INTRODUCTION

Across socially monogamous species, levels of extra-pair paternity (EPP) show that infidelity (i.e., extra-pair mating) occurs frequently, yet the evolution of this behaviour remains enigmatic (Griffith, Owens, & Thuman, 2002; Taylor, Price, & Wedell, 2014; Westneat & Stewart, 2003). Despite being associated with costs, extra-pair mating is expected to benefit males by increasing their reproductive success (Jennions & Petrie, 2000). However, in females, extra-pair fertilizations do not necessarily increase immediate reproductive success, so if and how females benefit from infidelity is still unclear, despite considerable research and debate (Forstmeier, Nakagawa, & Stewart, 2003).
Indeed, very few studies have attempted to disentangle within- from consistently different ability to gain EPP (van de Pol & Verhulst, 2006). The reproductive population, respectively) of individuals with converging age and (b) between-individual changes, driven by the selection, namely: (a) within-individual changes in EPP with age, but cannot distinguish the processes that may underlie such analyses, which can capture population-level associations between age and EPP, whereas little is known about how the rate of extra-pair reproduction changes within individuals with age, even though this may be key to understanding the benefits of infidelity for females.

According to the influential “good genes” hypothesis, extra-pair reproduction enables socially constrained females to acquire higher quality paternal genes for their young (Hamilton & Zuk, 1982). Furthermore, if male age reflects good genes via demonstrated viability (Kokko, 1998; Trivers, 1972), females should seek extra-pair fertilizations from older males, especially when paired with young males. This should result in between-male differences in paternity gain and loss in relation to age (i.e., higher EPP success and lower within-pair paternity loss in older males). Alternatively, the “competitive ability” hypothesis (Nakagawa, Schroeder, & Burke, 2015) posits that males increase their ability to gain paternity as they age. This may be due to physiological changes (e.g., improved body/ejaculate condition) or experience-enhanced behavioural changes (e.g., improved mate-guarding, timing of copulations, increased ability to force copulations; Curio, 1983; Hsu et al., 2015; Morton et al., 1990; Westneat & Stewart, 2003). Contrary to the good genes hypothesis, the competitive ability hypothesis predicts a within-male age effect and does not imply any indirect genetic benefits for females. However, these two hypotheses are not mutually exclusive because both within- and between-individual differences in EPP in relation to age may occur at the same time.

Most research on EPP so far has relied on cross-sectional analyses, which can capture population-level associations between age and EPP, but cannot distinguish the processes that may underlie such associations, namely: (a) within-individual changes in EPP with advancing age and (b) between-individual changes, driven by the selective appearance/disappearance (i.e., the age of entry into/exit from the reproductive population, respectively) of individuals with consistently different ability to gain EPP (van de Pol & Verhulst, 2006). Indeed, very few studies have attempted to disentangle within- from between-individual effects on EPP (Hsu et al., 2017; Schroeder et al., 2016). Clearly, more longitudinal studies are needed if we are to understand the factors that shape male age-dependent variation in EPP and, therefore, better understand the evolution of infidelity.

The relationship between age and extra-pair reproduction in females remains markedly understudied and unclear. Many reasons have been suggested as to why females may seek extra-pair copulations, including the acquisition of direct benefits (e.g., fertility assurance; Sheldon, 1994) or indirect genetic benefits (e.g., high-quality or compatible genes in offspring; Brown, 1997; Hamilton & Zuk, 1982; Zeh & Zeh, 1996). Older females may have fewer extra-pair offspring because they are more capable of obtaining a better quality social male (Wagner, Schug, & Morton, 1996), and thus do not need to seek extra-pair copulations. Alternatively, they may be more experienced at avoiding or resisting unwanted copulation attempts (Morton & Derrickson, 1990). On the other hand, older females may have more extra-pair offspring because they are better at avoiding mate-guarding, and at obtaining copulations with other males—for “good genes” or other reasons (Bouwman & Komdeur, 2005). Additionally, older females may be more likely to produce extra-pair offspring because they are better at overcoming constraints imposed by male retaliation to perceived paternity loss (the “constrained female” hypothesis; Dixon, Ross, O’Malley, & Burke, 1994; Gowaty, 1996). The few studies that have investigated the relationship between female age and the production of extra-pair offspring have provided contrasting results, showing a positive relationship (Bouwman & Komdeur, 2005; Dietrich, Schmoll, Winkel, Epplen, & Lubjuhn, 2004; Kempenaers, Congdon, Boag, & Robertson, 1999), a negative relationship (Moreno et al., 2015; Ramos et al., 2014; Stutchbury et al., 1997) or no relationship (Lubjuhn, Gerken, Brün, & Schmoll, 2007; Wagner et al., 1996). However, none of these studies distinguished within- and between-individual age effects.

Senescence—the progressive deterioration in performance in late life (Medawar, 1952; Williams, 1957)—is an interesting and important within-individual process related to age. There have been numerous studies assessing the fitness consequences of senescence, most of which have focused on declines in survival and reproduction with age (reviewed by Nussey, Froy, Lemaître, Gaillard, & Austad, 2013). However, to our knowledge only one study has addressed, albeit not explicitly, senescence in EPP (Hsu et al., 2017), and has focused only on males.

A within-individual senescent decline in EPP success in late life is compatible with both the good genes and the competitive ability hypotheses. Specifically, in a good genes scenario, the oldest males are the most attractive because they are of highest intrinsic quality (as evidenced by highest longevity) and thus are preferred by females; however, if senescence causes lower fertilization ability (e.g., because of lower sperm competitiveness; Dean et al., 2010), very old males may gain less extra-pair (and total) paternity than younger (less attractive) males. In this case, annual EPP success would be impacted by both a between-individual age effect, represented by a positive association between annual EPP and longevity (because males that live longer are preferred by females),
and by a within-individual age effect, resulting in a decline in EPP in all males in late life (because at very old ages males are in lower physical condition). According to the competitive ability hypothesis, all males initially increase their reproductive success as they age (due to increasing experience and/or body condition) but, if senescence occurs, this initial increase in reproduction with age is expected to turn into a decline in late life. In this case, annual EPP would be predicted only by a within-individual age effect, resulting in an increase in EPP at young ages followed by a decline at old ages (while in the absence of senescence EPP should asymptote when added experience, or condition, does not lead to further improvements in the ability to gain EPP). Given that the acquisition of EPP may change with age and/or experience, and also show senescence, it is likely that the contribution of EPP to total reproductive success will vary considerably with age. Numerous studies have investigated how EPP alters male reproductive success (e.g., Albrecht et al., 2007; Lebigre, Arcese, Sardell, Keller, & Reid, 2012), but only a few have done so in relation to age (e.g., Girndt, Chng, Burke, & Schroeder, 2018; Hsu et al., 2017). To our knowledge, only one of these studies disentangled within- and between-individual age effects (Hsu et al., 2017), although this study did not explicitly test for post-peak senescence in EPP.

Here, we investigate extra-pair offspring production in relation to male and female age in the Seychelles warbler (Acrocephalus sechellensis). This long-lived passerine has a mean life expectancy of 5.5 years after fledgling (Komdeur, 1991) and a maximum observed lifespan of 19 years (Hammers & Brouwer, 2017). Seychelles warblers display facultative cooperative breeding (Komdeur, 1992; Richardson, Burke, & Komdeur, 2007): dominant pairs occupy a territory on their own or (in ~30% of territories) are joined by subordinates of either sex (Hammers et al., 2019). Clutches typically consist of one egg, but ~13% of nests contain one or two additional eggs, often laid by subordinate females (Richardson, Burke, & Komdeur, 2002; Richardson, Jury, Blaakmeer, Komdeur, & Burke, 2001). Individuals are socially monogamous, but ~44% of young are sired by males other than the social male (Hadfield, Richardson, & Burke, 2006; Richardson et al., 2001). Over 97% of all paternity is gained by dominant males (Hadfield et al., 2006; Raj Pant, Komdeur, Burke, Dugdale, & Richardson, 2019; Richardson et al., 2001) either in their own territory (within-group paternity: WGP) or with females from another territory (extra-group paternity: EGP). Thus, in this species EGP is virtually the equivalent of EPP.

In the Seychelles warbler, females that live in larger groups are more likely to produce extra-group offspring (EGO, i.e., offspring sired by extra-group males; Raj Pant et al., 2019). In subordinate mothers, but not in dominant mothers (which account for over 85% of offspring), the likelihood of producing EGO is positively linked to the genetic relatedness to the dominant male in the group (Raj Pant et al., 2019). However, this potential inbreeding avoidance mechanism in subordinate females does not prevent the occurrence of inbreeding in the population (Richardson, Komdeur, & Burke, 2004). In Seychelles warblers, there is evidence that dominant males actively seek EGP during extra-territorial forays (Komdeur, Kraaijeveld-Smit, Kraaijeveld, & Edelaar, 1999). Past research has also shown that females initiate successful copulations, are able to resist mating attempts (Komdeur et al., 1999) and are more likely to gain extra-group fertilizations from more MHC (major histocompatibility complex)-diverse males when paired with males of low MHC diversity (Richardson, Komdeur, Burke, & von Schantz, 2005), which suggests that female choice plays an active role in EGP. However, the relative role of female choice (pre- or post-copulatory) and male–male competition (including sperm competition) in determining patterns of extra-group fertilizations remains unknown in this system.

We use an 18-year longitudinal data set from the Seychelles warbler to determine the patterns of extra-group reproduction in relation to age in males (dominant) and females (dominant or subordinate). Our isolated study population on Cousin Island provides an excellent system for such investigations: inter-island migration is virtually absent (Komdeur, Burke, Dugdale, & Richardson, 2016; Komdeur, Piersma, Kraaijeveld, Kraaijeveld-Smit, & Richardson, 2004), extrinsic mortality is low (Hammers et al., 2015) and >96% of individuals have been DNA-sampled and individually colour-ringed since 1997 (Brouwer et al., 2010). Accurate parentage assignment and precise estimates of survival and individual reproductive output (throughout life) are therefore available. We assess how patterns of extra-group reproduction are affected by within-individual changes with age and between-individual selective appearance and disappearance effects. We also test for declines in extra-group reproduction in late life (senescence). Finally, we quantify the relative contribution of EGP and WGP success to annual reproductive success in males. By undertaking the analyses outlined above, we provide evidence to distinguish between different hypotheses as to why females engage in extra-pair mating and improve our understanding of the factors driving the evolution of infidelity.

2 | MATERIALS AND METHODS

2.1 | Study system

The Seychelles warbler is an insectivorous passerine endemic to the Seychelles archipelago. The population on Cousin Island (29 ha, 04°20’S, 55°40’E) has been monitored as part of a long-term study, which started in 1981 and intensified in 1997 (Hammers et al., Komdeur, 1992; Richardson, Komdeur, & Burke, 2003). Since then, virtually all breeding attempts have been followed each year during the main breeding season (June–September). As many birds as possible were captured every year, either using mist-nets or as nestlings. Newly caught individuals were assigned a unique combination of three colour rings and a British Trust for Ornithology metal ring.

All caught individuals were blood sampled (~25 µl) using brachial venipuncture and DNA from these samples was used for molecular sexing (following Griffiths, Double, Orr, & Dawson, 1998) and genotyping based on 30 microsatellite loci (see: Richardson et al., 2001; Spurgin et al., 2014). Parentage was assigned to 1,554 offspring (1991–2015) using MasterBayes 2.52 and used to build a
genetic pedigree (see: Edwards, Dugdale, Richardson, Komdeur, & Burke, 2018; Sparks et al., 2020).

Inter-island dispersal is virtually absent (<0.1%) in the Seychelles warbler (Komdeur et al., 2004, 2016) and individual resighting probability per season on Cousin Island is very high (~92–98%, Brouwer et al., 2010); therefore individuals not seen over two consecutive seasons can safely be assumed to be dead (Hammers, Richardson, Burke, & Komdeur, 2013).

During each breeding season, group membership, social status and territory boundaries were assigned for all birds using observations of foraging and singing locations, non-aggressive social interactions, and aggressive territorial interactions (e.g., Bebbington et al., 2017). Within groups, dominant pairs were identified via pair and courtship behaviours. Subordinate birds, which are often offspring that have delayed dispersal (Kingma, Bebbington, Hammers, Richardson, & Komdeur, 2016), are classified as “helpers” or “non-helpers” based on their participation in incubation (females only) and in feeding offspring (both males and females; Hammers et al., 2019; Komdeur, 1994).

Reproduction is seasonally constrained by invertebrate prey availability and offspring are fed for up to three months after fledging (Komdeur, 1991). We refer to the dominant male in a group as the “social male” of any reproductively mature females in his group (dominant and subordinate), as males can fertilize both dominant and subordinate females in their territory.

2.2 Data selection

We gathered previously generated parentage data for 934 Seychelles warblers that were assigned parentage with high confidence (>80%) and that hatched on Cousin Island during main breeding seasons in the period 1997–2014 (Edwards et al., 2018; Hadfield et al., 2006; Richardson et al., 2001; Sparks et al. 2020). We used these data to assess the age-dependent production of EGO by females and the age-dependent risk of cuckoldry for their social male partner (the dominant male in the group). We first tested if the likelihood that an offspring was sired by a male outside the breeding group (“EGP likelihood”) was related to the age of the mother (dominant or subordinate) and/or the age of the dominant male. The ages of dominant males and females are only weakly correlated in the Seychelles warbler (r = 0.16; Hammers et al., 2019). Given that Seychelles warbler females do not lay eggs in nests outside their own territory (Richardson et al., 2002), EGP likelihood will capture female infidelity.

We compiled 1,364 annual records of all dominant males alive between 1997 and 2014 that were genetically assigned at least one offspring across the whole data period (including entries of males siring no young in single years, n = 237 males). For each male, we determined the annual number of EGO (i.e., EGP success) and within-group offspring (i.e., WGP success). We then estimated the annual proportion of EGO sired by each dominant male (535 annual records from 233 males, excluding cases in which a male had an annual reproductive success of zero). In our paternity measures, we included only offspring that survived for at least three months to remove any potential bias on annual reproductive estimates resulting from differing catching efforts across years (which cause offspring to be caught at different ages in different years). Using these data, we assessed the relationship between male age and WGP, EGP, and the annual proportion of EGO sired by each male (i.e., the contribution of EGP to annual reproductive success).

2.3 Statistical analyses

We quantified within-individual effects of age on the production of EGO (i.e., longitudinal changes throughout an individual’s lifetime). To separate out between-individual (population-level cross-sectional) effects of age (i.e., due to selective appearance and disappearance), we employed the method developed by van de Pol and Verhulst (2006). By including age of first reproduction and age of last reproduction (or longevity) in addition to age in the same mixed model, this approach allows us to quantify the within-individual effect of age (expressed by the age term) while controlling for selective appearance and disappearance (expressed by age of first and last reproduction/longevity, respectively). Here, we modelled selective appearance using the age of first dominance for males, to account for when they could potentially start breeding (virtually all paternity is obtained by dominant males in the Seychelles warbler; Raj Pant et al., 2019). Because females can reproduce before gaining dominance, we used the age at which females were first assigned an offspring as subordinates or the age of first dominance—whichever came first (subsequently termed “age of first dominance” for simplicity). Age at death (longevity) was used to model selective disappearance for both males and females. Individuals of unknown longevity (i.e., birds translocated to other islands or those that had not died by the end of 2018) were excluded from the analyses. The chronologically age of individuals was always included as a fixed effect alongside age of first dominance and longevity so that it represents the within-individual effect of age on EGP. Chronological age, age of first dominance and longevity were all measured in years (integers).

Reproductive performance can change shortly before death, independently of age (Bowers et al., 2012; Coulson & Fairweather, 2001). Therefore, to avoid confounding any age-related effects with an age-independent terminal effect, we included a binary variable in models indicating whether a bird died before the subsequent breeding season.

We performed statistical analyses in R (3.6.3) with generalized linear mixed models (GLMMs) fitted using the lme4 (1.1–20) package (Bates, Mächler, Bolker, & Walker, 2015). We built separate GLMMs to analyse the following variables (summarized in Table 1): (1) offspring EGP likelihood—i.e., the likelihood that the offspring was sired by a male outside the group (yes/no binary variable), in relation to the age of the mother (n = 852 offspring) or the age of the dominant male in the group (n = 848 offspring); (2) annual paternity obtained by each male (n = 1,364 male/year) split into (2a)
TABLE 1  Summary of the response variables addressed in our analyses of age-related changes in the reproduction of the Seychelles warbler, highlighting the age-effects we detected

| Variable name                        | Description                                                                 | Age-effects assessed for | Detected age-effects                                           |
|--------------------------------------|-----------------------------------------------------------------------------|--------------------------|---------------------------------------------------------------|
| Extra-group paternity (EGP) likelihood| The likelihood that the offspring is sired by an extra-group male (yes/no binary variable) | Females (the mothers)    | Within-individual: increase in early life, senescent decline |
|                                      |                                                                            | The dominant male in the natal group | Within-individual: decrease in early life (before levelling off) |
| EGP success                          | The annual number of extra-group offspring (EGO) sired                       | Males                    | Within-individual: increase in early life, senescent decline |
| WGP success                          | The annual number of within-group offspring (WGO) sired                       | Males                    | Within-individual: increase in early life, senescent decline |
| Annual reproductive success (ARS)     | The annual number of offspring sired (EGO + WGO)                             | Males                    | Within-individual: increase in early life, senescent decline |
| Proportion of EGO                    | Annual EGP success over ARS                                                 | Males                    | Within-individual: increase in early life (before levelling off) |

EGP success (number of EGO sired), (2b) WGP success (number of within-group offspring sired) and (2c) total reproductive success (i.e., extra-group + within-group young sired); and (3) annual proportion of EGO (i.e., the number of EGO over the number of total offspring per male per year) sired by each dominant male that was assigned at least one offspring in a given year (n = 535). Models analysing EGP likelihood and the annual proportion of EGO sired by each male were constructed with a binomial error structure and logit link function, while models of paternity success (EGP/ WGP/annual reproductive success) were built with a Poisson error structure and log link function. Furthermore, we built a GLMM to perform a population-level comparison between the age of the social father (i.e., the cuckolded male) and the age of the genetic father (i.e., the extra-group sire) of each EGO (n = 395). The GLMM was built with a Poisson error structure and log link function. We checked for collinearity between fixed effects using the variance inflation factor (VIF) and found none (VIF ≤ 3). We standardized (mean-centred and scaled to one standard deviation) continuous predictors and used the “BOBYQA” nonlinear optimization (Powell, 2009) to aid convergence of models. The statistical significance of model terms was assessed using parametric bootstrap p-values (nsim = 3,000) from full models containing all biologically meaningful predictors of interest.

We assessed female and social male age effects on offspring EGP likelihood in separate models to avoid any potential bias caused by the non-independence of female and social male age over time (within our data set, 51% of females that reproduced in multiple years and 53% of social males that raised offspring in multiple years had more than one social partner). Both female and social male models contained four random effects (random intercepts): year, territory, mother’s identity and social male’s identity (pair identity explained zero variance and was not included as an additional random effect). Female and social male models also included two fixed effects in common: the age difference between the mother and her social male and the offspring’s natal group size (offspring EGP likelihood is higher in larger groups; Raj Pant et al., 2019).

The model analysing female age effects on EGP likelihood included as additional fixed effects the mother’s age (linear and quadratic), age of first dominance, longevity and terminal effect. To check for any potential bias caused by inbreeding avoidance effects occurring in subordinate females (Raj Pant et al., 2019), we re-ran this model on offspring produced only by dominant females (n = 759) and compared results with those from the model run on the full data set (i.e., offspring produced by dominant and subordinate females, n = 852).

The model analysing social male age effects on EGP likelihood contained the corresponding additional fixed effects of social male instead of female age traits (i.e., a social male’s age, age of first dominance, longevity and terminal effect). Furthermore, we built a model to test for any differences between the age of cuckolded males and the age of the extra-group sires that cuckolded them (population-level analysis). This model featured male age as the response variable, male status (i.e., extra-pair or cuckolded male) as a fixed effect and three random effects (random intercepts: social father, genetic father and mother identity).

Models analysing annual paternity success (EGP/WGP/reproductive success) per male and the annual proportion of EGO sired by each male included five fixed predictors—male age (linear and quadratic), age of first dominance, longevity and a terminal effect—and three random effects (random intercepts)—year, territory and male identity. Because annual EGP and WGP may affect one another, when analysing EGP and WGP, we also included as fixed effects either WGP or EGP respectively, and the interaction between WGP/EGP and male age.

A negative quadratic relationship between reproductive components and age does not necessarily indicate that a late-life decline in these components exists but may just represent that an increase early in life levels off at later ages (Bouwhuis, Sheldon, Verhulst, & Charmantier, 2009). To determine whether EGP likelihood and paternity success exhibit true late-life declines consistent with senescence, we tested for linear age effects after the peak age for each of these components. We estimated peak ages from the linear and
quadratic coefficients of age, as \((-β_{\text{linear}})/(2 \times β_{\text{quadratic}})\), from models we built with non-standardized data. We compiled subsets of individuals with ages ≥ the peak age for offspring EGP likelihood (female age effects: \(n = 319\); social male age effects: \(n = 346\)), paternity success (within-group: \(n = 598\); extra-group and total: \(n = 360\)), or the proportion of EGO sired (\(n = 97\)) per male. We ran models regressing EGP likelihood, paternity success (extra-group/within-group/total) or the proportion of EGO sired over the linear age (post-peak) of individuals and other predictors included in previous models, except the quadratic age term. For simplicity, when analysing EGP/WGP in the post-peak subsets we also excluded WGP/EGP and the interaction between WGP/EGP and male age, which were all nonsignificant predictors in full data set analyses (see Table 4).

3 | RESULTS

3.1 | Offspring EGP likelihood and female age

The proportion of offspring sired by an extra-group male was 42% in the population. There was a negative quadratic effect of maternal age on offspring EGP likelihood, which increased from a predicted ~29% for mothers in their first year to ~46% when the mother was 5.6 years old, after which it decreased to ~10% for the oldest mothers (Figure 1 and Table 2). Furthermore, the older a female was relative to the dominant male in her group, the higher the likelihood was that she produced an EGO (Table 2). Regarding senescent effects, the EGP likelihood of offspring produced by females ≥6 years old (i.e., past the peak age of EGP likelihood) declined with female age (\(β ± SE = −0.41 ± 0.20, p = .039\); Table S1). EGP likelihood was not affected by the mother’s age of first dominance, longevity or a terminal effect (Table 2). EGP likelihood was positively related to group size (Table 2). All results remained quantitatively similar when repeating the analysis of EGP likelihood using only offspring produced by dominant mothers (\(n = 759\); Table S2). This indicates that any in-breeding avoidance effect occurring via extra-group reproduction by subordinate females does not bias our results on age-dependent production of EGO by females.

3.2 | Offspring EGP likelihood and social male age

When analysing offspring EGP likelihood—i.e., the probability that a male was cuckolded (on an offspring by offspring case)—in relation to male age, we found a positive quadratic effect of male age. The likelihood of being cuckolded decreased within individuals, from a predicted ~44% in young males to ~34% in males of 6.2 years of age (Figure 2 and Table 3). Despite the positive quadratic effect of age there was no post peak senescent effect—i.e., that males were increasingly more likely to be cuckolded when they were ≥6 years old (\(β ± SE = 0.05 ± 0.19, p = .808\); Table S3). Males that lost WGP were on average 1 year younger than the extra-group
males that cuckolded them; this difference was significant (GLMM: \( \beta = 0.23 \pm 0.03, p < .001 \); Figure S1). The positive relationship between the probability of WGP loss and the female–social male age difference did not reach statistical significance (\( p = .067 \)). The probability of being cuckolded was not associated with male age of first dominance or a terminal effect, and only showed a non-significant tendency to decrease with male longevity (\( p = .063 \)). The probability of WGP loss was positively associated with group size (Table 3).

### 3.3 | Annual paternity success and male age

When analysing annual paternity success (extra-group, within-group and total) in relation to male age, we found an early-life increase and a late-life decline within individuals (Table 4). Specifically, there was a negative quadratic effect of male age on EGP success; the predicted number of extra-group offspring sired per annum increased from \(-0.06\) in males in their first year to peak at \(-0.24\) at 7.8 years and decreased thereafter to \(-0.03\) in the oldest males (Figure 3 and Table 4). There was also a negative quadratic effect of male age on annual WGP gained; the predicted number of within-group offspring sired increased from \(-0.23\) per annum in males in their first year to \(-0.32\) at 6.1 years and declined to \(-0.09\) in the oldest males (Figure 3 and Table 4).

As a result of the age-related changes in EGP and WGP outlined, total predicted annual reproductive success increased with male age from \(-0.29\) offspring in first-year males up to \(-0.58\) at 7.7 years, before declining to \(-0.14\) in the oldest males (Figure 3 and Table 4). The post-peak reduction in male reproduction in late life was confirmed by the significant negative linear relationships between age and annual EGP in males \(\geq 6\) years (\( \beta = -0.32 \pm 0.16, p = .013 \)) and total paternity success in males \(\geq 8\) years (\( \beta = -0.27 \pm 0.10, p = .001 \)) and total paternity success in males \(\geq 8\) years (\( \beta = -0.25 \pm 0.11, p = .021 \); Table S4). Male annual extra-group, within-group and total reproductive success were not affected by male longevity, age of first dominance or a terminal effect (Table 4). We found no evidence of a trade-off between EGP and WGP: WGP and its interaction with male age were not related to EGP success, and EGP and its interaction with male age did not predict WGP success (Table 4). When analysing the proportion of a male’s annual reproductive output obtained outside his own group, this increased with age, from a predicted \(-19\%\) in first-year males to a peak of \(-49\%\) at 8.6 years (Figure 4 and Table 5).

Despite finding a significant negative quadratic effect of age, there was no significant senescent decline in the proportion of reproductive success resulting from EGP with age in males \(\geq 9\) years old (\( \beta = -0.23 \pm 0.25, p = .343 \); Table S5). This suggests that the proportion of EGO sired remained relatively stable after peaking, probably as the decline in the amount of extra- and within-group offspring sired were similar in late life. The annual proportion of EGO sired was not influenced by male longevity, age of first dominance or a terminal effect (Table 5).

### 4 | DISCUSSION

Our results show that in both male and female Seychelles warblers, extra-group reproduction changed in relation to age within individuals, increasing in early life and declining in late life. In males, there were similar within-individual changes with age in WGP gained.
and—as a result of EGP and WGP patterns—in total reproductive success. Moreover, the likelihood of being cuckolded decreased within males from early to midlife. Extra-group reproduction accounted for ~50% of annual reproduction for males at their reproductive peak. No age-dependent differences among individuals, due to selective appearance or disappearance, were detected in relation to extra-group reproduction in either males or females, or in relation to within-group and total reproductive success in males. We detected
FIGURE 3  Extra-group paternity (EGP, top panel), within-group paternity (WGP, middle panel) and total paternity (bottom panel) gained by dominant male Seychelles warblers per year in relation to their age (n = 1,364 annual observations from 237 males). Means of raw data (points) and their standard error (bars) are shown with associated sample sizes. Black lines represent the predictions from the GLMMs (Table 4) and the areas shaded in grey indicate the 95% confidence intervals (estimated with the *predict* function in R package ggplot2, version 3.3.0).
4.1 | Age-dependent female extra-group reproduction

The likelihood of producing EGO changed with age within females, increasing until females were 5.6 years old and declining thereafter (Figure 1), but there were no selective appearance or disappearance effects (between-female age effects). Our findings are consistent with some cross-sectional studies that have found a positive association between female age and infidelity (Bouwman & Komdeur, 2005; Dietrich et al., 2004; Kempenaers et al., 1999), while other cross-sectional studies have shown a negative relationship (Moreno et al., 2015; Ramos et al., 2014; Stutchbury et al., 1997) or no relationship with age (Cordero, Wetton, & Parkin, 1999; Li & Brown, 2000; Lubjuhn et al., 2007; Veiga & Boto, 2000; Wagner et al., 1996). To our knowledge, no other studies have separated within- from between-individual age effects on the production of EGO by females.

The age-related increase in female extra-group reproduction we observed may be due to increases in experience and/or body condition of females with age (until they approach 6 years). In female Seychelles warblers, breeding and helping experience, which accrue with age, enhance the number of offspring raised to independence (Komdeur, 1996). Moreover, female reproductive success increases until they reach 6.5 years of age (Hammers, Richardson, Burke, & Komdeur, 2012), suggesting that a female's physical condition (and experience) improves until this point. It is possible that females at this peak of reproduction are more attractive to males seeking EGP (which may perceive them as successful reproducers) and that they are targeted for extra-group fertilizations. In that case, the detected within-female change in reproduction may be mostly, or even entirely, male-driven. Another possibility is that, as they grow older, females improve their ability to avoid mate-guarding and engage in extra-group copulations, thanks to experience or improved body condition (Bouwman & Komdeur, 2005). Additionally, greater experience and/or condition may allow older females to cope better with any reduction in paternal investment that may occur when males perceive a loss of paternity, thus allowing females to gain more extra-group fertilizations (the constrained female hypothesis; Gowaty, 1996). However, indirect evidence suggests that female extra-group reproduction is not constrained by male retaliation in the Seychelles warblers. In territories with cooperative breeding, helpers provide load-lightening to the dominant pair (van Boheemen et al., 2019; Hammers et al., 2019) and this might liberate dominant females from the costs imposed by male retaliation (Mulder, Dunn,
Contrary to the expectation based on this logic, the presence of helpers is not associated with higher dominant female extra-group reproduction in the Seychelles warbler (Raj Pant et al., 2019). When analysing female age effects, we also found that the older a female was compared to the dominant male in her group, the higher the likelihood was that she would produce an EGO. This is in accordance with other studies in which the production of extra-pair offspring was based on the combination of the female’s age and that of her social male (Bouwman & Komdeur, 2005; Dietrich et al., 2004; Ramos et al., 2014; Rätti, Lundberg, Tegelström, & Alatalo, 2001)—but see Lubjuhn et al. (2007) and Moreno et al. (2015). This result further suggests that females may be targeted more by extra-group males—and/or more easily avoid mate-guarding—when socially

**TABLE 5** Parameter estimates from a GLMM of the proportional contribution of extra-group paternity (EGP) to the annual reproductive success of dominant male Seychelles warblers (siring ≥ 1 offspring), in relation to male age (n = 535)

| Fixed effects        | Proportion of EGP (n = 535) |
|----------------------|-----------------------------|
|                      | β   | SE  | p    |
| Intercept            | −0.37 | 0.13 | .005 |
| Male age             | 0.38 | 0.14 | .005 |
| Male age²            | −0.18 | 0.07 | .012 |
| Male AFD             | −0.02 | 0.10 | .835 |
| Male longevity       | −0.01 | 0.13 | .941 |
| Male terminal effect | 0.50 | 0.30 | .117 |

| Random effects       | σ²  | n   |
|----------------------|-----|-----|
| Male ID              | 0.00 | 233 |
| Territory            | 0.51 | 131 |
| Year                 | 0.00 | 18  |

Note: Coefficient estimates (β), standard errors (SE) and p-values (p) are shown for each fixed effect. Variance (σ²) and number of observations (n) are shown for each random effect. Significant (p < .05) terms are shown in bold.

Abbreviation: AFD, age of first dominance.

Cockburn, Lazenby-Cohen, & Howell, 1994).
paired with a male that is considerably younger than the female (i.e., a young male that is not skilled at mate-guarding and/or defending his territory from intruders).

4.2 | Female benefits of infidelity

One key hypothesis suggests that females may seek extra-group fertilizations to obtain good paternal genes for their offspring (Hamilton & Zuk, 1982) and age is expected to reflect individual quality via viability (Trivers, 1972). Consequently, the fact that many (cross-sectional) studies have shown that older males gain more paternity through extra-pair reproduction than younger males (Ackay & Roughgarden, 2007; Hsu et al., 2015) has often been put forward as support for the good genes hypothesis (Forstmeier et al., 2014). In the Seychelles warbler, we found that male paternity gain (and loss) varied with age within individuals, and that age-related changes were not explained by selective appearance or disappearance effects. Similar results were found in the two other studies that have separated within- from between-individual age effects on EPP success and within-pair paternity loss (Hsu et al., 2017; Schroeder et al., 2016). This lack of any between-individual male age effect on EPP success is important because it undermines key evidence put forward to support the good genes model, as preferred or more successful sires do not appear to be of higher quality, at least as evidenced through greater longevity.

At first glance, this finding may appear to be in contrast to previous studies on the Seychelles warbler, which have provided evidence that extra-pair mating can offer genetic benefits. Those studies showed that dominant female Seychelles warblers paired with males exhibiting low MHC diversity gain extra-pair fertilizations (with males of higher MHC diversity) to produce more MHC-diverse offspring, with improved juvenile survival (Brouwer et al., 2010; Richardson et al., 2005). However, any female (pre-/post-copulatory) preference for more MHC-diverse extra-pair males would not cause a between-individual effect of male age on EPP in the Seychelles warbler, because the survival benefit of higher MHC diversity is only observed in juveniles. In adult males, there is no differential survival linked to MHC diversity (older adult males are not more MHC-diverse than younger adult males).

Further work is now required to understand the mechanisms through which males improve their ability to gain EGP with age, and whether this also provides any benefits to females. Females may also engage in extra-pair mating to gain other types of benefits (Forstmeier et al., 2014), such as fertilization assurance (Sheldon, 1994) in case they are socially paired with truly infertile males (Hasson & Stone, 2009). In the Seychelles warbler, individual males both gain EGP and lose WGP. This indicates that males that become cuckolded are not infertile but does not rule out that extra-group copulations could have evolved to guard against any rare cases of infertility (although totally infertile males have never been identified in the Seychelles warbler). Another reason why females may seek extra-pair fertilizations is to acquire indirect genetic non-additive benefits (e.g., compatible genes in offspring; Brown, 1997; Zeh & Zeh, 1996). However, unlike “good genes” benefits, other benefits are not normally expected to be signaled by male viability.

Alternatively, it is possible that infidelity may not provide any benefits for females and instead may have evolved as a by-product of positive selection on genetically correlated traits in males (between-sex correlation) or in females themselves (Arnvist & Kirkpatrick, 2005; Forstmeier, Martin, Bolund, Schielzeth, & Kempeaers, 2011; Forstmeier et al., 2014; Halliday & Arnold, 1987). This idea, which has received little attention so far, may constitute a promising avenue in unveiling the evolution of infidelity in socially monogamous species, but assessing this hypothesis is beyond the scope of the current study.

4.3 | Male age-dependent paternity gain and loss

Both extra-group and within-group paternity success increased within individual male Seychelles warblers in early life and declined in late life. Moreover, the likelihood of being cuckolded decreased within males at young ages and remained stable from midlife onward. These within-male changes in reproduction and cuckoldry, coupled with the lack of between-male differences due to selective appearance and disappearance, do not provide evidence for the good genes hypothesis (Hamilton & Zuk, 1982) but support the competitive ability hypothesis (Nakagawa et al., 2015). This hypothesis argues that the improvement in male paternity success with age is due to increasing experience (Hsu et al., 2015; Morton et al., 1990; Westneat & Stewart, 2003) or body condition (Nakagawa et al., 2015) causing, for instance, improvements in ejaculate competitiveness, timing of copulations, mate-guarding ability, and effectiveness in finding and copulating with fertile extra-pair females. Even though EGP success changed within individuals with age and was unrelated to longevity (which is expected to reflect an individual’s intrinsic quality through viability), the correlative nature of our study calls for further work to distinguish between effects arising due to age per se and effects caused by the difference in genetic or phenotypic quality among individuals. A starting point would be to determine which variables improve with age in the Seychelles warbler.

Gaining EPP enables males to increase their reproductive output without suffering costs associated with rearing additional offspring. In the Seychelles warbler, this ability showed marked changes with age in males. EGP success increased in males up to 7.8 years, before displaying a decline consistent with senescence in the oldest males. The relative contribution of EGP to total paternity increased until 8.6 years of age, with no evidence for a post-peak decline (i.e., both EGP and WGP declined similarly in late life, so the proportion of paternity explained by EGP remained constant). Consequently, EGP is a very important source of total paternity gain, contributing ~49% of annual reproductive success in older males. These results concur with those from numerous cross-sectional studies that have shown a...
positive correlation between EPP success and male age (reviewed by Cleasby & Nakagawa, 2012) and with a longitudinal study on house sparrows which identified age-dependent increases in EPP and within-pair paternity success (Hsu et al., 2017). However, our results are particularly insightful because they clearly show that these age-related changes in extra- and within-group paternity occur within individuals and not as a result of preferred males living longer.

There was no evidence of a trade-off between WGP and EGP gain in Seychelles warblers and both WGP and EGP success increased in early life and declined in late life (Figure 3); this indicates that when males obtain more WGP success, they do not do so at the expense of EGP gains, and vice versa. The combined result of EGP and WGP is that annual reproductive success changes with male age, increasing until 7.7 years and declining thereafter (Figure 3). Such within-individual variation in reproductive success (an increase in early life followed by a decline in late life) is common in vertebrates (Nussey et al., 2013). In the Seychelles warbler, annual EGP success displayed a particularly steep increase at young ages, thus strongly intensifying the spike in annual male reproductive success at 7.7 years of age (Figure 3).

Age-dependent changes in the contribution of EPP to male reproductive success have been shown in several species. Most of these (largely cross-sectional) studies have found an increase in the contribution of EPP to reproductive success with male age (e.g., Brekke, Ewen, Clucas, & Santure, 2015; Girndt et al., 2018; Richardson & Burke, 1999) (but see e.g., McDonald, Spurgin, Fairfield, Richardson, & Pizzari, 2017). To our knowledge, the only study that has disentangled within- from between-individual age effects found that the contribution of EPP to reproductive success increased within males until midlife (in agreement with our findings), but that it also varied between individuals with age (Hsu et al., 2017). Our results are important because they show that EGP may play a very substantial role in male fitness, especially later in life. More longitudinal studies across a diversity of taxa are needed to fully understand how important EPP is in terms of overall reproductive success and, in particular, how that may change in relation to age.

It is widely recognized that senescence is an important age-related process occurring in the wild. Numerous studies have assessed senescence in multiple traits (Hayward et al., 2015), including survival (e.g., Cameron & Siniff, 2004) and reproductive output (e.g., Dugdale, Pope, Newman, MacDonald, & Burke, 2011), across a number of species. In the Seychelles warbler, senescence in female reproductive success has been detected in the past (Hammers et al., 2012). To our knowledge, however, no studies have assessed senescence in extra-pair reproduction in females, and only one study has addressed (and found) senescence in extra-pair reproduction in males (Hsu et al., 2017), although this study did not explicitly test for senescent post-peak declines in EPP. Here, we analysed senescent post-peak declines in extra-group reproduction and found these to occur in both male and female Seychelles warblers. In males, we also assessed and found evidence for senescence in WGP and total paternity success. Our results highlight the importance of the role that senescence plays in the alternative pathways to reproductive success in this and possibly other species.

5 | CONCLUSIONS

The lack of between-male age effects on extra-group reproduction emerging from our study undermines the often cited suggestion that male age-dependent patterns of EPP success support the good genes hypothesis for the evolution of female infidelity. Our results provide support for the idea that infidelity may be important to females for other reasons, such as the acquisition of compatible genes in offspring, or that infidelity evolved because of genetic constraints (i.e., genetic correlation between infidelity and traits under positive selection). Our analyses also provide, to our knowledge, the first explicit evidence that senescence in extra-group reproduction occurs not only in males, but also in females. Finally, our work shows that EGP explains a large proportion of the annual reproductive success of males, and that age-specific changes in EGP amplify age-dependent patterns of reproduction. Further work is now needed to understand how this affects male variance of reproductive success and therefore selection for infidelity.

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AUTHOR CONTRIBUTIONS

D.S.R., H.L.D., S.R.P. and M.H. conceived and designed the study. D.S.R., M.H. and S.R.P. contributed significantly to fieldwork, and H.L.D performed parentage assignment. S.R.P. undertook the analysis (with substantial input from M.H.) and wrote the manuscript. S.R.P., M.H., D.S.R. and H.L.D. contributed to the interpretation of results and conclusions. All authors discussed the results and contributed to the manuscript. D.S.R., J.K., T.B. and H.L.D. manage the
long-term warbler project and data set. D.S.R. and J.K. raised the PhD studentship funding for S.R.P.

DATA AVAILABILITY STATEMENT
Analyses reported in this article can be reproduced using the data provided by Raj Pant, Komdeur, Burke, Dugdale, and Richardson (2020). The data used in this study are available in DRYAD (https://doi.org/10.5061/dryad.3jkd51fs).

REFERENCES
Ackay, E., & Roughgarden, J. (2007). Extra-pair reproductive activity in birds: Review of the genetic benefits. Evolutionary Ecology Research, 9(5), 855–868.

Albrecht, T., Schnitzer, J., Kreisinger, J., Exnerova, A., Bryja, J., & Munclinger, P. (2007). Extrapair mating and the opportunity for sexual selection in long-distance migratory passerines. Behavioral Ecology, 18(2), 477–486. https://doi.org/10.1093/beheco/arm001

Arnvist, G., & Kirkpatrick, M. (2005). The evolution of infidelity in socially monogamous passerines: The strength of direct and indirect selection on extrapair copulation behavior in females. American Naturalist, 165, S26–S37. https://doi.org/10.1086/429350

Bates, D., Mächler, M., Bolker, B., & Walker, S. (2015). Fitting linear mixed-effects models using lme4. Journal of Statistical Software, 67(1), 1–48. https://doi.org/10.18637/jss.v067.i01

Bebington, K., Kingma, S. A., Fairfield, E. A., Dugdale, H. L., Komdeur, J., Spurgin, L. G., ..., Strassmann, J. E. (2017). Kinship and familiarity mitigate costs of social conflict between Seychelles warbler neighbors. Proceedings of the National Academy of Sciences of the United States of America, 114(43), E9036–E9045. https://doi.org/10.1073/pnas.1704350114

Bouwman, K., Komdeur, J., & Dugdale, H. (2010). Old female reed buntings (Emberiza schoeniclus) increase extrapair paternity in their broods when mated to young males Karen. Behaviour, 147(11), 1449–1463. https://doi.org/10.1163/156853907X1253391891

Brown, J. L. (1997). A theory of mate choice based on heterozygosity. Behavioral Ecology, 8(1), 60–65. https://doi.org/10.1093/beheco/8.1.60

Cameron, M. F., & Siniff, D. B. (2004). Age-specific survival, abundance, and immigration rates of a Weddell seal (Leptonychotes weddellii) population in McMurdo Sound. Antarctica. Canadian Journal of Zoology, 82(4), 601–615. https://doi.org/10.1139/z04-025

Cleasby, I. R., & Nakagawa, S. (2012). The influence of male age on within-pair and extra-pair paternity in passerines. Ibis, 154(2), 318–324. https://doi.org/10.1111/1471-919X.2011.01209.x

Cordero, P. J., Metton, J. H., & Parkin, D. T. (1999). Extra-pair paternity and male badge size in the house sparrow. Journal of Avian Biology, 30(1), 97. https://doi.org/10.2307/3677248

Cousson, J. C., & Fairweather, J. A. (2001). Reduced reproductive performance prior to death in the Black-legged Kittiwake: Senescence or terminal illness? Journal of Avian Biology, 32(2), 146–152. https://doi.org/10.1034/j.1600-048X.2001.320207.x

Curio, E. (1983). Why do young birds reproduce less well? Ibis, 125, 400–404. https://doi.org/10.1111/j.1471-919X.1983.tb03130.x

Dean, R., Cornwallis, C. K., Levlie, H., Worley, K., Richardson, D. S., & Pizzari, T. (2010). Male reproductive senescence causes potential for sexual conflict over mating. Current Biology, 20(13), 1192–1196. https://doi.org/10.1016/j.cub.2010.04.059

Dietrich, V., Schmolz, T., Winkel, W., Epplen, J., & Lubjuhn, T. (2004). Pair identity – An important factor concerning variation in extra-pair paternity in the coal tit (Perus ater). Behaviour, 141(7), 817–835. https://doi.org/10.1163/156853904265644

Dixon, A., Ross, D., O’Malley, S. L. C., & Burke, T. (1994). Paternal investment inversely related to degree of extra-pair paternity in the reed bunting. Nature, 371(6499), 698–700. https://doi.org/10.1038/371698a0

Dugdale, H. L., Pope, L. C., Newman, C., MacDonald, D. W., & Burke, T. (2011). Age-specific breeding success in a wild mammalian population: Selection, constraint, restraint and senescence. Molecular Ecology, 20(15), 3261–3274. https://doi.org/10.1111/j.1365-294X.2011.05167.x

Edme, A., Munclinger, P., & Krist, M. (2016). Female collared flycatchers choose neighbouring and older extra-pair partners from the pool of males around their nests. Journal of Avian Biology, 47(4), 552–562. https://doi.org/10.1111/jav.00839

Edwards, H. A., Dugdale, H. L., Richardson, D. S., Komdeur, J., & Burke, T. (2018). Extra-pair parentage and personality in a cooperatively breeding bird. Behavioral Ecology and Sociobiology, 72(3), 37. https://doi.org/10.1007/s00265-018-2448-z

Forstmeier, W., Martín, K., Bolund, E., Schielzeth, H., & Kempenaers, B. (2011). Female extrapair mating behavior can evolve via indirect selection on males. Proceedings of the National Academy of Sciences of the United States of America, 108(26), 10608–10613. https://doi.org/10.1073/pnas.101395108

Forstmeier, W., Nakagawa, S., Griffith, S. C., & Kempenaers, B. (2014). Female extra-pair mating: Adaptation or genetic constraint? Trends in Ecology and Evolution, 29(8), 456–464. https://doi.org/10.1016/j.tree.2014.05.005

Girndt, A., Chng, C. W. T., Burke, T., & Schroeder, J. (2018). Male age is associated with extra-pair paternity, but not with extra-pair mating behaviour. Scientific Reports, 8(1), 8378. https://doi.org/10.1038/s41598-018-26649-1

Gowaty, P. A. (1996). Parental care: evolution, mechanisms, and adaptable significance. In P. Slater, & M. Milinski (Eds.), Advances in the study of behavior (1st ed., Vol. 25, pp. 477–531). San Diego, CA: Academic Press.

Griffith, S. C., Owens, I. P. F., & Thuman, K. A. (2002). Extra-pair paternity in birds: A review of interspecific. Molecular Ecology, 11, 2195–2212.

Griffiths, R., Double, M. C., Orr, K., & Dawson, R. J. (1998). A DNA test to sex most birds. Molecular Ecology, 7(8), 1071–1075. https://doi.org/10.1046/j.1365-294X.1998.00389.x
Halliday, T., & Arnold, S. J. (1987). Multiple mating by females: A perspective from quantitative genetics. *Animal Behaviour*, 35(3), 939–941.

Hamilton, W. D., & Zuk, M. (1982). Heritable true fitness and bright birds: A role for parasites? *Science*, 218, 384–387. https://doi.org/10.1126/science.7123238

Hammers, M., & Brouwer, L. (2017). Rescue behaviour in a social bird: Removal of sticky ‘bird-catcher tree’ seeds by group members. *Behaviour*, 154(4), 403–411. https://doi.org/10.1163/15685 39X-00003428

Hammers, M., Kingma, S. A., Bebbington, K., van de Crommenacker, J., Spurgin, L. G., Richardson, D. S., & Komdeur, J. (2015). Senescence in the wild: Insights from a long-term study on Seychelles warblers. *Experimental Gerontology*, 71, 69–79. https://doi.org/10.1016/j. exger.2015.08.019

Hammers, M., Kingma, S. A., Spurgin, L. G., Bebbington, K., Dugdale, H. L., Burke, T., & Komdeur, J. (2019). Breeder helpers differ in age and their genetics when helping: A study of parentage assignment in the Seychelles warbler. *Behavioral Ecology*, 30(5), 1075–1081. https://doi.org/10.1093/beheco/arz081

Hammers, M., Kingma, S. A., Bebbington, K., Dugdale, H. L., Burke, T., & Komdeur, J. (2012). The impact of reproductive and extra-pair fertilizations on extra-pair offspring: A meta-analytic examination. *Royal Society B: Biological Sciences*, 279(1728), 384–387. https://doi.org/10.1098/rspb.2011.2204

Hasson, O., & Stone, L. (2009). Male infertility, female fertility and extra-pair copulations. *Biological Reviews*, 84(2), 225–244. https://doi.org/10.1111/j.1469-185X.2008.00068.x

Hayward, A. D., Moorad, J., Regan, C. E., Berenos, C., Pilkington, J. G., Pemberton, J. M., & Nussey, D. H. (2015). Asynchrony of senescence among phenotypic traits in a wild mammal population. *Experimental Gerontology*, 71, 56–68. https://doi.org/10.1016/j. exger.2015.08.003

Hsu, Y. H., Schroeder, J., Winney, I., Burke, T., & Nakagawa, S. (2015). Are extra-pair males different from cuckolded males? A case study and a meta-analytic examination. *Molecular Ecology*, 24(7), 1558–1571. https://doi.org/10.1111/mec.13124

Hsu, Y. H., Simons, M. J. P., Schroeder, J., Girndt, A., Winney, I. S., Burke, T., & Nakagawa, S. (2017). Age-dependent trajectories differ between within-pair and extra-pair paternity success. *Journal of Evolutionary Biology*, 30(5), 951–959. https://doi.org/10.1111/jeb.13058

Jennions, M. G., & Petrie, M. (2000). Why do females mate multiply? A review of the genetic benefits. *Biological Reviews of the Cambridge Philosophical Society*, 75(1), 21–64. https://doi.org/10.1017/323199005423

Kempenaers, B., Congdon, B., Boag, P., & Robertson, R. J. (1999). Extra-pair paternity and egg hatchability in tree swallows: Evidence for the genetic compatibility hypothesis? *Behavioral Ecology*, 10(3), 304–311. https://doi.org/10.1093/beheco/10.3.304

Kempenaers, B., Verheyen, G. R., & Dhondt, A. (1997). Extra-pair paternity in the blue tit (*Parus caeruleus*): Female choice, male characteristics, and offspring quality. *Behavioral Ecology*, 8(5), 481–492. https://doi.org/10.1093/beheco/8.5.481

Kingma, S. A., Bebbington, K., Hammers, M., Richardson, D. S., & Komdeur, J. (2016). Delayed dispersal and the costs and benefits of different routes to independent breeding in a cooperatively breeding bird. *Evolution*, 70(11), 2595–2610. https://doi.org/10.1111/evo.13071

Kokko, H. (1998). Good genes, old age and life-history trade-offs. *Evolutionary Ecology*, 12(6), 739–750. https://doi.org/10.1023/A:1006541701002

Komdeur, J. (1991). Cooperative breeding in the Seychelles warbler. Cambridge, UK: University of Cambridge.

Komdeur, J. (1992). Importance of habitat saturation and territory quality for evolution of cooperative breeding in the Seychelles warbler. *Nature*, 358, 493–495. https://doi.org/10.1038/355242a0

Komdeur, J. (1994). Experimental evidence for helping and hindering by previous offspring in the cooperative-breeding Seychelles warbler *Acrocephalus sechellensis*. *Behavioral Ecology and Sociobiology*, 34, 175–186. https://doi.org/10.1007/4600929

Komdeur, J. (1996). Influence of helping and breeding experience on reproductive performance in the Seychelles warbler: A translocation experiment. *Behavioral Ecology*, 7(3), 326–333. https://doi.org/10.1093/beheco/7.3.326

Komdeur, J., Burke, T., Dugdale, H. L., & Richardson, D. S. (2016). Seychelles warblers: The complexities of the helping paradox. In W. D. Koenig, & J. L. Dickson (Eds.), *Cooperative breeding in vertebrates: Studies of ecology, evolution, and behavior* (pp. 197–217). Cambridge, UK: Cambridge University Press.

Komdeur, J., Kraaijeveld-Smit, F., Kraaijeveld, K., & Edelaar, P. (1999). Explicit experimental evidence for the role of mate guarding in minimizing loss of paternity in the Seychelles warbler. *Proceedings of the Royal Society B: Biological Sciences*, 266(1433), 2075–2081.

Komdeur, J., Piersma, T., Kraaijeveld, K., Kraaijeveld-Smit, F., & Richardson, D. S. (2004). Why Seychelles Warblers fail to re-colonize nearby islands: Unwilling or unable to fly there? *Ibis*, 146(2), 298–302. https://doi.org/10.1111/j.1474-919X.2004.00255.x

Lebigre, C., Arcese, P., Sardell, R. J., Keller, L. F., & Reid, J. M. (2012). Extra-pair paternity and the variance in male fitness in song sparrows (*Melospiza melodia*). *Evolution*, 66(10), 3111–3129.

Li, S.-H., & Brown, J. L. (2000). High frequency of extrapair fertilization in a plural breeding bird, the Mexican jay, revealed by DNA microsatellites. *Animal Behaviour*, 60(6), 867–877. https://doi.org/10.1006/anbe.2000.1554

Lubjuhn, T., Gerken, T., Brün, J., & Schmall, T. (2007). Yearling male great tits, *Parus major*, suffer more strongly from cuckoldry than older males. *Zoology*, 110(5), 387–397. https://doi.org/10.1016/j.zool.2007.07.005

McDonald, G. C., Spurgin, L. G., Fairfield, E. A., Richardson, D. S., & Pizzari, T. (2017). Pre- and postcopulatory sexual selection favor aggressive, young males in polyandrous groups of red junglefowl. *Evolution*, 71(6), 1653–1669. https://doi.org/10.1111/evo.13242

Medawar, P. B. (1952). An unsolved problem of biology. *Evolution in health and disease*. London, UK: Inaugural Lecture Delivered at University College London.

Moreno, J., Martínez, J. G., González-Braojos, S., Cantarero, A., Ruiz-de-Castañeda, R., Precioso, M., & López-Arrabé, J. (2015). Extra-pair paternity declines with female age and wing length in the pied flycatcher. *Ethology*, 121(5), 501–512. https://doi.org/10.1111/eth.12364

Morton, E. S., & Derrickson, K. C. (1990). The biological significance of age-specific return schedules in breeding purple martins. *Condor*, 92(4), 1040–1050. https://doi.org/10.2307/1368740

Morton, E. S., Forman, L., & Braun, M. (1990). Extrapair fertilizations and the evolution of colonial breeding in purple martins. *The Auk*, 107(2), 275–283. https://doi.org/10.2307/4087610

Mulder, R. A., Dunn, P. O., Cockburn, A., Lazenby-Cohen, K. A., & Howell, M. J. (1994). Helpers liberate female fairy-wrens from constraints on extra-pair mate choice. *Proceedings of the Royal Society B: Biological Sciences*, 255(1344), 223–229. https://doi.org/10.1098/rspb.1994.0032

Nakagawa, S., Schroeder, J., & Burke, T. (2015). Sugar-free extrapair mating: A comment on Arct et al *Behavioral Ecology*, 26(4), 971–972. https://doi.org/10.1093/beheco/arv041
Nussey, D. H., Froy, H., Lemaitre, J. F., Gaillard, J. M., & Austad, S. N. (2013). Senescence in natural populations of animals: Widespread evidence and its implications for bio-gerontology. Ageing Research Reviews, 12(1), 214–225. https://doi.org/10.1016/j.arr.2012.07.004

Powell, M. J. D. (2009). The \textsc{bobyqa} algorithm for bound constrained optimization without derivatives. NA Report NA2009/06. Cambridge, UK: University of Cambridge.

Raj Pant, S., Komdeur, J., Burke, T., Dugdale, H. L., & Richardson, D. S. (2019). Socio-ecological conditions and female infidelity in the Seychelles warbler. Behavioral Ecology, 30(5), 1254–1264. https://doi.org/10.1093/beheco/azr072

Raj Pant, S., Komdeur, J., Burke, T., Dugdale, H. L., & Richardson, D. S. (2020). Data for MEC-20-0592: Age-dependent changes in infidelity in Seychelles warblers. Dryad Digital Repository, https://doi.org/10.5061/dryad.3j9kd51fs

Ramos, A. G., Nunziata, S. O., Lance, S. L., Rodriguez, C., Faircloth, B. C., Richardson, D. S., Burke, T., & Komdeur, J. (2002). Direct benefits and the evolution of female-biased cooperative breeding in Seychelles warblers. Evolution, 56(11), 2313–2321. https://doi.org/10.1111/j.0014-3820.2002.tb00154.x

Richardson, D. S., Burke, T., & Komdeur, J. (2007). Grandparent helpers: The adaptive significance of older, post-dominant helpers in the Seychelles warbler. Evolution, 61(12), 2790–2800. https://doi.org/10.1111/j.1558-5646.2007.00222.x

Richardson, D. S., Burke, T., & Komdeur, J. (2003). Avian behaviour: Altruism and infidelity among warblers. Nature, 422(6932), 580. https://doi.org/10.1038/422580a

Richardson, D. S., Komdeur, J., & Burke, T. (2004). Inbreeding in the Seychelles warbler: Environment-dependent maternal effects. Evolution, 58(9), 2037–2048. https://doi.org/10.1111/j.0014-3820.2004.tb00488.x

Richardson, D. S., Komdeur, J., Burke, T., & von Schantz, T. (2005). MHC-based patterns of social and extra-pair mate choice in the Seychelles warbler. Proceedings of the Royal Society B: Biological Sciences, 272(1564), 759–767. https://doi.org/10.1098/rspb.2004.3028

Schroeder, J., Hsu, Y.-H., Winney, I., Simons, M., Nakagawa, S., & Burke, T. (2016). Predictably philandering females prompt poor paternal provisioning. American Naturalist, 188(2), 219–230. https://doi.org/10.1086/687243

Sherdon, B. C. (1994). Male phenotype, fertility, and the pursuit of extra-pair copulations by female birds. Proceedings of the Royal Society B: Biological Sciences, 257(1348), 25–30. https://doi.org/10.1098/rspb.1994.0089

Sparks, A. M., Spurgo, L. G., van der Velde, M., Fairfiled, E. A., Komdeur, J., Burke, T., ... Dugdale, H. L. (2020). Telomere heritability and parental age at conception effects in a wild avian population. Ecol Evol, https://ecolevirox.org/eq2zf/

Spurgo, L. G., Wright, D. J., van der Velde, M., Collar, N., Komdeur, J., Burke, T., & Richardson, D. S. (2014). Museum DNA reveals the demographic history of the endangered Seychelles warbler. Evolutionary Applications, 7(9), 1134–1143. https://doi.org/10.1111/eva.12191

Stutchbury, B. J. M., Piper, W. H., Neudorf, D. L., Tarof, S. A., Rhymer, J. M., Fuller, G., & Fleischer, R. C. (1997). Correlates of extra-pair fertilization success in hooded warblers. Behavioral Ecology and Sociobiology, 40(2), 119–126. https://doi.org/10.1007/s002650050324

Taylor, M. L., Price, T. A. R., & Wedell, N. (2014). Polyandry in nature: A global analysis. Trends in Ecology and Evolution, 29(7), 376–383. https://doi.org/10.1016/j.tree.2014.04.005

Trivers, R. L. (1972). Parental investment and sexual selection. In B. Campbell (Ed.), Sexual selection and the descent of man, 1871–1971 (pp. 136–179). Chicago, IL: Aldine.

van Boeheemen, L. A., Hammers, M., Kingma, S. A., Richardson, D. S., Burke, T., Komdeur, J., & Dugdale, H. L. (2019). Compensatory and additive helper effects in the cooperatively breeding Seychelles warbler (Acrocephalus sechellensis). Ecology and Evolution, 9(5), 2986–2995. https://doi.org/10.1002/ece3.4982

van de Pol, M., & Verhulst, S. (2006). Age-dependent traits: A new statisti- cal model to separate within- and between-individual effects. American Naturalist, 167(5), 766–773. https://doi.org/10.1086/503331

Veiga, J. P., & Boto, L. (2000). Low frequency of extra-pair fertilisations in House Sparrows breeding at a high density. Journal of Avian Biology, 31(2), 237–244. https://doi.org/10.1034/j.1600-048X.2000.310215.x

Wagner, R. H., Schug, M. D., & Morton, E. S. (1996). Condition-dependent control of paternity by female purple martins: Implications for coloniality. Behavioral Ecology and Sociobiology, 38(6), 379–389. https://doi.org/10.1007/s002650050255

Westneat, D. F. (1990). Genetic parentage in the indigo bunting: A study using DNA fingerprinting. Behavioral Ecology and Sociobiology, 27(1), 67–76. https://doi.org/10.1007/BF00183315

Westneat, D. F., & Stewart, I. R. K. (2003). Extra-pair paternity in birds: Causes, correlates, and conflict. Annual Review of Ecology, Evolution, and Systematics, 34(1), 365–396. https://doi.org/10.1146/annurev.ecolsys.34.011802.132439

Williams, G. C. (1957). Pleiotropy, natural selection, and the evolution of life tables. Nature, 179(4628), 766–773. https://doi.org/10.1038/179766a0

Zeh, J., & Zeh, D. W. (1996). The evolution of polyandry I: Intragenomic conflict and genetic incompatibility. Proceedings of the Royal Society B: Biological Sciences, 263(1377), 1711–1717. https://doi.org/10.1098/rspb.1996.0250

SUPPORTING INFORMATION
Additional supporting information may be found online in the Supporting Information section.

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