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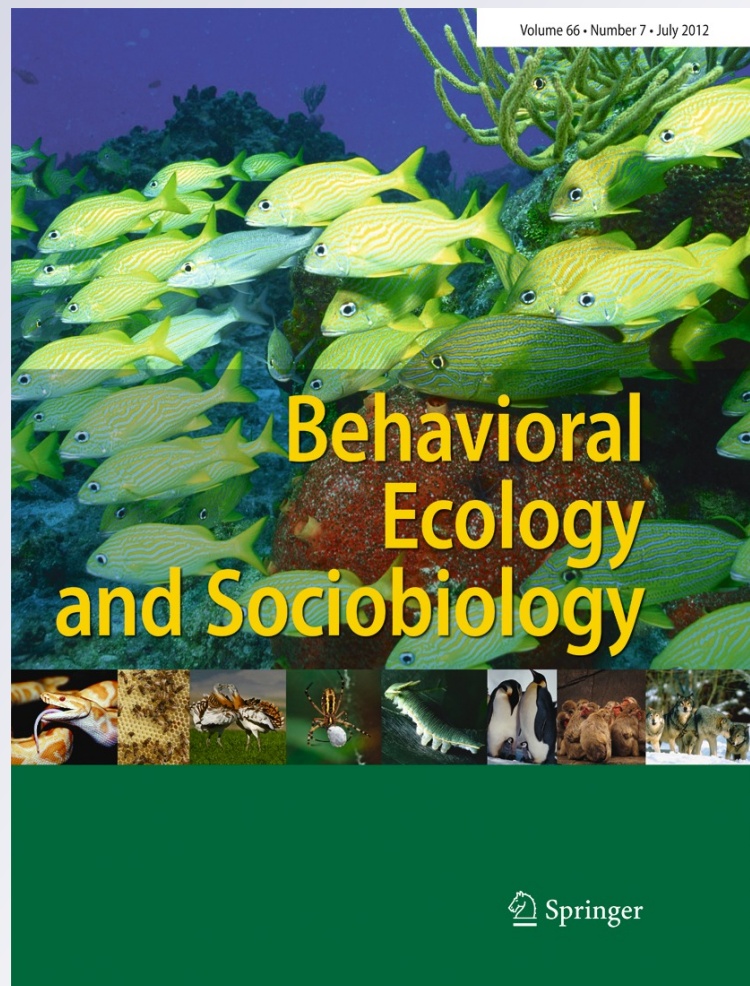
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Breeding success of a brood parasite is associated with social mating status of its host

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Abstract Reproductive success of brood parasites varies considerably both among and within host species, mainly due to differences in host egg-rejection rates and survival of parasitic chicks. Here, we investigated the breeding success of the cuckoo (*Cuculus canorus*) in one of its major hosts, the great reed warbler (*Acrocephalus arundinaceus*), with respect to host social mating status. In this passerine, polygynous males provide less parental care to their young per nest than monogamous males. Consequently, their less-assisted females may fledge lower numbers of nestlings than monogamous females. This may be especially true for secondary females, which often receive limited or no paternal help with young at all. Based on these findings, we expected higher cuckoo reproductive success in nests of socially monogamous than polygynous great reed warbler males. More specifically, we predicted lower fledging success of cuckoo young in nests of secondary than primary or monogamous females. In line with the prediction, we found higher cuckoo fledging success in nests of monogamous

than polygynous males, monogamous nests being more than twice as successful as secondary nests. We detected, however, only a tendency to lower cuckoo success in primary compared to monogamous nests and no differences between primary and secondary nests. Moreover, neither parasitism nor host egg-rejection rates differed among the nests of different status. Our results show, for the first time, that the social mating status of a host may influence the overall reproductive success of a brood parasite and thus should be considered in further studies.

Keywords *Acrocephalus arundinaceus* · Brood parasitism · Cuckoo · *Cuculus canorus* · Great reed warbler · Polygyny · Reproductive success

Introduction

Avian brood parasitism is an alternative reproductive strategy where parasitic females lay their eggs into the nests of other individuals (hosts), avoiding thus subsequent parental duties (Payne 1977; Rothstein 1990). In this way, brood parasites may dramatically reduce host fitness (Øien et al. 1998; Hauber 2003). Therefore, during their long-term interactions, both the brood parasites and their hosts have evolved intricate adaptations and counter-adaptations, which make the system an illustrative example of coevolution (Schulze-Hagen et al. 2009; Davies 2011).

However, not only do the brood parasites affect host fitness but also hosts may influence the reproductive output of brood parasites. Several previous studies have shown that the reproductive success of brood parasites varies considerably both among and within host species (Brooke and Davies 1987; Soler et al. 1995; Scott and Lemon 1996; Rutila et al. 2002; Mermoz and Reboreda 2003; Grim et

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al. 2011). This variation can be ascribed mainly to the differences in host egg-rejection rates (Davies 2000; Stokke et al. 2008) and in predation pressure on host nests (Dearborn 1999; Kleven et al. 2004; Hannon et al. 2009; de Mársico and Reboresda 2010). Furthermore, because parasites are dependent on hosts, their reproductive success may also relate to host parental quality. Chicks of many brood parasites are larger and require longer periods of host parental care compared to host own young (Wyllie 1981; Kilpatrick 2002; Numerov 2003). Raising the alien nestling may thus be more exhausting for the foster parents, especially for small passerines, than raising their own offspring (but see Brooke and Davies 1989). It is then natural to expect that survival of parasitic chicks will vary with the amount and quality of parental care provided by the hosts.

Generally, the intensity of parental care varies considerably among species (Trivers 1972; Clutton-Brock 1991). A lot of this variation can be explained by the differences in species-specific mating systems (Thomas and Székely 2005; Olson et al. 2008). In many polygynous birds with biparental nestling care, females sharing the same male usually receive less male assistance with parental duties than monogamous females (Webster 1991; Pinxten and Eens 1994; Trnka and Prokop 2010), resulting in their lower direct reproductive success (Johnson et al. 1993; Lubjuhn et al. 2000; Pribil 2000). Based on this scenario, survival of parasitic chicks in host nests may also vary with host social mating status.

A well-known example of a cuckoo (*Cuculus canorus*) host is the great reed warbler (*Acrocephalus arundinaceus*) (Moskát and Honza 2002; Kleven et al. 2004). It is a facultatively polygynous, open-nesting passerine with biparental nestling care (Cramp 1992; Hasselquist 1998). The rate of polygyny in this species varies between 8 and 43 % (Hasselquist 1998; Leisler and Wink 2000; Trnka et al. 2010; Honza et al. 2011), and cuckoo parasitism rate ranges between 0 and 68 % (e.g. Moksnes et al. 1993; Moskát et al. 2002; Moskát and Honza 2002). Previous studies have demonstrated that polygynous great reed warbler males provide significantly less parental care (feeding as well as nest defence) to their offspring than monogamous males (Dyrce 1986; Bensch and Hasselquist 1994; Sejberg et al. 2000; Trnka and Prokop 2010).

While a recent study of Trnka and Prokop (2011) compared frequencies of hatched cuckoos among great reed warbler nests of different status, here we made an important step forward. In the present study, we investigated whether there is an association between great reed warbler social mating status and cuckoo reproductive success. We used within-species variability in presumed host quality in contrast to between-species variability that was examined in previous studies (Kleven et al. 2004; Grim 2006; Sklepowicz and Haľupka 2009; Remeš 2010).

Some studies have shown that cuckoos may parasitize nests non-randomly within a host population according to nest position, host parental quality, egg appearance or nesting activity (Moskát and Honza 2000; Avilés et al. 2006; Cherry et al. 2007; Polačiková et al. 2009; but see Avilés et al. 2009; Antonov et al. 2012). As these characteristics can differ between monogamous and polygynous hosts, we separately explored cuckoo parasitism rate, host egg-rejection rate and survival of cuckoo chicks in great reed warbler nests of different status. We expected higher cuckoo reproductive success in nests of monogamous than polygynous males. Since secondary great reed warbler females often fledge a lower number of young than monogamous females (Catchpole et al. 1985; Dyrce 1986; Bensch and Hasselquist 1991), we specifically predicted lower fledging success of cuckoo young in secondary than in monogamous or primary nests. According to our best knowledge, the influence of host social mating status on the reproductive output of a brood parasite has not yet been studied.

Methods

Study sites and populations

The study was carried out on two colour-ringed great reed warbler populations: at fishponds near Štúrovo (47°51' N 18°36' E) in Slovakia (site 1) and in a fishpond area between Hodonín (48°51' N 17°07' E) and Mutěnice (48°54' N 17°02' E), Czech Republic (site 2). The two sites are located about 155 km apart. For their detailed descriptions, see Trnka and Prokop (2010) and Honza et al. (2002), respectively. The studied populations consist of 40–60 (site 1) and 80–100 breeding pairs (site 2). On both sites, great reed warblers breed in relatively narrow belts of littoral vegetation surrounding the fishponds. The overall rate of male polygyny (i.e. percent of polygynous males) was 26 % (21–29 %) and did not differ between years within sites (site 1: $\chi^2=0.04$, $df=1$, $P=0.849$; site 2: $\chi^2=1.22$, $df=1$, $P=0.270$) or between the sites ($\chi^2=0.28$, $df=1$, $P=0.595$).

Data collection

We conducted the fieldwork during the breeding seasons of 2009 and 2010. From early May to late July, we systematically searched for great reed warbler nests and checked them daily from the beginning of egg laying until clutch completion. During the nest controls, we numbered each egg with a waterproof pen according to the laying sequence. These regular checks enabled us to ascertain the incidence of cuckoo parasitism: if the nest contained a cuckoo egg, we considered it as parasitized. We regarded cuckoo eggs as accepted if hosts did not reject them (by ejection or nest

desertion) within 6 days after parasitism (see also Øien et al. 1998; Moskát and Honza 2002). After clutch completion, we checked each active nest two to three times during egg incubation until hatching and then other two to three times until fledging or nest failure.

In total, we found 335 nests (site 1, 102; site 2, 233). We determined the social mating status of each female and her mate based on their captures at the nests during the nest-building or egg-laying stages and confirmed it repeatedly throughout a season by direct observations of colour-ringed birds defending their nests or feeding their young. Male and female mating status may change in time due to settlement of other females or because of nest predation/desertion (Bensch 1996; Sejberg et al. 2000; Trnka and Prokop 2011). Therefore, we used the current mating status of nest owners either during egg laying and early incubation (to assess its effect on parasitism and host egg-rejection rates) or when caring for young (to examine its consequences for cuckoo fledging success). At both of these stages, we classified each nest as belonging to a monogamous, primary or secondary female. A monogamous female was the only female of a monogamous male. A primary female was the first female of a polygynous male, which started her breeding at the time when the secondary female has already been present in the same territory. Analogically, the secondary female was the second female of the polygynous male, breeding after, but overlapping in time with, the primary female (see also Bensch 1996). Out of 335 nests included into analyses, 231 were initially monogamous, 41 primary and 63 secondary. In 33 nests, the initial status changed later in the course of breeding season (15 from monogamous to primary, 5 from primary to monogamous, 2 from primary to secondary, 9 from secondary to monogamous and 2 from secondary to primary).

Previous studies have shown that females of polygynous males in some species may be of lower quality than monogamous females or, more specifically, that secondary females are of lower quality than primary females (Griggio et al. 2003). On the other hand, the opposite may also be true (Forstmeier et al. 2001a, b) or there may be no differences in female quality at all (Honza et al. 2011; Trnka and Prokop 2011; Trnka et al. 2012). To check for the potential effect of this confounding variable, we compared body condition (expressed as residuals from the linear regression of body mass on wing length) of the monogamous, primary and secondary females ($n=114$) caring for 8- to 10-day-old nestlings. We found no differences in female condition in relation to her mating status (after controlling for female identity, generalised linear mixed-effects model (GLMM): $F=1.04$; $df=2, 17$; $P=0.375$).

Statistical analysis

We fitted GLMMs to test whether (1) cuckoo parasitism (1 = parasitized, 0 = non-parasitized; $n=335$), (2) host rejection

of parasitic eggs (1 = rejected, 0 = accepted; $n=116$ parasitized clutches) and (3) cuckoo fledging success (1 = fledged, 0 = not fledged; $n=50$ nests with accepted cuckoo eggs) differ in relation to nest status (monogamous, primary and secondary; a fixed effect) and laying date (date of the first egg laid per clutch, a covariate). To control for the possible relationship between nest status and laying date, we centred the laying date around the mean in each nest status category. Apart from laying date, in model 2, we also used actual number of host eggs in the nest at time of parasitism (as a covariate), as this factor was shown to have a pronounced effect on great reed warbler egg-rejection rates (Moskát and Hauber 2007; Moskát et al. 2010). To cope with the issue of pseudoreplications, in models 1 and 3, we included both male and female identity as random effects. Only female identity as a random effect entered the model 2 because it is the female who is responsible for the rejection of parasitic eggs in the great reed warbler (Požgayová et al. 2009, 2011). As we did not have any year- and site-specific predictions, we included year and study site as categorical random effects (in all models) to take potential spatiotemporal variation in the data into account. We fitted the models with binary response variables using the `glmer` function from the package `lme4` (Bates et al. 2008) and estimated the GLMM parameters using Laplace approximation (Bolker et al. 2009).

Based on the initial models 1–3 (Table 1), we specified sets of candidate models with all possible combinations of fixed effects and identical random effects. Then, we determined the best-fit model based on Akaike's information criterion (AIC), corrected for low sample size (AIC_c) where appropriate, following the recommendations of Burnham and Anderson (2002). Models were ranked from the best to the worst using Δ_i ($\Delta_i = \text{AIC}_{c(i)} - \text{AIC}_{c(\min)}$), and the Akaike weights (w_i) were calculated to give the relative support for a given model compared with the others. For the top candidate models that provided substantial support ($\Delta_i \leq 2$), we applied model averaging to identify the relative importance of each model term in predicting the response variable and to estimate effect sizes of the predictors. Model selection and averaging procedures were carried out in the package `MuMIn` (Bartoń 2011). All statistical analyses were performed in R 2.12.0 (R Development Core Team 2010).

Results

Out of 335 great reed warbler nests, 128 (38 %) were parasitized. Hosts rejected the parasitic egg in 66 (57 %) nests, most frequently by ejection (45 cases). In 20 cases, the parasitized clutch was deserted; one cuckoo egg was buried into nest lining. The cuckoo egg was accepted in 50 (43 %) nests; in 12 other nests, host reaction could not be

Table 1 Fixed effects from the initial generalised linear mixed-effects models

| | | Estimate | SE | Z value | P value |
|--------------------------|-----------|----------|-------|---------|---------|
| Model 1 | | | | | |
| Intercept | | -0.605 | 0.146 | -4.132 | 0.000 |
| Nest status ^a | Primary | 0.154 | 0.364 | 0.424 | 0.672 |
| | Secondary | 0.299 | 0.303 | 0.988 | 0.323 |
| Laying date ^b | | -0.011 | 0.010 | -1.079 | 0.281 |
| Model 2 | | | | | |
| Intercept | | -0.165 | 0.507 | -0.325 | 0.745 |
| Nest status ^a | Primary | -0.950 | 0.617 | -1.539 | 0.124 |
| | Secondary | -0.623 | 0.491 | -1.269 | 0.204 |
| Laying date ^b | | -0.045 | 0.019 | -2.388 | 0.017 |
| No. of eggs ^c | | 0.274 | 0.167 | 1.643 | 0.100 |
| Model 3 | | | | | |
| Intercept | | 1.280 | 0.471 | 2.717 | 0.007 |
| Nest status ^d | Primary | -1.530 | 0.859 | -1.781 | 0.075 |
| | Secondary | -2.047 | 0.809 | -2.530 | 0.011 |
| Laying date ^b | | 0.012 | 0.032 | 0.388 | 0.698 |

Model 1 cuckoo parasitism ~ nest status + laying date + (female identity) + (male identity) + (site) + (year), model 2 egg rejection ~ nest status + laying date + number of host eggs in nest + (female identity) + (site) + (year), model 3 cuckoo fledging success ~ nest status + (female identity) + (male identity) + (site) + (year). Variable names in parentheses denote random effects. Note that estimates of nest status (categorical predictor with three levels: monogamous, primary, secondary) in the models are in the form of treatment contrasts

^a During egg laying and early incubation stages

^b Clutch initiation date centred around zero (see “Methods”)

^c Number of host eggs at time of parasitism

^d In nestling phase

evaluated due to nest failure within our temporal criterion of egg acceptance. For site- and year-specific data, see Table 2.

Incidence of cuckoo parasitism was best explained by two models: one that included no fixed term and the other that included only laying date. However, model averaging showed that laying date was not an important predictor of cuckoo parasitism (95 % confidence interval (CI) of its model-averaged estimate contained zero, results not shown).

Two models explained the variation in host rejection of parasitic eggs. The best model included only laying date, and the second one, laying date and host clutch size at time of parasitism. Model averaging showed that only laying date was important in predicting host egg-rejection behaviour (model-averaged estimate \pm SE = -0.043 ± 0.019 , 95 % CI -0.080 to -0.006). Thus, host egg-rejection probability decreased with advancing clutch initiation date. Host clutch size at time of parasitism was not important as 95 % CI of its model-averaged estimate included zero.

Only one model substantially explained the variation in cuckoo fledging success. This model contained only nest status and showed that cuckoo young were less likely to fledge from nests of polygynous than monogamous males (estimate \pm SE = -1.805 ± 0.671 , 95 % CI -2.912 to -0.442). The most pronounced differences in cuckoo fledging

success were between monogamous and secondary nests, monogamous nests being more than twice as successful as secondary nests (Table 3, Fig. 1). Moreover, there was a tendency to lower cuckoo success in primary compared to monogamous nests and no differences between primary and secondary nests (Table 3, Fig. 1).

Out of 50 nests with accepted cuckoo eggs, 30 (60 %) were successful. Of the seven unsuccessful monogamous nests, three nest failures were due to predation, one due to starvation and three cases of nest failure were due to other causes. Of the five unsuccessful primary nests, two were predated, two cuckoo chicks starved to death and one case of nest failure was due to other causes. Of the eight unsuccessful secondary nests, four were predated, two cuckoo chicks starved to death and two cases were due to other causes. There were no significant differences among the three category of nests with respect to the cause of nest failure (Fisher exact probability test: $P=0.822$).

Discussion

In the present study, we detected 60 % overall fledging success of cuckoo young in nests of their great reed warbler

Table 2 Site- and year-specific parasitism rates and host responses towards cuckoo eggs

| | Year | Parasitism rate | Ejection rate | Desertion rate | Others ^a |
|--------|------|-----------------|---------------|----------------|---------------------|
| Site 1 | 2009 | 19/59 | 9/19 | 1/19 | 0/19 |
| | 2010 | 16/43 | 6/16 | 7/16 | 0/16 |
| Site 2 | 2009 | 43/121 | 15/43 | 6/43 | 4/43 |
| | 2010 | 50/112 | 15/50 | 6/50 | 9/50 |

^aThese include one case of egg burial and 12 instances when host responses could not be determined (see the text)

hosts, which is comparable to 69.2 % reported by Kleven et al. (2004). Nevertheless, we found high variation in cuckoo fledging success with respect to the host social mating status. As we predicted, cuckoo chicks in nests of monogamous males had higher fledging success than chicks in the nests of polygynous males. We found that cuckoos had more than two times higher fledging success in monogamous nests than in secondary nests (i.e. 76 and 33 %, respectively; Fig. 1). However, cuckoo fledging success between monogamous and primary nests only tended to differ, and we detected no difference when comparing primary and secondary nests.

In the light of studies conducted by Sejberg et al. (2000) and Trnka and Prokop (2010), we suppose that lower fledging success of cuckoos in secondary nests may result from their higher mortality, caused by starvation and predation due to insufficient assistance with feeding of young or nest defence rendered by polygynous males. Starvation of cuckoo chicks may, however, also be a result of host anti-parasite defence (Grim et al. 2003; Grim 2007). Although it may hardly apply to the cuckoos raised only in secondary nests (see “Results”), this possibility should be tested more rigorously in future studies. In addition, differences in growth rates among the cuckoo chicks raised in nests of different status may also stand behind lower cuckoo fledging success in the secondary nests. Neither our study nor the others, unfortunately, have explored growth rates, parental food provisioning and defence of cuckoo chicks in host nests of different status. Only Honza et al. (2010) showed that great reed warblers defended cuckoo chicks with similar intensity as their own offspring, but they did not consider host social mating status. Moreover, there may be other variables related to host social mating status that could explain our results,

Table 3 The most likely model explaining the variation in cuckoo fledging success in great reed warbler nests of different status

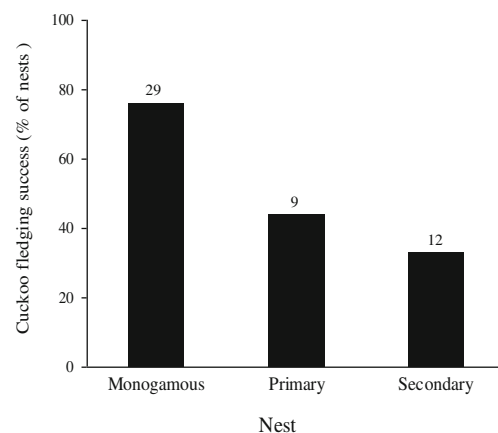
| | Estimate | SE | Z value | P value | |
|--------------------------|-----------|--------|---------|---------|-------|
| Intercept | 1.269 | 0.468 | 2.713 | 0.007 | |
| Nest status ^a | Primary | -1.517 | 0.853 | -1.778 | 0.075 |
| | Secondary | -2.033 | 0.805 | -2.525 | 0.012 |

Note that estimates of nest status (categorical predictor with three levels: monogamous, primary, secondary) in the models are in the form of treatment contrasts. Treatment contrast between primary and secondary nests: -0.517 ± 0.967 (SE), $Z=-0.534$, $P=0.593$

^a In nestling phase

including female physical condition or phenotypic quality, female age, nest location or territory quality (Verner and Willson 1966; Orians 1969; Slagsvold and Lifjeld 1994; Forstmeier et al. 2001a, b; Grønseth et al. 2003). Such relationships, however, have not been found either in our or other great reed warbler populations (Leisler et al. 1995; Bensch 1996; Hansson et al. 2000; Honza et al. 2011; Trnka and Prokop 2011; Trnka et al. 2012, this study). Therefore, to support our assumptions mentioned above, further field studies are needed. Only carefully designed experiments may discern between the contribution of different components of host parental care to the differential survival rates of cuckoo nestlings in nests of monogamous and polygynous hosts. Such studies would greatly improve our understanding of the costs of polygyny for the brood parasites as well as for their hosts.

In contrast to the variation in fledging success of cuckoos fostered by hosts of different social mating status, rejection rates of cuckoo eggs were similar on monogamous, primary and secondary nests. We found, however, that early breeders (regardless of their mating status) exhibited higher egg-rejection rates than later breeding birds. Previous studies on the closely related oriental reed warbler (*Acrocephalus orientalis*) demonstrated that rejection rates of parasitic eggs relate to host age (Lotem et al. 1992, 1995). Young breeders are usually naive (inexperienced), start breeding later in the season (i.e. they have lower re-nesting potential) than old and more experienced birds. Thus, younger great reed warbler females could be also more likely to accept cuckoo eggs than older females. However, due to the difficulty of ageing

**Fig. 1** Cuckoo fledging success in great reed warbler nests of different status (total number of nests above bars)

great reed warblers after complete moult and because only a few ringed nestlings returned to our study sites, we could not determine the age of all breeding birds and thus evaluate its effect on egg-rejection behaviour. Therefore, we could not decide which factor is more relevant to egg rejection, whether host age (in terms of egg recognition abilities) or time constraints associated with late breeding.

Although several previous studies documented non-random distribution of parasitized nests within a host population (e.g. Soler et al. 1995; Avilés et al. 2006; Polačiková et al. 2009, but see Antonov et al. 2012), we did not find any differences in parasitism rates among the nests of different status. This seems to contradict the previous study of Trnka and Prokop (2011), where nests of polygynous great reed warbler males suffered more from cuckoo parasitism than monogamous nests. However, Trnka and Prokop (2011) considered only successfully parasitized nests, i.e. those where the cuckoo chicks hatched and evicted host offspring. In the present study, on the other hand, we investigated the levels of initial cuckoo parasitism (i.e. the percentage of naturally parasitized nests) in monogamous and polygynous hosts. When we re-analysed both data sets using the same measure of parasitism (i.e. the rate of successful cuckoo parasitism), we found the same pattern as above. In Trnka and Prokop's study, successful cuckoo parasitism differed with nest status (Fisher exact probability test: $P=0.003$), while in the present study, it did not ($\chi^2=4.22$, $df=2$, $P=0.121$). It seems that these results are driven by the site-specific differences as rates of successful parasitism differed at site 1 ($\chi^2=14.61$, $df=2$, $P<0.001$; data from both studies pooled), but not at site 2 ($\chi^2=1.96$, $df=2$, $P=0.375$).

Considering that cuckoo females that parasitize monogamous great reed warbler nests achieve much higher breeding success than those parasitizing nests of polygynous males, we suppose that monogamous pairs are more suitable as cuckoo hosts than polygynous. However, cuckoos most probably do not differentiate between the nests of different status and parasitize them perhaps following a simple nest-visibility rule (see also Moskát and Honza 2000; Avilés et al. 2009; V. Jelínek et al. in prep.). Nonetheless, our study suggests that the actual level of social polygyny in a host population may considerably influence the overall reproductive success of a local cuckoo population. Therefore, social mating status of a host may be an important confounding variable affecting the outcomes of studies on cuckoo parasitism and population dynamics of cuckoo–host associations and, as such, should be controlled for in field experiments.

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Ethical standards The work described here was done under licence and complied with the current laws of the countries in which it was performed. Licences to conduct the research and bird ringing in Slovakia were issued by the Ministry of Environment of the Slovak Republic (licences number 269/132/05-5.1pil and 7230/2008-2.1pil). The fieldwork in the Czech Republic adhered to the Animal Care Protocol of the Academy of Sciences of the Czech Republic (licence number 0008/98-M103) and current Czech Law on the Protection of Animals against Mistreatment.

Conflict of interest The authors declare that they have no conflict of interest.

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