestbox, most of their time was spent building the nest or being vigilant (Appendix Table A5). On the level of the pair, birds that spent more of their time in the nestbox together spent more time being vigilant ( $\rho=0.906, t_{60}=16.547, P<0.001$ ), but not more time building ( $\rho=0.139, t_{60}=1.090, P=0.280$ ). Conversely, pairs in which only one individual occupied the nestbox for longer spent more time building ( $\rho=0.840, t_{60}=11.996, P<0.001$ ), but not more time being vigilant ( $\rho=0.175, t_{60}=1.379, P=0.173$ ). The time females spent building and the time males spent being vigilant were positively correlated ( $\rho=0.271, t_{60}=2.184, P=0.033$ ). No behaviour was repeatable in 16 pairs for which repeated measures were
available within and/or between years (Appendix Table A9). Jackdaws' behaviour also varied between years (Appendix Table A7).

## Behaviours and Correlates of Reproductive Success (Predictions 3-6)

The majority of jackdaw females laid their first egg in the middle of April ( $17.05 \pm 3.20$ days where $1=1$ April; $5.19 \pm 2.92$ days after the first clutch was initiated per site). The lay date of pairs (relative to the first lay date per site) was repeatable (Appendix Table A10). Females laid a mean of $4.43 \pm 0.87$ eggs, and clutch size was not repeatable for those pairs observed in both years (Appendix Table A10). The mean volume of the first and the third egg was $11.42 \pm 0.85 \mathrm{~cm}^{3}$ and $11.23 \pm 1.07 \mathrm{~cm}^{3}$, respectively. The volume of the first egg and the third egg females laid was repeatable across years (Appendix Table A10).

Overall activity levels and intensity of behaviours (Prediction 3)
The intensity of behaviours $\left(\mathrm{PC}_{\mathrm{AlI}}\right)$ and the amount of time birds invested in nest-building behaviours compared to other behaviours, such as vigilance ( $\mathrm{PC}_{\mathrm{All}}$ ), was not associated with any proxy of reproductive success (relative lay date, clutch size, egg volume; Appendix Table A11).

## Direct investment in the nest ('effort'; Prediction 4)

Nest building effort ( $\mathrm{PC} 1_{\mathrm{Effort}}$ ) did not predict variation in reproductive success (relative lay date, clutch size, egg volume; Appendix Table A11).

## Relative investment by females ('division of labour', 'DoL’; Prediction 5)

Females that contributed relatively more to bringing nest material laid their first egg relatively earlier (Fig. 3, Appendix Table A12). In this model, female age had no effect, so it was removed (GLMM: female age: $\beta \pm \mathrm{SE}=0.086 \pm 0.082, X^{2}{ }_{1}=1.108$, $P=0.292,95 \% \mathrm{Cl}[-0.074,0.246])$. Including a proxy for female body condition did not improve the model ( $X^{2}{ }_{1}<0.001, P>0.99$ ). We found no relationship between 'DoL' and the fitness proxies clutch size and egg volume (Appendix Table A11).

Time spent together in the nestbox ('synchrony'; Prediction 6)
The volume of both the first and the third egg was smaller in pairs that spent more time together in the box (Fig. 4, Appendix Table A12). This relationship remained after excluding an influential data point (a pair that spent more than $60 \%$ of the time together in the nestbox; LMM, synchrony, first egg: $\beta \pm S E=-0.027 \pm 0.012$, $X^{2}{ }_{1}=5.590, P=0.018,95 \%$ CI $[-0.051,-0.003]$; third egg: $\beta \pm \mathrm{SE}=-0.046 \pm 0.014, \quad X^{2}{ }_{1}=10.648, \quad P=0.001, \quad 95 \% \quad$ CI $\quad[-$ $0.075,-0.016])$. In the models examining synchrony there was no effect of female age on egg volume (LMM: age, first egg: $\beta \pm \mathrm{SE}=0.102 \pm 0.094, X^{2}{ }_{1}=1.169, P=0.280,95 \%$ CI $[-0.076$, 0.290 ]; third egg: $\beta \pm \mathrm{SE}=0.005 \pm 0.115, X^{2}{ }_{1}=0.002, P=0.962$, $95 \%$ CI [-0.212, 0.228]). Including female body condition did not improve the model examining the relationship between synchrony and first egg volume ( $X^{2}{ }_{1} 0.113, P=0.945$ ) or between synchrony and third egg volume ( $X^{2}{ }_{1} 0.013, P=0.909$ ). Pairs that spent more time being vigilant had smaller first eggs and third eggs (LMM: vigilance, first egg: $\beta \pm \mathrm{SE}=-0.013 \pm 0.007, X^{2}{ }_{1}=3.919, P=0.048$, $95 \% \mathrm{Cl}[-0.003,0.000]$; third egg: $\beta \pm \mathrm{SE}=-0.022 \pm 0.008$, $X^{2}{ }_{1}=6.934, P=0.008,95 \%$ CI [-0.038, -0.005$]$ ). Variation in lay date and clutch size was not associated with variation in synchrony (Appendix Table A11).

## DISCUSSION

Our findings demonstrate that in monogamous jackdaws, nest building entails substantial investment from both partners, and may have important fitness consequences. Although both sexes had a similar behavioural repertoire and cooperated to create their nest, there were some sex differences in investment, with females building more and males tending to be more vigilant. Moreover, some aspects of behaviour, such as the relative female contribution to transporting material and time spent together in the box, were associated with measures of reproductive success (lay date and egg volume).

Consistent with our Prediction 1, jackdaws cooperated during nest construction, with the two sexes behaving broadly similarly, with both investing in bringing material, building the nest and forming the cup. In jackdaw pairs fitness outcomes are interdependent due to repeated mating opportunities with the same partner across years and low rates of divorce and successful extrapair copulations (Gill et al., 2020; Wechsler, 1989). Under these circumstances, conflicts of interest between mates may be minimized, particularly if biparental care is necessary to successfully rear offspring, ultimately favouring cooperation. Although nest building by jackdaws clearly requires a substantial cooperative investment from both partners, our findings suggest moderate division of labour could facilitate cooperation, which is in accordance with Prediction 2 (cf. Iserbyt, Fresneau, Kortenhoff, Eens, \& Müller, 2017). Females built more than males and were therefore more responsible for the nest structure. In contrast, males dedicated more time to vigilance than to building, which may be particularly important in colonially nesting jackdaws, where intraspecific competition over nest cavities is severe and can constrain reproduction (Henderson \& Hart, 1993; Röell, 1978; Verhulst \& Salomons, 2004). Vigilant residents may not only anticipate threatening nonresident competitors searching for a nest cavity,


 lines show the $95 \%$ confidence intervals around the fitted line (solid) from the model output.


Figure 4. Relationship between the first egg volume per female and the percentage of time mates spent together in the nestbox. The continuous fitted line corresponds to the model output; dotted lines show the $95 \%$ confidence intervals.
but their bright eye colour has also been shown to deter intruders (Davidson, Clayton, \& Thornton, 2014). Males may prioritize vigilance because the risks of vigilance and defence may be more costly for females as they need to stay in a good condition for later stages of breeding, such as incubation. Additionally, males may invest relatively more in vigilance than in building due to their slightly larger body size (Fletcher \& Foster, 2010), a trait that impacts contests in this species (Verhulst, Geerdink, Salomons, \& Boonekamp, 2014). There was no significant difference in the amount of time males and females spent in vigilance, and male vigilance was independent of days until his partner's fertile window (beginning 5 days prior to the lay date; Gill et al., 2020), suggesting that vigilance serves primarily to defend the nest site rather than as a form of mate guarding. Males cooperated, for example through vigilance and transporting nest material, despite contributing less to building the nest by arranging material in the nestbox. By increasing their own nest-building activity, females may be able to partially compensate for this. Females may also spend more time building than males because they may be better informed about their own requirements for incubating the clutch. The mechanisms through which partners acquire and act upon information to respond to each other's behaviour and coordinate division of labour remains unknown. Elucidating these mechanisms will be vital to understanding the cognitive demands of pair bonding, such as the need to track and respond to another's behaviour (Emery et al., 2007).

Our results suggest substantial variation in behaviour and time budgets between pairs. Furthermore, no behavioural variable was significantly repeatable within pairs, indicating there may also be considerable behavioural variation within pairs. It is possible that the lack of repeatability within pairs is an artefact of differences in sampling between years because videos were recorded significantly closer to the lay date in 2019, which could have affected the behaviour. For instance, pairs may have seemingly built less in 2019, but this could have been because the video was recorded closer to the lay date. Given the limited amount of data per pair and the fact that not all pairs were observed repeatedly, our
repeatability analyses may lack power to detect repeatable behaviour; therefore, these results must be interpreted with caution. Nevertheless, our findings raise the possibility that there may be substantial phenotypic plasticity in jackdaw nest-building behaviour, in keeping with recent evidence that nest-building behaviour may be less 'fixed' than previously thought (Walsh et al., 2013). Indeed, we found intensity of behaviours varied significantly across 2 years, implying that environmental variables may affect behaviour and measures of reproductive success. Given that videos were recorded significantly closer to the lay date in 2019 than in 2018, it is possible that the effect of 'year' may reflect an effect of the proximity to the lay date; that is, the behaviour may change as the breeding season proceeds. However, this seems unlikely as the number of days the video had been recorded before the lay date had no effect on either nest building or vigilance in our analyses. Instead, our findings suggest that variation across years may be linked to differences in weather conditions or resource availability. Given that birds spent a relatively small proportion of observation time building their nests, the robustness of our estimates of between-year variation may be limited. While observation periods of 1 h can provide relatively accurate insights into parental behaviour at the nest (Murphy, Chutter, \& Redmond, 2015), longer durations and more observations across the nest-building phase may reveal clearer patterns of investment by the two sexes as well as temporal variation.

Some behaviours during nest construction were associated with proxies for reproductive success, raising the possibility that selection pressures may act on how pairs cooperate and how they spend their time. The relative contribution of females to bringing nest material was associated with an earlier lay date. Given that early laying can reduce competition for food when provisioning offspring and is often linked to elevated reproductive success in birds (Perrins, 1970), this suggests the female contribution to nest building may have important fitness consequences. We had hypothesized that more equal contributions by both partners could enable an earlier lay date by reducing the time and energy needed to build the nest (Prediction 5, 'division of labour', PCA $A_{\text {DoL }}$ ),
potentially important for females to save energy for costly egg production (Williams, 2005). Instead, we found that the time females spent building and the time males spent being vigilant were positively correlated, suggesting that greater investment in vigilance by the male may allow the female to invest more energy in building the nest and thus lay earlier. Contrary to our Prediction 3, the overall behavioural intensity and the relative investment in building behaviour compared to other behaviours ( $\mathrm{PCA}_{\text {All }}$ ) were not associated with any proxy for reproductive success. Similarly, we did not find support for our Prediction 4 that nest-building effort ( $\mathrm{PCA}_{\text {Effort }}$ ) would be linked to an earlier lay date. These results suggest that investing more time in nest building does not necessarily translate to increased fitness. Given that nest building is costly there may instead be advantages to building more efficiently or to starting building far in advance of laying and spreading the costs of building over a long period to minimize daily expenditures.

The amount of time partners spent together was also linked to fitness outcomes, but in the opposite direction to our Prediction 6. Whereas we had predicted that greater synchrony (more time spent together in the nestbox) would reflect compatibility between partners and be linked to reproductive benefits (Spoon et al., 2006), we actually found more synchronous pairs laid smaller eggs. One possible explanation for this is that the pairs that spent more time together in the nest were those that faced greater competition, as both partners are required to successfully guard a nest site in this species (Röell, 1978; Verhulst \& Salomons, 2004). Indeed, we found that pairs that spent more time together invested more time in vigilance but not in building the nest. Moreover, when additionally analysing the relationship between egg volume and vigilance directly, we found that birds that spent more time being vigilant had smaller first and third eggs. This suggests a competitive and stressful period where the need to defend the nestbox detracts from investment in nest building (Röell, 1978). There is evidence from other species, such as house sparrows, Passer domesticus, that investment in parental care, and consequent reproductive success, can be impaired by chronically elevated stress hormone levels (Ouyang, Sharp, Dawson, Quetting, \& Hau, 2011). While including morphological measures of female body condition did not improve our models, measures of current energetic and physiological state may prove more useful in future studies.

Together, our findings indicate that nest building in monogamous birds provides an important, but understudied, model system to investigate the evolution and proximate mechanisms of cooperation. How much a partner invests in nest building may be a source of information used by individual birds to assess how much their partner could be willing to invest later during the breeding attempt. This may be critical for individuals to estimate and to adjust their own effort. In the future, finer-scale analyses may also allow us to understand whether and how individuals respond strategically to each other's behaviour, for example by taking turns (cf. Johnstone \& Savage, 2019; Savage, Browning, Manica, Russell, \& Johnstone, 2017). Given growing evidence that nest building improves with experience (Muth \& Healy, 2011), it is also important to establish whether pairs learn and refine their cooperative nestbuilding strategies over time. Although there was little evidence that age was an important factor in our analyses, future work will be vital to determine whether and how the prior history of specific partners shapes their behaviour and reproductive success. Finally, investigations of nest building may also contribute to our understanding of animal architecture. Most research on cooperatively built architecture has focused on the nests and mounds of eusocial insects, where the colony is the unit of reproduction. Cooperative nest building in birds may provide useful opportunities to understand how variation in conflicts of interest influences the adaptive
value of cooperating to build structures for mutual benefit, and the proximate mechanisms through which this is achieved.

## Acknowledgments

We are grateful to the Gluyas family and staff at Pencoose Farm as well as to the people of Stithians who allowed us to carry out fieldwork. We thank Emily Cuff and Maude Quicray for collecting some of the video data in the field. This research was supported by a BBSRC David Phillips Fellowship (BB/H021817/2) and a Human Frontier Science Program grant (RG0049/2017) to A.T. R.H. was supported by a Natural Environment Research Council GW4 studentship (NERC 107672G).

## References

Bailey, I. E., Morgan, K. V., Bertin, M., Meddle, S. L., \& Healy, S. D. (2014). Physical cognition: Birds learn the structural efficacy of nest material. Proceedings of the Royal Society B: Biological Sciences, 281, 20133225. https://doi.org/10.1098/ rspb.2013.3225
Bailey, I. E., Morgan, K. V., Oschadleus, H. D., Deruiter, S. L., Meddle, S. L., \& Healy, S. D. (2016). Nest-building males trade off material collection costs with territory value. Emu-Austral Ornithology, 116(1), 1-8. https://doi.org/10.1071/ MU15022
Bailey, I. E., Muth, F., Morgan, K., Meddle, S. L., \& Healy, S. D. (2015). Birds build camouflaged nests. The Auk: Ornithological Advances, 132(1), 11-15. https:/| doi.org/10.1642/auk-14-77.1
Barta, Z., Székely, T., Liker, A., \& Harrison, F. (2014). Social role specialization promotes cooperation between parents. The American Naturalist, 183(6), 747-761. https://doi.org/10.5061/dryad.h4g74
Bates, D., Mächler, M., Bolker, B. M., \& Walker, S. C. (2015). Fitting linear mixedeffects models using lme4. Journal of Statistical Software, 67(1), 1-48. https:// doi.org/10.18637/jss.v067.i01
Birkhead, T. (2010). The magpies: the ecology and behaviour of black-billed and yellow-billed magpies. London, U.K.: T \& A D Poyser.
Bolker, B., \& R Core Team. (2017). bbmle: Tools for General Maximum Likelihood Estimation. R package version $1.0 .20 \mathrm{https}: / / c r a n . r-$ project.org/package= $=$ bbmle.
Borgia, G. (1995). Why do bowerbirds build bowers? American Scientist, 83(6), 542-547.
Breen, A. J., Guillette, L. M., \& Healy, S. D. (2016). What can nest-building birds teach us? Comparative Cognition and Behavior Reviews, 11(1), 83-102. https://doi.org/ 10.3819/ccbr.2016.110005

Brooks, M. E., Kristensen, K., van Benthem, K. J., Magnusson, A., Berg, C. W., Nielsen, A., et al. (2017). glmmTMB balances speed and flexibility among packages for zero-inflated generalized linear mixed modeling. $R$ Journal, 9(2), 378-400. https://doi.org/10.32614/rj-2017-066
Budaev, S. V. (2010). Using principal components and factor analysis in animal behaviour research: Caveats and guidelines. Ethology, 116(5), 472-480. https:|/ doi.org/10.1111/j.1439-0310.2010.01758.x
Campbell, B. L., Hurley, L. L., \& Griffith, S. C. (2018). Behavioural plasticity under a changing climate; how an experimental local climate affects the nest construction of the zebra finch Taeniopygia guttata. Journal of Avian Biology, 49(4), 1-8. https://doi.org/10.1111/jav. 01717
Chapman, T., Arnqvist, G., Bangham, J., \& Rowe, L. (2003). Sexual conflict. Trends in Ecology \& Evolution, 18(1), 41-47. https://doi.org/10.1016/S0169-5347(02) 00004-6
Cockburn, A. (2006). Prevalence of different modes of parental care in birds. Proceedings of the Royal Society B: Biological Sciences, 273, 1375-1383. https:/| doi.org/10.1098/rspb. 2005.3458
Collias, N. E. (1964). The evolution of nests and nest-building in birds. American Zoologist, 4(2), 175-190.
Collias, N. E., \& Collias, E. C. (1978). Cooperative breeding behavior in the whitebrowed sparrow weaver. The Auk, 95(3), 472-484. https://doi.org/10.1093/ AUK/95.3.472
Collias, N.E., \& Collias, E. C. (1984). Nest building and bird behavior. Princeton, NJ: Princeton University Press.
Cuff, E., \& Quicray, M. (2019). Prospecting behaviour of jackdaws. Unpublished raw data.
Davidson, G. L., Clayton, N. S., \& Thornton, A. (2014). Salient eyes deter conspecific nest intruders in wild jackdaws (Corvus monedula). Biology Letters, 10(2), 20131077. https://doi.org/10.1098/rsbl.2013.1077

Deeming, D. C., \& Reynolds, S. J. (Eds.). (2015). Nests, eggs, and incubation: new ideas about avian reproduction. Oxford, U.K.: Oxford University Press.
Diniz, P., MacEdo, R. H., \& Webster, M. S. (2019). Duetting correlates with territory quality and reproductive success in a suboscine bird with low extra-pair paternity. Auk, 136(1). https://doi.org/10.1093/auk/uky004
Disciullo, R. A., Thompson, C. F., \& Sakaluk, S. K. (2019). Perceived threat to paternity reduces likelihood of paternal provisioning in house wrens. Behavioral Ecology, 30(5), 1336-1343. https://doi.org/10.1093/beheco/arz082

Edwards, S. C., Shoot, T. T., Martin, R. J., Sherry, D. F., \& Healy, S. D. (2020). It's not all about temperature: Breeding success also affects nest design. Behavioral Ecology, 31(4), 1065-1072. https://doi.org/10.1093/beheco/araa052
Emery, N. J., Seed, A. M., Von Bayern, A. M. P., \& Clayton, N. S. (2007). Cognitive adaptations of social bonding in birds. Philosophical Transactions of the Royal Society B: Biological Sciences, 362(1480), 489-505. https://doi.org/10.1098/ rstb.2006.1991
Fletcher, K., \& Foster, R. (2010). Use of external biometrics to sex carrion crow Corvus corone, rook C. frugilegus and western jackdaw C. monedula in Northern England. Ringing and Migration, 25(1), 47-51. https://doi.org/10.1080/ 03078698.2010.9674414

Fox, J., \& Weisberg, S. (2011). An $\{R\}$ Companion to Applied Regression (2nd ed.). Thousand Oaks, CA: Sage http://socserv.socsci.mcmaster.ca/jfox/Books/ Companion.
Friard, O., \& Gamba, M. (2016). BORIS: A free, versatile open-source event-logging software for video/audio coding and live observations. Methods in Ecology and Evolution, 7(11), 1325-1330. https://doi.org/10.1111/2041-210X. 12584
Gill, L. F., van Schaik, J., von Bayern, A. M. P., \& Gahr, M. L. (2020). Genetic monogamy despite frequent extrapair copulations in 'strictly monogamous' wild jackdaws. Behavioral Ecology, 31(1), 247-260. https://doi.org/10.1093/beheco/arz185
Graham, M. H. (2003). Confronting multicollinearity in ecological multiple regression. Ecology, 84(11), 2809-2815. https://doi.org/10.1890/02-3114
Greggor, A. L., Spencer, K. A., Clayton, N. S., \& Thornton, A. (2017). Wild jackdaws' reproductive success and their offspring's stress hormones are connected to provisioning rate and brood size, not to parental neophobia. General and Comparative Endocrinology, 243, 70-77. https://doi.org/10.1016/j.ygcen.2016.11.006
Griffith, S. C. (2019). Cooperation and coordination in socially monogamous birds: Moving away from a focus on sexual conflict. Frontiers in Ecology and Evolution, 7, 1-15. https://doi.org/10.3389/fevo.2019.00455
Griffiths, R., Double, M. C., Orr, K., \& Dawson, R. J. G. (1998). A DNA test to sex most birds. Molecular Ecology, 7(8), 1071-1075. https://doi.org/10.1046/j.1365294x.1998.00389.x
Habig, B. (2020). The evolution of nest building behavior in Ploceidae weaverbirds. In The 57th Annual Conference of the Animal Behavior Society. July 28-31.
Hansell, M. (2000). Bird nests and construction behaviour. Cambridge, U.K.: Cambridge University Press.
Hansell, M. (2005). Animal architecture. Oxford, U.K.: Oxford University Press.
Hansell, M. (2007). Built by animals: the natural history of animal architecture. Oxford, U.K.: Oxford University Press.
Harrison, X. A. (2014). Using observation-level random effects to model overdispersion in count data in ecology and evolution. PeerJ, 2014(1), Article e616. https://doi.org/10.7717/peerj. 616
Harrison, F., Barta, Z., Cuthill, I., \& Székely, T. (2009). How is sexual conflict over parental care resolved? A meta-analysis. Journal of Evolutionary Biology, 22(9), 1800-1812. https://doi.org/10.1111/j.1420-9101.2009.01792.x
Harrison, X. A., Donaldson, L., Correa-Cano, M. E., Evans, J., Fisher, D. N., Goodwin, C. E. D., et al. (2018). A brief introduction to mixed effects modelling and multi-model inference in ecology. PeerJ, 6, Article e4794. https://doi.org/ 10.7717/peerj. 4794

Hartig, F. (2019). DHARMa: Residual diagnostics for hierarchical (multi-level/mixed) regression models. R package version 0.2 .6 https://cran.r-project.org/ package $=$ DHARMa.
Healy, S. D., Walsh, P. T., \& Hansell, M. (2008). Nest building by birds. Current Biology, 18(7), R271.
Henderson, I. G., \& Hart, P. J. B. (1993). Provisioning, parental investment and reproductive success in jackdaws Corvus monedula. Scandinavian Journal of Ornithology, 24(2), 142-148.
Hinde, C. A., \& Kilner, R. M. (2007). Negotiations within the family over the supply of parental care. Proceedings of the Royal Society B: Biological Sciences, 274(1606), 53-60. https://doi.org/10.1098/rspb.2006.3692
Hooper, R. (2019). Pair-bond strength in jackdaws, 2019 (Unpublished manuscript).
Horn, J. L. (1965). A rationale and test for the number of factors in factor analysis. Psychometrika, 30(2), 179-185. https://doi.org/10.1007/BF02289447
Hunt, G. R. (1996). Manufacture and use of hook-tools by New Caledonian crows. Nature, 379, 249-251.
Iserbyt, A., Farrell, S., Eens, M., \& Müller, W. (2015). Sex-specific negotiation rules in a costly conflict over parental care. Animal Behaviour, 100, 52-58. https:// doi.org/10.1016/j.anbehav.2014.11.014
Iserbyt, A., Fresneau, N., Kortenhoff, T., Eens, M., \& Müller, W. (2017). Decreasing parental task specialization promotes conditional cooperation. Scientific Reports, 7(1), 1-10. https://doi.org/10.1038/s41598-017-06667-1
Johnstone, R. A., Manica, A., Fayet, A. L., Stoddard, M. C., Rodriguez-Gironés, M. A., \& Hinde, C. A. (2014). Reciprocity and conditional cooperation between great tit parents. Behavioral Ecology, 25(1), 216-222. https://doi.org/10.1093/beheco/ art109
Johnstone, R. A., \& Savage, J. L. (2019). Conditional cooperation and turn-taking in parental care. Frontiers in Ecology and Evolution, 7, 1-11. https://doi.org/10.3389/ fevo.2019.00335
Krist, M. (2011). Egg size and offspring quality: A meta-analysis in birds. Biological Reviews, 86(3), 692-716. https://doi.org/10.1111/j.1469-185X.2010.00166.x
Leighton, G. M. (2014). Sex and individual differences in cooperative nest construction of sociable weavers Philetairus socius. Journal of Ornithology, 155(4), 927-935. https://doi.org/10.1007/s10336-014-1075-3
Lorenz, K. (1931). Beiträge zur Ethologie sozialer Corviden. Journal Für Ornithologie, 1, 67-127.

Lv, L., Zhang, Z., Groenewoud, F., Kingma, S. A., Li, J., van der Velde, M., et al. (2019). Extra-pair mating opportunities mediate parenting and mating effort trade-offs in a songbird. Behavioral Ecology, 31(2), 421-431. https://doi.org/10.1093/ beheco/arz204
Mainwaring, M. C., \& Hartley, I. R. (2013). The energetic costs of nest building in birds. Avian Biology Research, 6(1), 12-17. https://doi.org/10.3184/ 175815512 X13528994072997
Massoni, V., Reboreda, J. C., López, G. C., \& Florencia Aldatz, M. (2012). High coordination and equitable parental effort in the rufous hornero. The Condor, 114(3), 564-570. https://doi.org/10.1525/cond.2012.110135
McIvor, G.E., Goumas, M., Alothyqi, N., Troscianko, J., Thornton, A. (n.d.) Egg colour in jackdaws and its links to maternal condition, offspring growth, and survival. (Manuscript in preparation).
Metz, M., Klump, G. M., \& Friedl, T. W. P. (2009). Male nest-building behaviour and mating success in the red bishop (Euplectes orix). Behaviour, 146(6), 771-794. https://doi.org/10.1163/156853909X446208
Morton, F. B., \& Altschul, D. (2019). Data reduction analyses of animal behaviour: Avoiding kaiser's criterion and adopting more robust automated methods. Animal Behaviour, 149, 89-95. https://doi.org/10.1016/j.anbehav.2019.01.003
Murphy, M. T., Chutter, C. M., \& Redmond, L. J. (2015). Quantification of avian parental behavior: What are the minimum necessary sample times? Journal of Field Ornithology, 86(1), 41-50. https://doi.org/10.1111/jofo. 12087
Muth, F., \& Healy, S. D. (2011). The role of adult experience in nest building in the zebra finch, Taeniopygia guttata. Animal Behaviour, 82(2), 185-189. https:// doi.org/10.1016/j.anbehav.2011.04.021
Nickell, W. P. (1958). Variations in engineering features of the nests of several species of birds in relation to nest sites and nesting materials. Butler University Botanical Studies, 13(2), 121-139.
Orians, G. H. (1969). On the evolution of mating systems in birds and mammals. The American Naturalist, 103(934), 589-603. https://doi.org/10.1086/282628
Ouyang, J. Q., Sharp, P. J., Dawson, A., Quetting, M., \& Hau, M. (2011). Hormone levels predict individual differences in reproductive success in a passerine bird. Proceedings of the Royal Society B: Biological Sciences, 278, 2537-2545. https:// doi.org/10.1098/rspb.2010.2490
Parrot, D. (1995). Social organization and extra-pair behaviour in the European blackbilled magpie Pica pica [PhD thesis]. Sheffield, U.K: University of Sheffield.
Perrins, C. M. (1965). Population fluctuations and clutch size in the great tit, Parus major. Journal of Animal Ecology, 34(3), 601-647.
Perrins, C. M. (1970). The timing of birds' breeding seasons. Ibis, 112(2), 242-255. https://doi.org/10.1111/j.1474-919X.1970.tb00096.x
R Core Team. (2020). R: A language and environment for statistical computing. Vienna, Austria: R Foundation for Statistical Computing.
Remeš, V., Freckleton, R. P., Tökölyi, J., Liker, A., \& Székely, T. (2015). The evolution of parental cooperation in birds. Proceedings of the National Academy of Sciences of the United States of America, 112(44), 13603-13608. https://doi.org/10.1073/ pnas. 1512599112
Revelle, W. (2018). psych: Procedures for Personality and Psychological Research. https://cran.r-project.org/package=psych Version $=1.8 .12$.
Röell, A. (1978). Social behaviour of the jackdaw Corvus monedula, in relation to its niche [PhD thesis]. the Netherlands: University of Groningen.
Rosell, F., Bozser, O., Collen, P., \& Parker, H. (2005). Ecological impact of beavers Castor fiber and Castor canadensis and their ability to modify ecosystems. Mammal Review, 35(3-4), 248-276. https://doi.org/10.1111/j.1365-2907.2005.00067.x
Sakia, R. M. (1992). The box-cox transformation technique: A review. The Statistician, 41(2), 169. https://doi.org/10.2307/2348250
Savage, J. L., Browning, L. E., Manica, A., Russell, A. F., \& Johnstone, R. A. (2017). Turntaking in cooperative offspring care: By-product of individual provisioning behavior or active response rule? Behavioral Ecology and Sociobiology, 71, 162. https://doi.org/10.1007/s00265-017-2391-4
Soler, J. J., Møller, A. P., \& Soler, M. (1998). Nest building, sexual selection and parental investment. Evolutionary Ecology, 12(4), 427-441. https://doi.org/ 10.1023/A:1006520821219

Spoon, T. R., Millam, J. R., \& Owings, D. H. (2006). The importance of mate behavioural compatibility in parenting and reproductive success by cockatiels, Nymphicus hollandicus. Animal Behaviour, 71(2), 315-326. https://doi.org/ 10.1016/j.anbehav.2005.03.034

Stoffel, M. A., Nakagawa, S., \& Schielzeth, H. (2017). rptR: repeatability estimation and variance decomposition by generalized linear mixed-effects models. Methods in Ecology and Evolution, 8, 1639-1644. https://doi.org/10.1111/2041210X. 12797
Troscianko, J. (2014). A simple tool for calculating egg shape, volume and surface area from digital images. Ibis, 156(4), 874-878. https://doi.org/10.1111/ibi.12177
Venables, W. N., \& Ripley, B. D. (2002). Modern applied statistics with S (4th ed.). New York, NY: Springer.
Verhulst, S., Geerdink, M., Salomons, H. M., \& Boonekamp, J. J. (2014). Social life histories: Jackdaw dominance increases with age, terminally declines and shortens lifespan. Proceedings of the Royal Society B: Biological Sciences, 281, 20141045. https://doi.org/10.1098/rspb.2014.1045

Verhulst, S., \& Salomons, H. M. (2004). Why fight? Socially dominant jackdaws, Corvus monedula, have low fitness. Animal Behaviour, 68(4), 777-783. https:// doi.org/10.1016/j.anbehav.2003.12.020
Walsh, P. T., Hansell, M., Borello, W. D., \& Healy, S. D. (2013). Are elaborate bird nests built using simple rules? Avian Biology Research, 6(2), 157-162. https://doi.org/ 10.3184/175815513X13629302805186

Wechsler, B. (1989). Measuring pair relationships in jackdaws. Ethology, 80(1-4), 307-317. https://doi.org/10.1111/j.1439-0310.1989.tb00749.x
West, B., Welch, K., \& Galecki, A. (2014). Linear mixed models: A practical guide using statistical software. Journal of the American Statistical Association, 103(481), 427-428.
Williams, T. D. (2005). Mechanisms underlying the costs of egg production. BioScience, 55(1), 39-48.
Zann, R. A. (1996). The zebra finch: a synthesis of field and laboratory studies. Oxford, U.K.: Oxford University Press.

## Appendix

Table A1
Number of observations per pair in 2018 and 2019

| Pair ID | 2018 | 2019 |
| :---: | :---: | :---: |
| J1313UNM.X33.18 | 1 | 0 |
| J1318J494 | 0 | 2 |
| J1319J416 | 1 | 1 |
| J1323J1340 | 2 | 1 |
| J1337J1259 | 2 | 1 |
| J1342J293 | 1 | 0 |
| J1349J1896 | 2 | 2 |
| J1366J1890 | 2 | 0 |
| J1469J1388 | 0 | 1 |
| J164J831 | 2 | 1 |
| J1876J1888 | 1 | 0 |
| J1903J469 | 0 | 1 |
| J1908J1458 | 0 | 1 |
| J1915J1507 | 1 | 1 |
| J1979J1131B | 1 | 0 |
| J218J289 | 0 | 1 |
| J2375J837 | 0 | 1 |
| J2377J1463 | 0 | 1 |
| J2394J219 | 1 | 0 |
| J2403J2965 | 1 | 0 |
| J2409J2402 | 0 | 1 |
| J250J290 | 0 | 1 |
| J262J838 | 1 | 0 |
| J2951J2966 | 1 | 0 |
| J2979J2406 | 1 | 0 |
| J39J267 |  | 1 |
| J407J1359 | 1 | 0 |
| J41J323 | 1 | 1 |
| J42J1351 | 1 | 0 |
| J43J265 | 1 | 0 |
| J505J1480 | 1 | 1 |
| J510J516 | 0 | 1 |
| J572J257 | 1 | 0 |
| J692J899 | 1 | 1 |
| J792J1895 | 1 | 1 |
| J793J1352 | 1 | 1 |
| J796J1346 | 0 | 1 |
| J803J266 | , | 0 |
| J908J587 | 1 | 1 |
| J912J1360 | 1 | 1 |

The ID of the first pair indicates that the male was unmarked.

Table A2
Loadings of behavioural variables onto the first two principal components of the PCA $_{\text {All }}$ including nine different behaviours $(N=59)$

| Behaviour | PC1 | PC2 |
| :--- | :--- | :--- |
| In box (cumulative) | -0.456 | 0.175 |
| In box (both birds) | -0.354 | 0.391 |
| Build nest | -0.344 | -0.385 |
| Material | -0.309 | -0.449 |
| Material out | -0.224 | -0.478 |
| Form cup | -0.266 | -0.226 |
| Vigilance | -0.380 | 0.390 |
| Chatter | -0.297 | 0.048 |
| Calls | -0.317 | 0.180 |
| Variation explained | 46.54 | 25.22 |

Table A3
Loadings of behavioural variables related to nest building onto $\mathrm{PC}_{\text {Efffort }}$ and $P C 2_{\text {Efffort }}$ of the PCA Efffort $(N=61)$

| Behaviour | PC1 | PC2 |
| :--- | :--- | :--- |
| Build nest | -0.541 | 0.017 |
| Material | -0.543 | -0.123 |
| Material out | -0.496 | -0.558 |
| Form cup | -0.408 | 0.821 |
| Variation explained | 69.18 | 18.86 |

Table A4
Loadings of behavioural variables (relative female contribution) onto PC1 and PC2 of the PCA Dol $(N=61)$

| Behaviour | PC1 | PC2 |
| :--- | :--- | :--- |
| Build nest ㅇ | 0.700 | -0.118 |
| Material 9 | 0.116 | 0.992 |
| Vigilance 9 | -0.704 | 0.046 |
| Variation explained | 58.16 | 33.06 |

Table A5
Behaviours (state events) jackdaws showed in their nestbox ( $N=62$ observations; $N=60$ for chatter)

| Behaviour | \% Time (out of observation length) <br> Mean $\pm$ SD | \% Time (out of time in box) <br> Mean $\pm$ SD | Females: \% time (observation length) <br> Mean $\pm$ SD | Males: \% time (observation length) <br> Mean $\pm$ SD |
| :--- | :--- | :--- | :--- | :--- |
| In box (cumulative) | $37.96 \pm 30.50$ | - | $16.06 \pm 15.99$ | $15.50 \pm 15.04$ |
| Box occupied | $29.09 \pm 19.64$ | - | - | - |
| In box (both) | $8.99 \pm 13.56$ | $23.17 \pm 24.58$ | - | $7.57 \pm 10.47$ |
| Vigilance | $15.73 \pm 19.24$ | $37.83 \pm 19.32$ | $5.58 \pm 8.69$ | $3.05 \pm 3.48$ |
| Build nest | $9.59 \pm 8.39$ | $27.34 \pm 16.92$ | $4.66 \pm 4.95$ | $0.60 \pm 0.68$ |
| Form cup | $1.67 \pm 1.70$ | $4.71 \pm 4.01$ | $0.74 \pm 0.90$ | $0.003 \pm 0.014$ |
| Modify | $0.05 \pm 0.13$ | $0.15 \pm 0.52$ | $0.024 \pm 0.075$ | $0.33 \pm 0.77$ |
| Chatter | $0.92 \pm 1.70$ | $1.93 \pm 2.50$ | $0.33 \pm 0.80$ |  |

The second and third column summarize the percentage of time pairs showed each behaviour. The last two columns indicate the amount of time (percentage of observation length) both sexes exhibited a particular behaviour. The behaviours of the sexes do not always add up to the cumulative amount because in some instances a bird was not identifiable.

Table A6
Behaviours (point events) shown by jackdaw pairs ( $N=62$ observations)

| Behaviour | No. of events/observation <br> Mean $\pm$ SD | No. of events/h <br> Mean $\pm S D$ | Females: no. of events/h <br> Mean $\pm$ SD |
| :--- | :--- | :--- | :--- |
| Visits | $32.53 \pm 28.01$ | $10.06 \pm 8.07$ | $3.71 \pm 3.13$ |
| Material | $18.89 \pm 20.99$ | $5.71 \pm 6.09$ | $2.12 \pm 2.42$ |
| Material out | $4.68 \pm 7.93$ | $1.44 \pm 2.46$ | $0.67 \pm 1.42$ |
| Calls | $18.05 \pm 23.85$ | $5.52 \pm 5.85$ | $2.83 \pm 3.84$ |
| Food sharing $\left(\delta^{\top}\right)$ | $0.27 \pm 0.48$ | $0.09 \pm 0.18$ | - |

The second and third columns describe the total number of events per observation and per hour, respectively. The last two columns summarize the number of events per hour for both sexes separately. The behaviours of the sexes do not add up to the cumulative amount, as individuals were sometimes unidentifiable.


Figure A1. The number of pieces of material brought by individual birds (rounded, per hour) plotted against their age (horizontal jitter was used to make data points more distinguishable). The continuous fitted line corresponds to the model output; dotted lines show the $95 \%$ confidence intervals.

Table A7
Sex differences in behaviour

| Response variable (model) | Fixed effects | $\beta$ | SE ( $\beta$ ) | $\mathrm{X}^{2}$ | $d f$ | $\begin{aligned} & 95 \% \\ & \mathrm{CI} \text { (lower) } \end{aligned}$ | $\begin{aligned} & 95 \% \\ & \text { CI (upper) } \end{aligned}$ | P |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| In box (LMM) | Intercept | 2.682 | 0.183 |  | 1 |  |  |  |
|  | Sex | 0.017 | 0.162 | 0.011 | 1 | -0.301 | 0.335 | 0.916 |
|  | Year | -1.003 | 0.191 | 27.579 | 1 | -1.376 | -0.630 | < 0.001 |
| Random effects: Pair ID $\left(\sigma^{2}, \sigma\right)=0.498,0.706 ;$ Site $\left(\sigma^{2}, \sigma\right)=0.000,0.000$ |  |  |  |  |  |  |  |  |
| Build nest (LMM) | Intercept | 1.603 | 0.130 |  | 1 |  |  |  |
|  | Sex | -0.237 | 0.112 | 4.522 | 1 | -0.457 | -0.018 | 0.033 |
|  | Year | -0.580 | 0.133 | 19.008 | 1 | -0.845 | -0.319 | < 0.001 |
| Random effects: Pair ID $\left(\sigma^{2}, \sigma\right)=0.271,0.520$; Site $\left(\sigma^{2}, \sigma\right)=0.000,0.000$ |  |  |  |  |  |  |  |  |
| Vigilance (LMM) | Intercept | 1.591 | 0.161 |  | 1 |  |  |  |
|  | Sex | 0.259 | 0.138 | 3.520 | 1 | -0.012 | 0.530 | 0.060 |
|  |  | -0.787 | 0.165 | 22.795 | 1 | -1.109 | -0.464 | < 0.001 |
| Random effects: Pair ID $\left(\sigma^{2}, \sigma\right)=0.423,0.650$; Site $\left(\sigma^{2}, \sigma\right)=0.000,0.000$ |  |  |  |  |  |  |  |  |
| Build vs vigilance (LMM) | Intercept |  | 0.138 |  | 1 |  |  |  |
|  | Sex | -0.237 | 0.133 | 0.013 | 1 | -0.498 | 0.023 |  |
|  | Behaviour | -0.025 | 0.133 | $5.590$ | 1 | $-0.286$ | $0.235$ | $0.018$ |
|  | Sex*Behaviour | 0.497 | 0.189 | 6.924 | 1 | 0.128 | 0.865 | 0.009 |
|  | Year | -0.678 | 0.116 | 34.267 | 1 | -0.904 | - 0.453 | < 0.001 |
| Random effects: Pair ID $\left(\sigma^{2}, \sigma\right)=0.292,0.540 ;$ Site $\left(\sigma^{2}, \sigma\right)=0.000,0.000$ |  |  |  |  |  |  |  |  |
| Visits (GLMM) | Intercept | 1.567 | 0.130 |  | 1 |  |  |  |
|  | Sex | 0.001 | 0.121 | 0.000 | 1 | -0.237 | 0.238 | 0.995 |
|  |  |  | 0.162 | 37.655 | 1 | -1.313 | -0.677 | < 0.001 |
| Random effects: Pair ID $\left(\sigma^{2}, \sigma\right)=0.212 ; 0.461$, Site ( $\left.\sigma^{2}, \sigma\right)=0.000,0.000$ |  |  |  |  |  |  |  |  |
| Material (GLMM) | Intercept | 0.855 | 0.197 |  | 1 |  |  |  |
|  | Sex | 0.513 | 0.716 | 0.046 | 1 | -0.889 | 1.916 | 0.830 |
|  | Age | 0.257 | 0.124 | 4.135 | 1 | 0.013 | 0.500 | 0.042 |
|  | Sex*Age | -0.108 | 0.134 | 0.650 | 1 | -0.372 | 0.155 | $0.420$ |
|  |  |  | 0.289 | 30.311 | 1 | -2.154 | -1.023 | < 0.001 |
| Random effects: Pair ID $\left(\sigma^{2}, \sigma\right)=0.405,0.636$; Site $\left(\sigma^{2}, \sigma\right)=0.000,0.000$ |  |  |  |  |  |  |  |  |
| Material out (GLMM) | Intercept | -1.560 | 0.483 |  | 1 |  |  |  |
|  | Sex | -0.390 | 0.480 | $0.658$ | 1 | -1.331 | $0.552$ | $0.417$ |
|  | Year | -1.080 | 0.514 | 4.422 | 1 | -2.087 | -0.073 | 0.035 |
| Random effects: Pair ID $\left(\sigma^{2}, \sigma\right)=<0.001,<0.001$; Site $\left(\sigma^{2}, \sigma\right)=0.000,0.000$ |  |  |  |  |  |  |  |  |
| Form cup (LMM) | Intercept | $0.607$ |  |  | 1 |  |  |  |
|  | Sex | -0.061 | 0.053 | 1.340 | 1 | -0.165 | 0.042 | 0.247 |
|  | Year | -0.320 | 0.063 | 25.843 | 1 | -0.443 | -0.197 | < 0.001 |
| Random effects: Pair ID $\left(\sigma^{2}, \sigma\right)=0.062,0.248 ;$ Site $\left(\sigma^{2}, \sigma\right)=0.000,0.000$ |  |  |  |  |  |  |  |  |
| Chatter (LMM) | Intercept | 0.296 | 0.051 |  | 1 |  |  |  |
|  | Sex | 0.006 | 0.052 | 0.015 | 1 | -0.095 | 0.108 | 0.901 |
|  |  |  | 0.057 | 15.046 | 1 | -0.333 | -0.110 | < 0.001 |
| Random effects: Pair ID $\left(\sigma^{2}, \sigma\right)=0.022,0.147$; Site $\left(\sigma^{2}, \sigma\right)=0.000,0.000$ |  |  |  |  |  |  |  |  |
| Calls (GLMM) | Intercept | 1.026 | 0.239 |  | 1 |  |  |  |
|  | Sex | -0.717 | 0.233 | 9.466 | 1 | -1.175 | -0.261 | 0.002 |
|  | Year | $-0.582$ | 0.276 | 4.467 | 1 | -1.122 | -0.042 | 0.035 |
| Random effects: Pair ID ( $\sigma^{2}$, | 001, 0.026; Site | 0.000, 0. |  |  |  |  |  |  |

CI: confidence interval. Statistically significant results are in bold. Response variables for LMMs were log transformed and estimates for GLMMs (Conway-Maxwell-Poisson) are on the $\log$ scale. Intercepts refer to the values of females and to the year 2018. Observation level random effects accounted for zero inflation. $\sigma^{2}$ and $\sigma$ denote the variation and standard deviation attributed to random effects. Sex differences were not examined for rarer behaviours (modification of nest material).

Table A8
Relationship between behaviours shown by jackdaws and age

| Response variable (model) | Fixed effects | $\beta$ | SE ( $\beta$ ) | $\mathrm{X}^{2}$ | df | 95\% CI (lower) | 95\% CI (upper) | $P$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| In box (LMM) | Age | 0.091 | 0.112 | 0.398 | 1 | -0.127 | 0.310 | 0.528 |
|  | Sex*Age | -0.059 | 0.116 | 0.264 | 1 | -0.285 | 0.165 | 0.607 |
| Build nest (LMM) | Age | 0.054 | 0.079 | 0.573 | 1 | -0.100 | 0.208 | 0.449 |
|  | Sex*Age | -0.011 | 0.080 | 0.017 | 1 | -0.166 | 0.145 | 0.895 |
| Vigilance (LMM) | Age | 0.050 | 0.099 | 0.036 | 1 | -0.140 | 0.241 | 0.849 |
|  | Sex*Age | -0.060 | 0.099 | 0.367 | 1 | -0.251 | 0.132 | 0.545 |
| Build vs vigilance (LMM) | Age | 0.027 | 0.059 | 0.205 | 1 | -0.088 | 0.141 | 0.651 |
| Visits (GLMM) | Age | 0.165 | 0.081 | 2.380 | 1 | 0.005 | 0.324 | 0.123 |
|  | Sex*Age | -0.111 | 0.085 | 1.712 | 1 | -0.278 | 0.055 | 0.191 |
| Material out (GLMM) | Age | 0.222 | 0.243 | 0.275 | 1 | -0.255 | 0.698 | 0.600 |
|  | Sex*Age | -0.247 | 0.322 | 0.590 | 1 | -0.878 | 0.383 | 0.442 |
| Form cup (LMM) | Age | 0.048 | 0.037 | 1.380 | 1 | -0.025 | 0.120 | 0.240 |
|  | Sex*Age | -0.022 | 0.038 | 0.347 | 1 | -0.096 | 0.051 | 0.556 |
| Chatter (LMM) | Age | 0.059 | 0.032 | 2.112 | 1 | -0.002 | 0.121 | 0.146 |
|  | Sex*Age | -0.043 | 0.036 | 1.389 | 1 | -0.113 | 0.027 | 0.239 |
| Calls (GLMM) | Age | -0.044 | 0.148 | 1.046 | 1 | -0.334 | 0.247 | 0.306 |
|  | Sex*Age | 0.280 | 0.169 | 2.378 | 1 | -0.051 | 0.611 | 0.098 |

CI: confidence interval.

Table A9
Repeatability estimates for different behaviours of 16 pairs that were measured repeatedly

| Behaviour | Repeatability | SE | $2.5 \% \mathrm{CI}$ | $97.5 \% \mathrm{CI}$ |
| :--- | :--- | :--- | :--- | :--- | :--- |
| In box (cumulative) (SE) | 0 | 0.119 | 0 | 0.391 |
| In box (both birds) (SE) | 0 | 0.114 | 0.387 |  |
| Vigilance (SE) | 0 | 0.113 | 0.365 |  |
| Nest build (SE) | 0.080 | 0.140 | 0 | 0.99 |
| Material | 0 | 0.091 | 0.956 |  |
| Material out | 0.128 | 0.176 | 0.9 | 0.313 |
| Form cup (SE) | 0 | 0.118 | 0.99 |  |
| Chatter (SE) | 0 | 0.108 | 0.384 |  |
| Calls | 0 | 0.082 | 0 | 0.374 |
| Food sharing | Not converged | - | 0 | 0.397 |

CI: confidence interval. State events (SE) were Box-Cox transformed to approximate assumptions for Gaussian data.

Table A10
Repeatability estimates for different correlates of reproductive success

| Fitness measure | Repeatability | SE | $2.5 \% \mathrm{CI}$ | $97.5 \% \mathrm{CI}$ |  |
| :--- | :--- | :--- | :--- | :--- | :--- |
| Clutch size | 0 | 0.078 | 0 | 0.278 |  |
| Relative lay date | 0.643 | 0.202 | 0.025 | 0.826 |  |
| Volume first egg | 0.598 | 0.184 | 0.130 | 0.843 |  |
| Volume third egg | 0.531 | 0.193 | 0.045 | $\mathbf{0 . 0 2 3}$ | $\mathbf{0 . 8 9}$ |

CI: confidence interval.
Statistically significant results are in bold.

Table A11
Different models to examine the effect of behaviours on proxies for reproductive success

| Response variable | Concept (models) | Fixed effects | Error structure | Best model ( $\Delta$ AICc) | $\beta \pm$ SE ( $\beta$ ) | P |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Relative lay date | PCA ${ }_{\text {All }}$ (5) | $\mathrm{PC} 1_{\text {All }}, \mathrm{PC} 2_{\text {All }}$ | GLMM: COM-Poisson | Year, Day (3.3) | $-0.012 \pm 0.012$ | 0.286 |
|  | PCA $A_{\text {effort }}$ (1) | PC1 $1_{\text {effort }}$ |  |  |  |  |
|  | PCA ${ }_{\text {DoL }}$ (6) | PC1 $1_{\text {DoL }}$, Material by ${ }^{\text {f }}$ |  | Material 9 (4.7) |  |  |
|  | Synchrony (1) | Together in box |  |  | $0.002 \pm 0.003$ | 0.546 |
| Clutch size | PCA All (5) | $\mathrm{PC} 1_{\text {All }}, \mathrm{PC} 2_{\text {All }}$ | GLMM: COM-Poisson | Null model (0.0) | $0.002 \pm 0.003$ |  |
|  | PCA $\mathrm{Effort}_{\text {(1) }}$ | PC1 $1_{\text {Effort }}$ |  |  | $0.005 \pm 0.017$ | 0.753 |
|  | PCA ${ }_{\text {DoL }}$ (6) | PC1 ${ }_{\text {DoL }}$, Material by ${ }^{\text {f }}$ |  | Null model (3.6) |  |  |
|  | Synchrony (1) | Together in box |  |  | $-0.002 \pm 0.002$ | 0.424 |
| Volume of first egg | $\mathrm{PCA}_{\text {All }}(5)$ | PC1 $1_{\text {All }}, \mathrm{PC} 2_{\text {All }}$ | LMM: Gaussian | Null model (3.2) | $-0.032 \pm 0.071$ | 0.656 |
|  | PCA Effort $^{\text {(1) }}$ | PC1 $1_{\text {Effort }}$ |  |  |  |  |
|  | PCA ${ }_{\text {DoL }}$ (6) | $\mathrm{PC} 1_{\text {DoL }}$, Material by ${ }_{\text {o }}$ |  | Null model (3.4) |  |  |
|  | Synchrony (1) | Together in box |  |  | $-0.018 \pm 0.008$ | 0.019 |
| Volume of third egg | $\mathrm{PCA}_{\text {All }}$ (5) | $\mathrm{PC1} 1_{\text {All }}, \mathrm{PC} 2_{\text {All }}$ | LMM: Gaussian | Null model (9.8) | $-0.063 \pm 0.087$ | 0.469 |
|  | PCA Effort $^{\text {(1) }}$ | PC1 ${ }_{\text {Effort }}$ |  |  |  |  |
|  | PCA ${ }_{\text {DoL }}$ (6) | PC1 ${ }_{\text {DoL }}$, Material by ${ }^{\text {f }}$ |  | Null model (8.5) |  |  |
|  | Synchrony (1) | Together in box |  |  | $-0.026 \pm 0.010$ | 0.007 |

COM-Poisson: Conway-Maxwell-Poisson. In all models we accounted for the 'year', ‘day', 'food sharing' and 'female age' as fixed effects. 'Pair ID' and 'site’ were included as random effects in all models. The column 'best model' shows which model had the lowest AICc in cases where we performed model selection. The last two columns show the estimate, SE and $P$ value for the instances where we did not use AICc but constructed single models. Statistically significant results are in bold.

Table A12
Summary of the statistical analyses on the relationship between behaviours at the nest building phase and fitness proxies

| Response variable (model) | Fixed effects | $\beta$ | SE ( $\beta$ ) | $\mathrm{X}^{2}$ | df | 95\% CI (lower) | 95\% CI (upper) | P |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Relative lay date (GLMM) | Intercept | 1.168 | 0.415 |  | 1 |  |  |  |
|  | Material (9) | -0.388 | 0.167 | 5.395 | 1 | -0.715 | -0.061 | 0.020 |
|  | Food sharing ( ${ }^{\text {a }}$ ) | -0.546 | 0.299 | 3.330 | 1 | -1.131 | 0.040 | 0.068 |
|  | Year | -0.124 | 0.126 | 0.983 | 1 | -0.371 | 0.122 | 0.322 |
|  | Day | 0.020 | 0.007 | 8.140 | 1 | 0.006 | 0.034 | 0.004 |
|  | Random effect: Pair ID $\left(\sigma^{2}, \sigma\right)=0.318,0.564 ;$ Site $\left(\sigma^{2}, \sigma\right)<0.001,<0.001$ |  |  |  |  |  |  |  |
| Volume of first egg (LMM) | Intercept | 11.814 | 0.249 |  | 1 |  |  |  |
|  | Synchrony | -0.019 | 0.008 | 5.793 | 1 | -0.034 | -0.004 | 0.016 |
|  | Food sharing ( ${ }^{\text {a }}$ ) | 0.746 | 0.592 | 1.587 | 1 | -0.418 | 1.889 | 0.282 |
|  | Year | -0.329 | 0.186 | 3.118 | 1 | -0.685 | 0.026 | 0.077 |
|  | Day | 0.034 | 0.017 | 4.145 | 1 | 0.002 | 0.066 | 0.042 |
|  | Random effect: Pair ID $\left(\sigma^{2}, \sigma\right)=0.347,0.589$; Site $\left(\sigma^{2}, \sigma\right)<0.001,<0.001$ |  |  |  |  |  |  |  |
| Volume of third egg (LMM) | Intercept | 11.291 | 0.578 |  | 1 |  |  |  |
|  | Synchrony | -0.027 | 0.010 | 7.486 | 1 | -0.048 | -0.007 | 0.006 |
|  | Food sharing ( ${ }^{\text {® }}$ ) | 0.061 | 0.747 | 0.008 | 1 | -1.330 | 1.464 | 0.929 |
|  | Year | -0.211 | 0.282 | 0.557 | 1 | -0.663 | 0.243 | 0.455 |
|  | Day | 0.023 | 0.021 | 1.236 | 1 | -0.016 | 0.063 | 0.266 |
|  | Random effect: Pair ID $\left(\sigma^{2}, \sigma\right)=0.450,0.671$; Site $\left(\sigma^{2}, \sigma\right)<0.001,<0.001$ |  |  |  |  |  |  |  |

The fixed effect 'Material ( ()' reflects the relative female contribution to bringing nest material relative to the overall effort by both sexes. The reference year was 2018 and 'day' refers to the number of days the video was recorded before the lay date of the first egg. $\sigma^{2}$ and $\sigma$ show the variation and SD explained by random effects. Statistically significant results are shown in bold.


[^0]:    * Corresponding authors.

    E-mail addresses: luca.hahn8@gmail.com (L. G. Hahn), alex.thornton@exeter.ac. uk (A. Thornton).

