1	Lifetime offspring production in relation to breeding lifespan, attractiveness, and mating
2	status in male collared flycatchers
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# 20 Abstract

21 As a comprehensive fitness parameter, lifetime reproductive success (LRS) is influenced by many different environmental and genetic factors, among which longevity is one of the most 22 important. These factors can be reflected in secondary sexual characters, which may 23 contribute to the life-history of individuals via social relations with conspecifics. Facultative 24 polygyny in birds is another conspicuous reproductive trait that potentially increase male 25 reproductive success, but lifetime success data in relation to polygyny are scarce. Here we 26 used 17 years of breeding data to quantify the LRS of male collared flycatchers (Ficedula 27 albicollis) on the basis of lifetime recruitment of offspring. Breeding lifespan showed a 28 29 positive relationship with LRS, and it was also significantly associated with mean recruitment of offspring per breeding year. Body size and sexually selected forehead patch size did not 30 predict the number of recruits. Polygyny was positively associated with LRS, but probably 31 32 only due to the correlation between lifespan and polygyny. Our results demonstrate that the relationship between longevity and LRS is not explained by the larger number of reproductive 33 attempts when living longer, and question the adaptive value of polygyny in this population. 34 The lack of association between forehead patch size and recruitment suggests that forehead 35 patch is a poor indicator of phenotypic quality in our birds. 36

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38 Key words: reproductive success, recruitment, longevity, polygyny, Ficedula albicollis

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### 41 Introduction

In species with overlapping iteroparity, the most accurate method to estimate the contribution of their genes to the subsequent generations (i.e. fitness; Clutton-Brock 1988) is the calculation of lifetime reproductive success (LRS), which is given by the number of lifetime

recruits, i.e. sexually mature offspring contributing to the breeding population (Brommer et al. 45 2004). There are two main determinants of the number of lifetime recruits: lifespan and the 46 number of recruits per breeding attempt. Several studies have identified longevity or the 47 number of breeding attempts as an important determinant of LRS (birds: Gustafsson 1986; 48 Merilä and Sheldon 2000; Blums and Clark 2004; mammals: Clutton-Brock 1988; Bérubé et 49 al. 1999). However, a prolonged lifespan in itself is not sufficient to be successful, as a 50 considerable proportion of individuals do not produce any recruits despite their long 51 reproductive life (Gustafsson 1989; Blums and Clark 2004), and the successful individuals 52 also vary greatly in productivity (Newton 1989). In fast-living species, which live for a short 53 time but may produce numerous offspring per breeding attempt, the reproductive output in a 54 single year is more important for LRS than in slow-living species (Saether and Bakke 2000). 55

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57 Annual reproductive success can be affected by individual characteristics such as body size (Grant and Grant 2000) as well as sexually selected traits (Gustafsson et al. 1995; Hasselquist 58 et al. 1996). Individuals with more elaborate sexual traits are often of better quality (Møller 59 1994; Hasselquist et al. 1996). Hence, a positive relationship is also expected between the 60 elaboration of these characters and the number of recruits (Møller 1994; Petrie 1994, but see 61 Brooks 2000). Similarly, the number of mates also has an important role. Polygyny is usually 62 considered beneficial to immediate male reproductive success, but its effect on LRS is poorly 63 understood (Gustafsson 1989; Hasselquist 1998). In addition, care is needed when 64 interpreting the relationship between polygyny and fitness, as polygynous males may have a 65 high LRS because of their high quality and viability, irrespective of their mating status 66 (Hannon and Dobush 1997, also see Lambrechts and Dhondt 1986). 67

It is generally quite difficult to measure LRS in natural population, as long-term studies are 69 70 required to follow a sufficient number of individuals throughout their lives. In this study, using a 17-year dataset, we investigated potential determinants of male LRS in a small 71 passerine bird, the collared flycatcher (Ficedula albicollis). We measured LRS as the number 72 of lifetime recruits, and examined how individual variation in LRS was explained by 73 differences in breeding lifespan. We were also interested in the lifetime success consequences 74 of body size, forehead patch size (a sexually selected character; Hegyi et al. 2002), and 75 polygyny, which is a regularly encountered reproductive status of males (Garamszegi et al. 76 2004). 77

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### 80 Materials and methods

### 81 Study species and field methods

The collared flycatcher is a small, long-distance migratory, hole-nesting, insectivorous 82 passerine that breeds in deciduous woodlands of Central Europe. Our data were collected 83 between 1987 and 2003 in the Pilis Mountains, Hungary, in an oak-dominated forest, where 84 more than 750 nestboxes were placed. The nuptial plumage of collared flycatcher males is 85 black and white with a prominent white collar, forehead patch and wing patches. This species 86 is ideal for long-term studies of reproductive success. It shows a preference for nestboxes, can 87 easily be captured, and has high breeding site fidelity (Pärt and Gustafsson 1989; Könczey et 88 al. 1992; Hegyi et al. 2002) and considerable local recruitment rates (Pärt 1990; Török et al. 89 2004). Nestboxes were checked multiple times a week throughout the nesting period, so 90 breeding attempts were followed from nest building to fledging. Most parents were captured 91 92 and ringed when feeding young, but some females were caught during incubation.

The forehead patch of males is an important sexually selected trait which, however, shows 94 complicated links to individual life history. Studies in a population on Gotland, Sweden, 95 showed among others that the size of this trait was related to age and body condition during 96 the previous breeding season (Gustafsson et al. 1995; Qvarnström 1999). In addition, a 97 positive relationship was found between a male's mean lifetime patch size and his mean 98 recruitment of offspring per breeding attempt (Gustafsson et al. 1995; for more information 99 about forehead patch see Gustafsson and Qvarnström 2006 for a review). In our population, in 100 contrast, forehead patch size did not reflect the body condition of males, its dependence on 101 age and yearly environmental conditions was weak and there was no relationship between 102 103 forehead patch size and breeding lifespan either (Hegyi et al. 2002, 2006a). However, the trait is still a sexually selected character as an important determinant of social mating success: 104 males with a larger forehead patch find a mate more rapidly relative to their arrival date 105 106 (Hegyi et al. 2010). The yearly means of forehead patch sizes strongly varied among years in our population showing a linear temporal decline (Hegyi et al. 2006a), so patches of the same 107 size could be relatively small in earlier years and relatively large in later years. The forehead 108 patch size of males was estimated as a product of maximum height and maximum width. 109 Forehead patch dimensions and tarsus length (to estimate body size) were measured with a 110 calliper to the nearest 0.1 mm. The within-season repeatability of measurements between the 111 major measurers was r = 0.76 for tarsus and r = 0.60 for forehead patch. (We calculated r -112 the intra-class correlation coefficient – from variance components as described in Lessells and 113 Boag (1987), n = 32.) We did not mention here the other main secondary sexual character of 114 male flycatchers, the white wing patch, because of the more limited dataset available for that 115 trait. 116

The collared flycatcher is predominantly monogamous, but a fraction of males successfully 118 119 attract two females and become polygynous. During the study period 83 out of 1558 breeding males were polygynous in our population. Several studies have found that males that had two 120 mates divided their parental investment between the two nests with most effort devoted to the 121 primary brood (Král et al. 1996), which may increase LRS compared to monogamy. In 122 contrast, both primary and secondary nests experience similarly reduced reproductive success 123 124 in our population (Garamszegi et al. 2004), so the positive effect of polygyny on LRS should be weaker. 125

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## 127 Statistical analyses

We used a 17-year dataset, which contained data from 683 male flycatchers after excluding 128 individuals that were the subject of experiments that could have influenced their breeding 129 130 success. However, missing data for different variables resulted in different sample sizes among tests. In our population, returning male collared flycatchers occupy a nestbox within a 131 mean of 129 metres from the box that they used in the preceding year (Könczey et al. 1992) 132 and movement between plots is very rare, so it is possible to follow individuals throughout 133 their entire breeding lifespan. Only males with complete recapture records (that is, those that 134 were recaptured in each year between their first and last captures) were included in the 135 analyses (95.9% of non-manipulated males). 136

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The LRS of males was characterized by the number of lifetime recruits. As a significant proportion of recruits return only at the age of two or three years, males that bred after the year 2000 were excluded from the analyses, as their recruits may have returned after 2003, the end of the study period. Birds that were first captured in 1987 or 1988 as an adult (i.e. at least 2 years old, as indicated by the absence of subadult plumage) were also omitted, because very

few males had been trapped before 1987, so it was not known if these birds had bred prior to 143 the study period. In the morphological database, each individual had at most one 144 measurement. Males with records from multiple years were represented by the measurement 145 from their earliest year in the dataset. If there were more than one measurement from an 146 individual in a given year, we randomly selected one of them. As the yearly means of 147 forehead patch size in the population varied strongly among years, showing a linear temporal 148 149 decline, and because body size also declined during the study period (Hegyi et al. 2006a), we used year-standardized forehead patch size and tarsus length in the analyses (mean of 0, 150 standard deviation of 1). A male was considered polygynous if it was caught in two nestboxes 151 152 while feeding nestlings. It was possible that we did not detect polygyny in some cases, so the observed rate of polygyny (4.9 %) is an underestimate (but it is similar to that found in the 153 Swedish population with a similar approach (4.3 %); Qvarnström et al. 2003). Given the high 154 155 capture effort, polygynous males caught at only one nest probably allocated nearly all of their care to this nest (included here as a monogamous nest) while neglecting the other nest (not 156 used here due to the lack of the male). A secondary brood without the male caring for the 157 offspring presumably produces little reproductive output, so the misclassification of these 158 birds as monogamous is likely to bias polygynous LRS upwards. In this study, males were 159 included in the analyses as polygynous if they were polygynous during at least one year of 160 their entire lifespan. We adopted this binary categorization because only three males were 161 polygynous in more than one year. Breeding date was not considered in our analyses because 162 it was not repeatable within males (results not shown), so the timing of individual breeding 163 attempts would not directionally bias the estimates of LRS. Indeed, models controlling for 164 mean breeding date vielded the same conclusions as those reported here. 165

The breeding lifespan of a bird was defined as the number of consecutive years (see above) in 167 168 which it was caught as a potential breeder (irrespective of the actual breeding success). Because of the high site fidelity of breeding males (Könczey et al. 1992; Hegyi et al. 2002) 169 and the high capture effort in our population, birds that bred in one of our study plots in a 170 given year but were not recaptured in subsequent years were considered dead. We tested if 171 including cohorts (year of birth) in the analyses changed the results. Year of birth was obvious 172 173 in birds that ringed as a nestling or as one-year-old (which wear subadult plumage). In newly ringed adult males, the youngest possible age assignment (2-year-old) was used because 174 males that had been ringed as a nestling and bred first as an adult were mostly 2 years old (our 175 176 unpublished data).

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We found a significant relationship between breeding lifespan and mating status (polygynous 178 179 males had a longer lifespan; also see the Results section), so using both as independent variables in the same model would have led to questionable results (Graham 2003). We 180 resolved the situation in two steps. First, we ran two models that contained only one of these 181 two variables. This informed us about the relationship of one variable with LRS without 182 correction for the other. Second, to see whether the effect of mating status is due to its 183 correlation with lifespan, we assessed the effect of mating status on LRS among males of the 184 same breeding lifespan. We used the most common lifespans of 1 and 2 only, as for the other 185 values there were very few polygynous data. 186

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Individuals with a long lifespan can produce more recruits than those with a short lifespan simply because they have more breeding attempts. In connection with this, they have time to gain experience, and have more chance to become polygynous etc. In this case, when comparing individuals with the same breeding lifespan, we would not expect a difference

among them in terms of reproductive success. However, long-lived individuals can also
produce more recruits independently of their lifespan, for example, due to their more viable
offspring. In this case, they may realize higher reproductive success even on a yearly basis.
To clarify this issue, we computed the mean recruitment of offspring per breeding year by
dividing the number of lifetime recruits by the number of breeding years.

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198 Data on LRS were analysed in two generalized linear models with Poisson error and log link, containing the number of lifetime recruits as the dependent variable, forehead patch size and 199 tarsus length as continuous predictors, and either breeding lifespan as a continuous variable, 200 201 or mating status as a factor. Polygynous and monogamous males with the same breeding lifespan (1 or 2, see above) were compared with respect to LRS by using the number of 202 lifetime recruits as the dependent variable, mating status as a factor and forehead patch size 203 204 and tarsus length as continuous predictors. We used binomial error and logit link when comparing individuals producing versus not producing a recruit during their breeding 205 lifespan. In this analysis binary recruit production was the dependent variable and breeding 206 lifespan was a continuous variable. The dispersion parameters of the models were less than 207 1.34 and we corrected for them in the analyses. In all models we employed a backward 208 stepwise model selection procedure. Statistics presented for non-significant terms reflect their 209 reintroduction to the final model one by one. Since the mean recruitment of offspring per 210 breeding year could not be transformed to conform to any standard distribution, it was 211 analysed using non-parametric statistics (Spearman's rank correlation, Mann-Whitney U-212 test). All statistical tests were calculated in Statistica 5.5. Means are represented with their 213 standard errors. We report effect sizes estimated as Pearson's correlation coefficients and the 214 associated 95% confidence intervals as suggested previously (Nakagawa and Cuthill 2007). 215

# 218 Results

Individual males produced up to five recruits during their breeding lifespan of 1 to 6 years, 219 but 67.8% of males did not recruit any offspring. Mating status was significantly related to 220 breeding lifespan (polygynous males had a longer lifespan; Wald  $\chi^2_{(1)} = 13.44$ , P < 0.001, 221 n = 467,  $n_{mono} = 444$ ,  $n_{poly} = 23$ ; effect size r = 0.170 (0.080 / 0.256), Fig. 1), so we did not 222 enter the two parameters into the same model (see Methods). Breeding lifespan had a positive 223 effect on the number of lifetime recruits (Table 1, Fig. 2a). The probability of producing a 224 recruit also increased with lifespan (binary data, Wald  $\chi^2_{(1)} = 54.28$ , P < 0.001, n = 683; effect 225 size r = 0.282 (0.211 / 0.350), Fig. 2b), though there were several long-lived birds that did not 226 227 produce any breeding offspring. Forehead patch size and tarsus length were not correlated with the number of recruits (Table 1). Polygynous males had two clutches in at least one 228 season of their life, and so we could expect them to have more nestlings that fledged and more 229 offspring that returned to the breeding population. Indeed, polygyny, when assessed in 230 isolation from lifespan, had a positive effect on the LRS of male collared flycatchers 231 (Table 1). However, when we compared polygynous and monogamous males with the same 232 breeding lifespan, the success of polygynous males was no different from that of 233 monogamous males (breeding lifespan of 1: Wald  $\chi^2_{(1)} = 0.48$ , P = 0.49, n<sub>mono</sub> = 291, n<sub>poly</sub> = 9; 234 effect size r = 0.040 (-0.074 / 0.153); breeding lifespan of 2: Wald  $\chi^2_{(1)} = 1.97$ , P = 0.16, 235  $n_{mono} = 119$ ,  $n_{poly} = 8$ ; effect size r = 0.125 (-0.051 / 0.292)). The above results suggest that 236 237 polygyny is positively related to LRS, but this relationship is explained by the correlation between lifespan and polygyny. Including cohorts (year of birth) in the model did not affect 238 the outcome of the analysis. 239

The mean recruitment of offspring per breeding year did not differ between polygynous and strictly monogamous males (Mann–Whitney U-test: adjusted Z = -1.181, P = 0.24,  $n_{mono} = 444$ ,  $n_{poly} = 23$ ; effect size r = -0.055 (-0.145 / 0.036)), but it was positively related to breeding lifespan (Spearman's rank correlation: r = 0.142, P = 0.002, n = 467; effect size r =0.142 (0.052 / 0.230)). This finding means that the lifespan effect on LRS is not simply due to the larger number of breeding attempts by longer-lived males. Forehead patch size and tarsus length did not have any effects in this model either.

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### 250 Discussion

Here we found that the LRS of male collared flycatchers was mainly associated with their 251 breeding lifespan and that this was in a positive direction. Moreover, longevity was also 252 253 positively related to the mean number of recruits per breeding year. The morphological traits we considered (forehead patch size and tarsus length) were not related to the number of 254 lifetime recruits. There was a positive relationship between mating status and LRS, but this 255 association could not be detected when comparing polygynous and monogamous males with 256 the same breeding lifespan. Polygyny did not increase the mean yearly reproductive success 257 of males either. These findings show a more complex picture on the relation of breeding 258 lifespan and LRS than generally expected, and also have interesting implications for the 259 evolution of visual signals and alternative reproductive tactics in our population. 260

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In many bird species extra-pair paternity plays an important role in influencing the reproductive success of males. Unfortunately, we could not assess this component, because we did not have blood samples from individuals for most years of the study period. Given that paternity in the own nest is apparently not related to male ornaments or body size in our

population (Rosivall et al. 2009), a directional effect of extra-pair paternity on our results is 266 unlikely in this respect. However, the relationship between paternity and polygyny could be 267 negative (Pilastro et al. 2002), very weak (Pearson et al. 2006) or positive (Soukup and 268 Thompson 1998), so, our data on polygyny must be treated with caution. Studies conducted in 269 different populations of the sibling species pied flycatcher (*Ficedula hypoleuca*) consistently 270 showed that polygynous males had extra-pair young in their broods more frequently than 271 monogamous males (Brün et al. 1996; Lubjuhn et al. 2000; Drevon and Slagsvold 2005). 272 These findings suggest that considering extra-pair paternity would further reduce the 273 advantage of polygynous over monogamous males, thereby strengthening our conclusions. 274

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In species that breed more than once, breeding lifespan is often one of the most important 276 correlates of LRS (Newton 1989) and this holds true in our case as well: breeding lifespan has 277 278 a strong positive effect on the number of lifetime recruits. Such a relationship is expected because the presence or absence of a reproductive attempt often makes a numerically greater 279 difference to LRS than lower or higher reproductive success in a given season. However, we 280 also found that breeding lifespan positively predicted not only the number of lifetime recruits 281 but also the mean recruitment of offspring per breeding year, which means that individuals 282 with a long lifespan attained a higher LRS than expected from the number of their breeding 283 bouts. The higher yearly reproductive performance of long-lived individuals may be 284 explained by accumulating experience, that is, improving ability to raise offspring with 285 ageing, which experience could not be reached by short-lived individuals. This may either be 286 due to a better knowledge of the resource distribution and quality (i.e. foraging ability) or a 287 better ability to occupy a cavity in a favourable area (whereby e.g. reducing the risk of 288 predation). Alternatively, only birds with given genetic or phenotypic properties can survive 289 to a certain age (Forslund and Pärt 1995). These individuals may also better cope with the 290

costs of reproduction and may attain higher success independently of their lifespan. This 291 292 explanation may be more consistent with our results than improving experience as a large number of individuals produce no returning young despite breeding several times during their 293 life (also see Gustafsson 1989; Blums and Clark 2004). Females of many species apparently 294 prefer older males (Enstrom 1993; Richardson and Burke 1999), or traits that indicate the 295 expected lifespan of males (Jennions et al. 2001), thereby often enhancing the quality of 296 young they produce (Saetre et al. 1995; Hegyi et al. 2006b). In addition, females may also 297 invest preferentially in such offspring (Burley 1986; de Lope and Møller 1993), which may 298 further increase their mate's reproductive success. This implies that, in some cases, individual 299 300 attributes may influence both breeding lifespan and, indirectly, other aspects of fitness.

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The forehead patch is a well-studied secondary sexual character of male collared flycatchers. 302 303 It is sexually selected, but it seems that its information content differs between populations. Our results show that male forehead patch size did not predict lifetime offspring recruitment 304 and this result is consistent with those of earlier studies performed in this population, 305 suggesting that the forehead patch is a poor indicator of phenotypic quality in our birds 306 (Hegyi et al. 2002, 2006a, but see Hegyi et al. 2010), in contrast to the Swedish population 307 (Gustafsson et al. 1995; also see Gustafsson and Qvarnström 2006). It is possible that the 308 advantage of large forehead patch can be detected only in extra-pair paternity, but within-309 brood paternity at least was not robustly related to forehead patch size in this population 310 (Rosivall et al. 2009, but see Michl et al. 2002). Alternatively, large-patched males may be 311 successful in some years, but variation in year quality may swamp the overall effect (Török et 312 al. 2004). Long-term data on within- and extra-brood paternity would be helpful to further 313 clarify the selection pressures on forehead patch size. 314

Many studies have shown that polygyny increases seasonal reproductive success of male birds 316 317 due to the increased number of offspring from multiple broods (Davies and Houston 1986; Soukup and Thomson 1998). However, the increase may not be very large in cases when the 318 reduced male help impairs the success of secondary or both females (Slagsvold and Lifjeld 319 1994; Garamszegi et al. 2004). Our results imply that polygynous males realized a higher 320 LRS only because of their longer lifespan. This points out that, if polygyny is connected with 321 lifespan, a positive relationship between the occurrence of polygyny and LRS may have 322 nothing to do with the causal effect of polygyny on reproductive success. To our knowledge, 323 the only study to date that has examined the effects of mating status on LRS while correcting 324 325 for lifespan was conducted in a Swedish population of collared flycatchers. That study found that polygyny increased lifetime success irrespective of lifespan (Gustafsson 1989). In our 326 collared flycatcher population, which lives in more variable environmental conditions (Török 327 328 et al. 2004), the situation is different (also see Garamszegi et al. 2004).

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Our findings raise the fundamental question of whether it is adaptive for males to build 330 polygynous partnerships. It is possible that polygyny is not adaptive at present and the net 331 selection pressure operating on polygyny is very low. Indeed, as in collared flycatcher 332 polygynous males spend most of their life monogamously (also see Gustafsson 1989) the 333 potential benefits to polygynous males in terms of yearly reproductive success are expected to 334 become smaller when viewed across the whole breeding lifespan. Alternatively, polygyny 335 may be advantageous only in years of good food supply, but males may still try to become 336 polygynous every year because they cannot predict the food supply at the beginning of the 337 season (Lubjuhn et al. 2000). This explanation may easily apply in our population, where the 338 unpredictable among-year fluctuations of food availability even prevent the individual 339 optimization of clutch size (Török et al. 2004). Finally, fitness benefits to polygynous males 340

may also appear in the attractiveness of their offspring (Gwinner and Schwabl 2005; Huk and 341 342 Winkel 2006), which will increase the number of grandoffspring, a variable we did not assess here. Even data from the Swedish population did not suggest a reproductive advantage for the 343 offspring of polygynous males (Gustafsson and Qvarnström 2006), which makes such an 344 advantage unlikely in our population. Further investigations are currently underway to clarify 345 the determinants of polygyny in our population and its consequences for LRS in more detail. 346 Note that the potential detection failure of polygyny may lead to overestimated polygynous 347 LRS (see Methods). However, this supports rather than weakens our results, i.e. this likely 348 overestimated LRS is not higher than that of monogamous males. 349

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To summarize, our results show that the reproductive advantage of longer-living individuals does not always simply originate from their more breeding opportunities, and suggest that these individuals may also have other superior characteristics. The lack of effect of forehead patch size and polygyny on LRS indicates that the reproductive consequences of traits and strategies used in male mate acquisition are far from straightforward. Finally, our findings with polygyny and LRS highlight the need to consider the interrelations of various factors when assessing their importance in influencing LRS.

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- Fig. 1 The probability of becoming polygynous in relation to breeding lifespan in male
  collared flycatchers. Sample sizes are shown
- 536 Fig. 2 a) The number of lifetime recruits (mean  $\pm$  SE) and b) the probability of producing a
- 537 recruit in relation to breeding lifespan. In Fig. 2a the values of breeding lifespan are shown
- only up to 4 years for better visibility as only three males lived longer than this. Sample sizes
- 539 are shown
- 540

Fig. 1

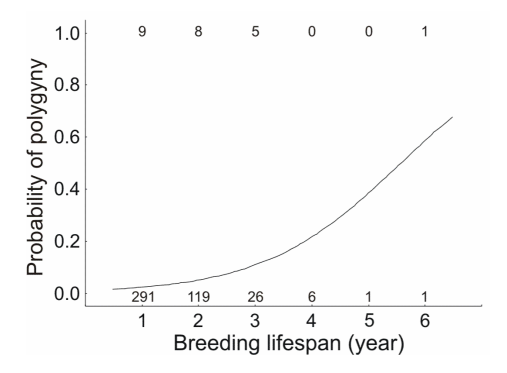
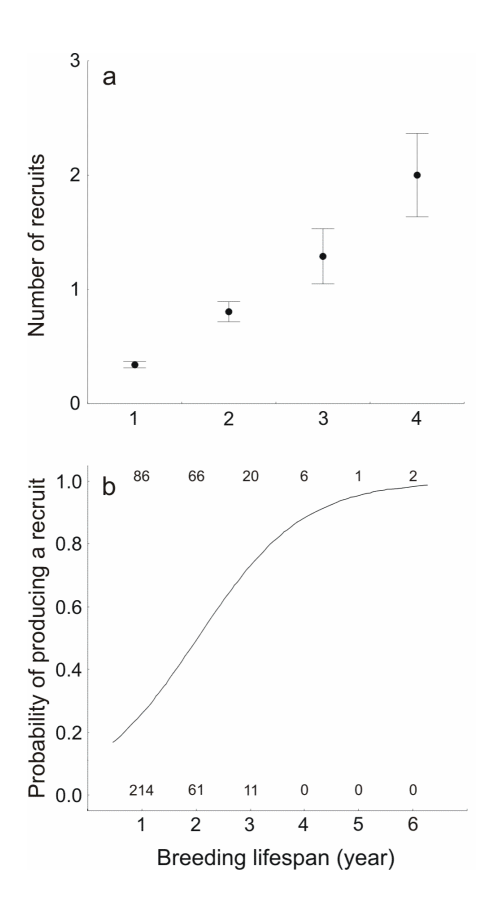


Fig. 2



	Breeding lifespan model					Mating status model					
	Wald $\chi^2$	Р	Effect size (r)	CI lower	CI upper	Wald $\chi^2$	Ρ	Effect size (r)	CI lower	CI upper	
Breeding lifespan	94.24	<0.001	0.449	0.374	0.519						
Mating status						6.70	0.01	0.120	0.029	0.208	
Forehead patch size	0.97	0.32	0.046	-0.045	0.136	1.24	0.27	0.052	-0.039	0.142	
Tarsus length	0.007	0.93	0.004	-0.087	0.095	0.07	0.79	0.012	-0.078	0.103	

Table 1 Correlates of lifetime reproductive success of male collared flycatchers. Generalized linear models with backward stepwise model selection. The number of degrees of freedom was 1 in all cases. n = 467 (monogamous 444, polygynous 23); CI, 95% confidence interval