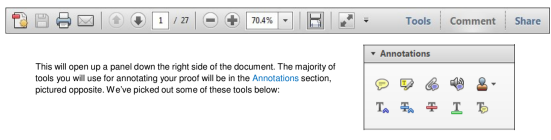
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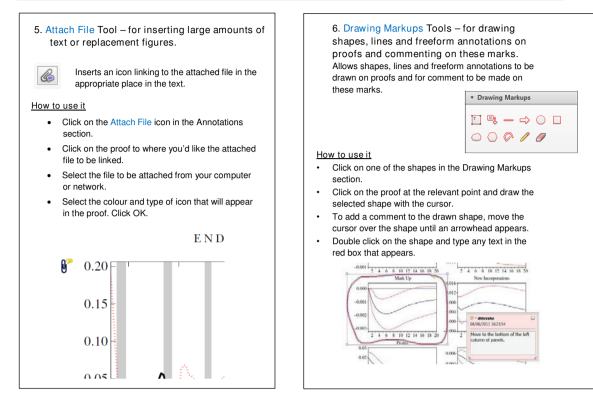
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Biological Journal of the Linnean Society, 2015, ••, ••-••.

Disappearance of eggs from nonparasitized nests of brood parasite hosts: the evolutionary equilibrium hypothesis revisited

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Received 28 April 2015; revised 15 October 2015; accepted for publication 15 October 2015

The evolutionary equilibrium hypothesis was proposed to explain variation in egg rejection rates among individual hosts (intra- and interspecific) of avian brood parasites. Hosts may sometimes mistakenly reject own eggs when they are not parasitized (i.e. make recognition errors). Such errors would incur fitness costs and could counter the evolution of host defences driven by costs of parasitism (i.e. creating equilibrium between acceptors and rejecters within particular host populations). In the present study, we report the disappearance of host eggs from nonparasitized nests in populations of seven actual and potential hosts of the common cuckoo *Cuculus canorus*. Based on these data, we calculate the magnitude of the balancing parasitism rate provided that all eggs lost are a result of recognition errors. Importantly, because eggs are known to disappear from nests for reasons other than erroneous host rejection, our data represent the maximum estimates of such costs. Nonetheless, the disappearance of eggs was a rare event and therefore incurred low costs compared to the high costs of parasitism. Hence, costs as a result of recognition errors are probably of minor importance with respect to opposing selective pressure for the evolution of egg rejection in these hosts. We cannot exclude the possibility that low or intermediate egg rejection rates in some host populations may be caused by spatiotemporal variation in the occurrence of parasitism and gene flow, creating a variable influence of opposing costs as a result of recognition errors and the costs of parasitism.

2 ADDITIONAL KEYWORDS: co-evolution – cuckoo – fitness cost – host defence – host-parasite interactions.

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[†]In memory of A. Antonov, who died much too young, 20 May 2012.

INTRODUCTION

In hosts of avian brood parasites, the costs of parasitism impose strong selection for the evolution of defensive traits because successful parasitism in many cases leads to a total loss of host reproductive output. Therefore, many hosts have evolved finetuned egg rejection abilities, which, in several cases, have been countered by parasites evolving mimetic eggs (Payne, 1967; Brooke & Davies, 1988; Davies & de Brooke, 1989a; Moksnes et al., 1991; Antonov et al., 2006a; Starling et al., 2006; Spottiswoode & Stevens, 2010; Stoddard & Stevens, 2010, 2011; Begum et al., 2011). In some species, there is apparently no variation in rejection abilities either within or between populations, and rejection rates are more or less fixed at 100% (i.e. all individuals are capable of rejection) as long as the appearance of the parasitic egg is cognitively recognizable for the individuals in question (Stokke, Moksnes & Røskaft, 2005). However, even rejection rates of non-mimetic eggs are only moderate in several other host species, often showing prominent temporal and/or spatial variation (Brooke, Davies & Noble, 1998; Soler et al., 1999; Stokke et al., 2008), relying on additional cues other than egg appearance when deciding to reject parasitic eggs (i.e. conditional responses; Brooke et al., 1998; Davies, 2000). Such co-occurrence of acceptors and rejecters (either as fixed or flexible strategies) within a single host population is often explained by the costs of making errors in the recognition and rejection of foreign eggs, which could outweigh the benefits of egg rejection under specific circumstances (Rothstein, 1982a; Marchetti, 1992; Lotem, Nakamura & Zahavi, 1995; Davies, Brooke & Kacelnik, 1996; Takasu, 1998). Specifically, host individuals attuned to reject foreign eggs may mistakenly reject one of their own eggs in nests that are not parasitized (Stokke et al., 2005; Røskaft et al., 2002a) or own eggs instead of parasitic eggs in parasitized nests. These costs obviously have fitness consequences because the resulting clutch size will be smaller than the original one. In theory, such errors are most likely to occur in hosts that have high variation in egg appearance within clutches (Davies & de Brooke, 1989b; Lotem et al., 1995; Stokke et al., 2007), making it difficult for hosts to recognize and hence reject parasitic eggs. Thus, if parasitism rates are low or variable, the costs of making recognition errors could potentially be higher than the costs of parasitism. Such a scenario could result in equilibrium between acceptors and rejecters within particular host populations or, if costs as a result of recognition errors and rejection costs are high, could even lead to acceptance being the optimal strategy

(the evolutionary equilibrium hypothesis; Lotem & Nakamura, 1998).

However, the importance of recognition errors as an opposing selective force to egg rejection, at least in hosts of evicting parasites, has been questioned (Røskaft et al., 2002a; Stokke et al., 2002). In brood parasites such as honeyguides (Indicatoridae) and cuckoos (Cuculidae), the parasitic chick gets rid of all host eggs or young from the nest soon after hatching (Davies, 2000; Anderson et al., 2009; Grim et al., 2009; Spottiswoode & Koorevaar, 2012), enforcing high costs on host reproduction and hence strong selection for the evolution of defences against parasitism (Medina & Langmore, 2015), depending on the level of parasitism (Davies et al., 1996). Furthermore, estimating the occurrence of recognition errors is not straightforward. It is well known that partial egg losses may be a result of jostling or partial predation (Rothstein, 1982b; Lerkelund et al., 1993). Hence, only the constant monitoring of nests throughout the egg-laying and incubation period can provide us with evidence for the occurrence of recognition errors. Without such monitoring, we cannot rule out the possibility that eggs may disappear for other reasons than erroneous host rejection and estimates of such costs are therefore in many cases probably higher than what is actually the case.

The present study aims to report the disappearance of own eggs in actual and potential host species of common cuckoo *Cuculus canorus*. Based on these data, we estimate the maximum costs of recognition errors and the parasitism rate that should balance these costs. We discuss our results in relation to current knowledge of co-evolutionary adaptations in common cuckoos and their hosts.

MATERIAL AND METHODS

EMPIRICAL DATA ON THE DISAPPEARANCE OF EGGS AND OTHER RELEVANT FACTORS

Data on the disappearance of own eggs in nonparasitized nests were retrieved from our own field studies in which nests were monitored throughout the egg-laying and incubation periods. We retrieved (1) number of host eggs lost from unparasitized host nests in the specific population; (2) mean clutch size in the specific population (F); and (3) rejection rate of experimentally added, non-mimetic eggs within the study populations. In addition, we also retrieved (4) the observed parasitism rate within each population (Table 1). The number of eggs lost in each population, termed the 'number of disappeared eggs', was calculated as the number of eggs lost from nonparasitized nests divided by total number of

Species	Locality	Number of disappeared eggs	RE1 (%)	RE2 (%)	RE3 (%)	RE4 (%)	Parasitism rate (%)	Rejection rate, RE4 (%)	Clutch size	Balancing parasitism, RE1	Balancing parasitism, non-mimetic eggs, Balancing parasitism, non-mimetic eggs Balancing parasitism, RE2	Balancing parasitism, RE3	Balancing parasitism, RE4
Acrocephalus	Hungary	4(54)	1.53(54)	3.07 (54)	6.13 (54)	1.61 (54)	59.4 (546)	95.2 (58)	4.83 (137)	1.9	3.9	7.7	2.0
arundınaceus Acrocephalus	Spain	6 (86)	1.50(86)	3.01(86)	6.01 (86)	$1.58(86)^{**}$	12.1 (116)	NA	4.64(59)	1.9	3.8	7.7	2.0^{**}
arunainaceus Acrocephalus	Spain	17~(229)	2.09 (229)	4.18(229)	8.36 (229)	16.73(229)	2.0 (402)	12.5(16)	3.55(273)	2.9	5.8	11.6	23.3
scirpaceus Acrocephalus	Bulgaria	1(157)	0.14 (157)	$0.27\ (157)$	0.54~(157)	0.15 (157)	28.3 (532)	88.2 (17)	4.68(159)	0.2	0.3	0.7	0.2
palustris Hippolais	Bulgaria	1(113)	0.23~(113)	$0.46\ (113)$	0.92~(113)	$0.25\ (113)$	26.6 (128)	91.3 (23)*	3.85(113)	0.3	0.6	1.2	0.3
pautaa Fringilla	Norway	0 (38)	0.00 (38)	0.00 (38)	0.00 (38)	0.00 (38)	0.0 (220)	87.0 (152)	4.75 (92)	0.0	0.0	0.0	0.0
coereos Fringilla	Norway	0 (57)	0.00 (57)	0.00 (57)	0.00 (57)	0.00 (57)	0.0(250)	78.4 (37)	6.02~(189)	0.0	0.0	0.0	0.0
montifringilla Miliaria calandra	Bulgaria	1 (118)	0.17 (118)	0.34~(118)	0.69 (118)	0.30 (118)	14.6 (356)	56.5 (108)	4.93 (119)	0.2	0.4	6.0	0.4

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nonparasitized nests (excluding nests that were naturally or experimentally parasitized). We only included nests with single host eggs lost because the disappearance of more than one egg could indicate partial predation rather than actual errors in recognition. Hence, in our marsh warbler Acrocephalus palustris population, we omitted cases where three out of four eggs (N = 1) and four out of five eggs (N = 2) disappeared. In our corn bunting *Miliaria* calandra population, we omitted cases where two out of three eggs (N = 1), three out of five eggs (N = 1). and four out of five eggs (N = 1) disappeared. We also omitted clutches where egg-laying did not follow the 'one egg per day' criterion. More specifically, this refers to two extraordinary cases in the chaffinch Fringilla coelebs population, with an irregular egg laying sequence (Stokke et al., 2002). Furthermore, the loss of all eggs in the clutch was considered to be caused by predation, and such nests were omitted from the calculations.

Nests were monitored daily from nest building until 6 days of incubation to allow estimates of disappearance of own eggs from nonparasitized nests. Eggs were marked with permanent ink in the sequence they were laid. Clutch size was estimated from completed, nonparasitized clutches. Nests used to calculate recognition errors and clutch size were different from those used to calculate rejection rate of experimentally added, non-mimetic eggs.

The disappearance of eggs from nonparasitized nests was estimated in eight study populations (Table 1): (1) great reed warblers Acrocephalus arundinaceus in Apaj, Hungary (1998-2007); (2) great reed warblers in Embalse del Hondo, Alicante, Spain (1995-2006); (3) reed warblers Acrocephalus scirpaceus in Embalse del Hondo, Alicante, Spain (1995-2006); (4) marsh warblers in Zlatia, Bulgaria (2002-2009); (5) olivaceous warblers *Hippolais pallida* in Zlatia, Bulgaria (2001–2009); (6) chaffinches in Stjørdal, Norway (1999-2001); (7) bramblings Fringilla montifringilla in Tana, Norway (2003-2004); and (8) corn buntings in Zlatia, Bulgaria (2002-2009). All these species are known to be parasitized by common cuckoos to various extents (Moksnes & Røskaft, 1995). Data on clutch sizes, parasitism rates, and rejection rates of experimentally added, non-mimetic eggs were retrieved from the literature for the same populations from which we obtained data on the disappearance of eggs (Moksnes et al., 1991; Moksnes, Røskaft & Solli, 1994; Bártol et al., 2002; Moskát & Honza, 2002; Stokke et al., 2002, 2004; Antonov 4 et al., 2006a,b, 2007a,b; Antonov et al. 2009; Hauber, Moskát & Bán, 2006; Moskát et al., 2008a,b,c, 2009; Avilés et al., 2009; Vikan et al., 2009, 2010, 2011). A few data from unpublished studies were also

included; in the Spanish reed warbler population (number 3 in the list above), rejection data of nonmimetic eggs refer to experimentally added eggs painted pale blue (i.e. resembling eggs laid by cuckoos utilizing redstarts *Phoenicurus phoenicurus*; Brooke & Davies, 1988).

E STIMATE OF PARASITISM RATE BALANCING MAXIMUM RECOGNITION ERRORS ESTIMATES

We acknowledge that the response to a parasitic egg may be conditional/plastic (Brooke et al., 1998: Lindholm & Thomas, 2000; Soler, Martín-Vivaldi & Fernández-Morante, 2012). Furthermore, we realistically assume that host egg rejection behaviour has a genetic basis (Martín-Gálvez et al., 2006). In the absence of parasitism, the frequency of rejecters in the population may decline as a result not only of recognition errors, but also other costs related to maintaining specific traits or because of stochasticity (Lahti, 2005, 2006). In populations experiencing parasitism above a certain threshold level, selection will likely lead to rejecters producing more offspring than acceptors because rejecters escape the costs of parasitism. We use the model presented by Davies & de Brooke (1989b) to derive the mean reproductive success of acceptor and rejecter pairs (RS_{acceptor} and RS_{rejecter}) and the corresponding balancing parasitism rate, p^* . Let p be the parasitism rate (probability of a host nest being parasitized). The mean reproductive success of acceptor pairs (both sexes are acceptors) is:

$$RS_{\text{acceptor}} = F(1-p) + c(F-1)p \tag{1}$$

where F is the mean host clutch size and c is the proportion of host young reared together with a parasitic chick ($0 \leq c \leq 1$). Typically, for evicting brood parasites such as *Cuculus* cuckoos, c = 0 (but see also Rutila, Latja & Koskela, 2002), whereas, for non-evicting parasites, c can be larger. We assume that cuckoo females remove one host egg from the nest when parasitizing the nest (Davies, 2000), even though the removal of more than one egg is not uncommon (Øien *et al.*, 1998). The mean reproductive success of rejecter pairs (at least one breeding individual is rejecter), where all parasite eggs are rejected is:

$$RS_{\text{rejecter}} = (F - \Delta F)(1 - p) + (F - \Delta F - 1)p \qquad (2)$$

where ΔF denotes recognition errors expressed as the number of host eggs lost by rejecters. Because only rejecters are likely to commit recognition errors, we need to take egg rejection rates within the population into account when calculating ΔF as:

erroneous egg rejection) show considerable variation

among species (range 0-23.3%) (Table 1). Because

only rejecters are assumed to erroneously reject own

eggs, and our estimate of recognition errors is one fixed value per population, the cost of recognition errors and the corresponding balancing parasitism rate will be higher when fewer individuals are classi-

fied as rejecters (Eqn 3). Hence, within particular

populations, estimates of recognition errors and bal-

ancing parasitism rates will generally be higher

when considering the rejection of mimetic eggs than non-mimetic eggs because the rejection rate for mimetic eggs in the population is generally lower than for non-mimetic eggs. If we assume that all individuals have the ability to reject eggs (100%

rejection rate), estimates of both recognition errors

and balancing parasitism rate can be regarded as

The balancing parasitism rates are generally in

the magnitude of 0-1.2%, except in great reed and

reed warblers, where it may reach 7.7% and 23.3%

respectively, depending on calculation of the proportion of individuals that are able to reject parasitic

eggs. In seven of the eight study populations,

observed parasitism rates are equal to or higher than

those required to balance the costs of making recog-

nition errors, indicating that egg rejection abilities should evolve and be maintained, which is in accor-

dance with the high rejection rates of non-mimetic

eggs generally found in the present study. Hence,

recognition errors should not be important as oppos-

ing selection pressure in these populations. The only exception among these seven populations is the corn

minimum estimates.

 $\Delta F = |$ Proportion of host eggs lost from unparasitized nests in population] [Mean clutch size in population]/ (3)

[Egg rejection rate in population]

At equilibrium $(RS_{acceptor} = RS_{rejecter})$, we obtain the parasitism rate, p^* , that balances the benefit of rejecting parasite eggs with the cost of rejecting own eggs in nonparasitized nests (recognition errors) as:

$$p* = \Delta F / ((F-1)/(1-c))$$
(4)

In some species, host individuals show phenotypic plasticity in their responses against parasites [based on social cues: Campobello & Sealy (2011); personality: Avilés & Parejo (2011; and perception of risk of parasitism: Welbergen & Davies (2009)] and parasitic eggs, and hosts will more likely reject eggs if they experience additional cues other than the egg itself, such as observing a cuckoo in the vicinity of the nest (Moksnes et al., 2000). Furthermore, the ability of individuals to reject parasitic eggs usually depends on the contrast between own and foreign eggs (i.e. egg mimicry) (Davies, 2000; Spottiswoode & Stevens, 2010). Hence, from Eqn 3, we obtain four estimates of ΔF ; assuming that (1) all (RE1), (2) 50% (RE2), (3) 25% (RE3), and (4) observed % (RE4) of individuals in the population are able to reject for-**5** eign eggs (where RE is the recognition error). The estimate RE4 is based on the rejection of experimental non-mimetic eggs in the specific study population (Table 1). We acknowledge that these four estimates only represent crude attempts to take phenotypic plasticity into account in our calculations but, first, we aim to keep our calculations as simple as possible, and, second, we lacked reliable quantitative estimates of phenotypic plasticity. By calculating four estimates, at least some of the phenotypic plasticity present at the individual level in specific populations is taken into account.

RESULTS

Disappearance of eggs from nonparasitized nests was most pronounced in great reed and reed warblers (7-7.4%). In the remaining species, eggs disappeared in only 0–0.9% of the nests (Table 1). Individuals in the populations included in our analyses experienced 0-16.7% loss of own eggs in nonparasitized nests depending on how rejection abilities were considered (RE1-RE4) (Table 1). Calculations of parasitism rates that would balance the costs of recognition errors (provided that all eggs lost were a result of

bunting population, which experiences a rather high parasitism rate but still only rejects non-mimetic eggs at an intermediate level. The remaining population, Spanish reed warblers, experiences a parasitism rate that is lower than the ones required maintaining rejection behaviour with all four estimates of recognition errors (Table 1).

DISCUSSION

Perceptual errors may cause costs that could oppose the evolution or maintenance of apparently optimal adaptations such as those involved in co-evolutionary arms races. Such costs may act as opposing selective pressures against the evolution of host defences against brood parasitism, as outlined in the evolutionary equilibrium hypothesis (Rothstein, 1982a; Lotem et al., 1995; Davies et al., 1996). In the present study, we have shown that the magnitudes of such errors are generally low across eight different host populations of the common cuckoo, even with our overestimated rates of recognition errors.

Furthermore, our estimates of recognition errors are comparable to those obtained from other species (Marchetti, 1992, 2000; Lotem *et al.*, 1995).

Several European passerines regarded as suitable common cuckoo hosts show strong rejection of experimentally added eggs (Davies & de Brooke, 1989a; Moksnes et al., 1991; Moskát, Szentpéteri & Barta, 2002; Honza et al., 2004; Lovászi & Moskát, 2004; Procházka & Honza, 2004; Rutila et al., 2006; Samaš et al., 2011) (Table 1). Such species are often characterized by having a low intraclutch variation in egg appearance (Øien, Moksnes & Røskaft, 1995; Soler & 6 Møller, 1996; Stokke, Moksnes & Røskaft, 2002b), reducing the risk of making recognition errors and enhancing the rejection of even moderately mimetic parasitic eggs (Stokke et al., 2007; Moskát et al., 2008a). Furthermore, several species may retain rejection behaviour in the absence of parasitism (Underwood, Sealy & McLaren, 2004; Lahti, 2006; Medina & Langmore, 2015), and even after speciation events (Bolen, Rothstein & Trost, 2000; Rothstein, 2001; Peer & Sealy, 2004), indicating that opposing selective pressures for egg rejection, such as recognition errors, are negligible in these species. Even with the existence of recognition errors, rejection behaviour may be retained without apparent interspecific parasitism for several reasons. First, there may be unaccounted benefits to egg rejection behaviour, such as resistance to intraspecific brood parasitism, that maintain rejection behaviour and even cause it to increase in frequency (Grim et al., 2011: Samaš et al., 2011). Second, interspecific parasitism may still occur at a low rate without researchers being able to detect it. Hence, parasitism by cuckoos laying non-mimetic eggs (i.e. from a gens with another main host) may happen from time to time but, because such eggs would be ejected quickly, the host population appears to be nonparasitized. Such 'accidental' layings are not uncommon (Capek, 1896; Chance, 1940), and the rate at which such events occur may be sufficient to retain rejection rates, as is apparent from the balancing parasitism rates reported in the present study. For example, Moksnes & Røskaft (1995) found 76 cuckoo eggs in chaffinches stored in European museums and, out of 58 000 cases of cuckoo parasitism from Europe collected by B. G. Stokke, 325 cases were recorded in chaffinch nests scattered all over Europe (unpubl. 7 data). The balancing parasitism rates estimated for chaffinches and bramblings in the present study are equal to zero, indicating that occasional parasitism by cuckoos is sufficient to retain high rates of egg rejection. Third, the retention of egg rejection in these species may be caused by immigration of rejecters from other populations that suffer high para-

sitism rates (Soler et al., 2001). Hence, there are

reports of chaffinches being utilized regularly by common cuckoos in the European parts of Russia (Malchevsky, 1960), and bramblings are favoured hosts in parts of Fennoscandia (Vikan *et al.*, 2011).

On the other hand, our results also indicate that perceptual errors may be influential for the evolution of egg rejection in some cases. Hence, one of our study populations experiences parasitism rates lower than the balancing parasitism rates. Reed warblers in Spain experience a parasitism rate of 2.0%, which is slightly lower than the calculated balancing parasitism rate (2.9-23.3%). In this population, egg rejection abilities should therefore deteriorate with time, based on our current estimates of recognition errors, provided that there is no immigration of rejecters from other populations, or as long as there are no large fluctuations in parasitism rate among years. Several studies have focused on the influence of recognition errors in reed warblers, although support for the importance of errors has been ambiguous (Davies & de Brooke, 1988; Davies et al., 1996; Røskaft et al., 2002a; Capek et al., 2010). This species shows marked spatial variation in egg rejection related to parasitism pressure in the specific population (Lindholm & Thomas, 2000: Stokke et al., 2008). indicating that there could be opposing selective pressures working against egg rejection in populations experiencing no or low parasitism. Alternatively, temporal variation in parasitism (Brooke et al., 1998) may also lead to the same pattern, with fluctuations in selective pressures depending on the current costs of parasitism. Furthermore, reed warblers have substantial intraclutch variation in egg appearance (Stokke et al., 1999; 2002b) and are parasitized by cuckoos laying mimetic eggs (Davies & de Brooke, 1988), making the recognition of parasitic eggs error prone. In such cases, hosts may rely on conditional stimuli in perceiving the risk of parasitism (Rothstein, 1982a; Davies & de Brooke, 1988, 1998; Stokke et al., 2005, 2007). Hence, reed warblers are more likely to reject parasitic eggs when they observe a cuckoo close to the nest, indicating an increased risk of parasitism (Davies & de Brooke, 1988; Moksnes, Røskaft & Korsnes, 1993; Moksnes et al., 2000). However, a recent study disclosed that the presence of a cuckoo does not necessarily lead to an increased risk of making recognition errors (Čapek et al., 2010). Careful investigations of reed warbler nests by utilizing video recordings should be undertaken to determine unambiguously if disappearance of eggs is a result of the erroneous rejection of own eggs.

Gene flow, not considered directly in the present study, may potentially slow down the process of evolving optimally expressed traits in particular populations or lead to local mal-adaptation (Nuismer, Thompson & Gomulkiewicz, 1999). Influx of rejecter or acceptor alleles may therefore influence the expression of egg rejection in local populations. However, this critically depends on spatiotemporal variation in selection regimes (Duffy & Forde, 2009), such as cuckoo parasitism and costs as a result of recognition errors, although, at present, such data are unavailable. Gene flow could also increase the frequency of rejecter alleles in nonparasitized or weakly parasitized populations (Røskaft et al., 2002b, 2006; Moskát et al., 2008b), thus causing errors to increase. Interestingly, there is low genetic differentiation among reed warbler populations in Europe, showing evidence of extensive gene flow among populations (Procházka et al., 2011). The intermediate rejection of non-mimetic eggs in reed warblers and possibly in corn buntings may therefore be caused by a combined effect of the costs of making recognition errors, gene flow, and spatiotemporal variation in the occurrence of parasitism, creating a mosaic of situations in which the opposing costs vary in relative magnitude. Interestingly, the few studies available on corn buntings indicate similar spatial variation in parasitism as in reed warblers. Hence, in Italy, only 1.4% (N = 208) corn bunting nests were parasitized (Campobello & Sealy, 2009), which is profoundly different from the relatively high parasitism rate at our Bulgarian study site.

It is important to acknowledge that recognition errors are probably rarer events than estimated in the present study because eggs may disappear from nests for several other reasons (Rothstein, 1982b; Lerkelund et al., 1993; Moksnes et al., 2000; Røskaft et al., 2002a). For example, Moksnes et al. (2000) and Røskaft et al. (2002a) reported that cuckoos visited and partially depredated 12% of reed warbler nests without actually parasitizing them; see also Wyllie (1975). Furthermore, cuckoos often remove one or two host eggs just prior to laying their own egg (Wyllie, 1975, 1981). If the host then rapidly ejects the parasitic egg, the loss of its own egg(s) will appear to be self-inflicted to the observer even when this was not actually the case. Even with daily monitoring of nests, such mistakes may take place, leading to overinflated estimates of recognition errors.

By using empirical data on disappearance of own eggs from nonparasitized nests, we have shown that costs opposing evolution of egg rejection in host species of the common cuckoo with intermediate-high rejection rates are generally small. Importantly, our estimates are most probably overestimates of true recognition errors, indicating that such costs in general are minute compared to the high costs of parasitism. Recognition errors appear to be most likely in species with specific characteristics such as a high intraclutch variation in egg appearance, an interme-

diate and variable rejection rate, spatiotemporal variation in occurrence of parasitism, and parasitism by brood parasites laying eggs mimicking host eggs at least to some extent, such as in reed warblers and perhaps corn buntings. Future studies on the importance of recognition errors should focus on long-term studies of 'low' and 'intermediate' rejecters at a spatiotemporal scale including several populations, thus taking gene flow into account, at the same time as also considering phenotypic plasticity in host antiparasite behaviour. By this approach, we should be able to obtain reliable estimates of variation in recognition errors, rejection rates, and parasitism rates and address the importance of the various costs for the evolution of egg rejection. Finally, the use of video cameras (Weidinger, 2010) would disclose the proportion of eggs that are actually lost by erroneous ejection of own eggs and not to other factors such as jostling or partial predation.

Feeney, Welbergen & Langmore (2014) called for a focus on the integration of co-evolved traits found at various stages in the breeding cycle to better understand interactions between brood parasites and their hosts. Our findings related to the egg-stage should be of importance for evaluating hypotheses proposing to explain variation in the expression of defences in hosts. Further research should focus on clarifying how recognition errors promote selection for low intraclutch variation, an important antiparasite defence in hosts of brood parasites, because host eggs with an extreme appearance are expected to be identified as parasitic eggs. We also suggest that future research should investigate how the frequency of recognition errors might be characteristic for specific stages of the arms race between hosts and brood parasites.

ACKNOWLEDGEMENTS

Comments made by John A. Allen and four anonymous referees significantly improved the manuscript. We would like to thank Manuel Martín-Vivaldi for access to unpublished data. Tomáš Grim provided constructive comments that significantly improved the manuscript. Permits to carry out data collection were issued by Conselleria de Infraestructuras, Territorio y Medio Ambiente in accordance with the relevant Spanish national guidelines for animal research (Real Decreto 1201/2005, de 10 de Octubre), the Norwegian Directorate for Nature Management (permit numbers 2008/1524 Art-VIID, 05/2580 ART-VI-ARES, 2007/1177 ART-VI-JAA), the Duna-Ipoly National Park in Hungary, and the Ministry of Environment and Water in Bulgaria. WL was funded by the National Natural Science Foundation of China (Nos. 31071938, 31272328, and 31472013). BGS and FF were funded by the Research Council of Norway (218144). CY was funded by the National Natural Science Foundation of China (Nos. 31101464 and 31260514) and New Century Excellent Talents in University (NCET-13-0761). CM was funded by the Hungarian National Science Fund (OTKA, No. 83217).

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